

This is a digital copy of a book that was preserved for generations on library shelves before it was carefully scanned by Google as part of a project to make the world's books discoverable online.

It has survived long enough for the copyright to expire and the book to enter the public domain. A public domain book is one that was never subject to copyright or whose legal copyright term has expired. Whether a book is in the public domain may vary country to country. Public domain books are our gateways to the past, representing a wealth of history, culture and knowledge that's often difficult to discover.

Marks, notations and other marginalia present in the original volume will appear in this file - a reminder of this book's long journey from the publisher to a library and finally to you.

Usage guidelines

Google is proud to partner with libraries to digitize public domain materials and make them widely accessible. Public domain books belong to the public and we are merely their custodians. Nevertheless, this work is expensive, so in order to keep providing this resource, we have taken steps to prevent abuse by commercial parties, including placing technical restrictions on automated querying.

We also ask that you:

- + *Make non-commercial use of the files* We designed Google Book Search for use by individuals, and we request that you use these files for personal, non-commercial purposes.
- + Refrain from automated querying Do not send automated queries of any sort to Google's system: If you are conducting research on machine translation, optical character recognition or other areas where access to a large amount of text is helpful, please contact us. We encourage the use of public domain materials for these purposes and may be able to help.
- + *Maintain attribution* The Google "watermark" you see on each file is essential for informing people about this project and helping them find additional materials through Google Book Search. Please do not remove it.
- + Keep it legal Whatever your use, remember that you are responsible for ensuring that what you are doing is legal. Do not assume that just because we believe a book is in the public domain for users in the United States, that the work is also in the public domain for users in other countries. Whether a book is still in copyright varies from country to country, and we can't offer guidance on whether any specific use of any specific book is allowed. Please do not assume that a book's appearance in Google Book Search means it can be used in any manner anywhere in the world. Copyright infringement liability can be quite severe.

About Google Book Search

Google's mission is to organize the world's information and to make it universally accessible and useful. Google Book Search helps readers discover the world's books while helping authors and publishers reach new audiences. You can search through the full text of this book on the web at http://books.google.com/

CATALYSIS

IN

ORGANIC CHEMISTRY

BY

PAUL SABATIER

MEMBER OF THE INSTITUTE
DEAN OF THE PACULTY OF SCIENCES OF TOULOUSE

Translated by E. EMMET REID

PROFESSOR OF ORGANIC CHEMISTRY
JOHNS HOPKINS UNIVERSITY



NEW YORK
D. VAN NOSTRAND COMPANY
EIGHT WARREN STREET
1922



c :·

COPYRIGHT; 1922 BY D. VAN NOSTRAND COMPANY

Printed in the United States of America

PREFACE

By his remarkable investigations on catalysis, Professor Sabatier has opened up new fields rich in scientific interest and fruitful in technical results. Catalytic hydrogenation will ever be an important chapter in chemistry. He is a teacher as well as an investigator and has done an important service in collecting from scattered sources a vast amount of information about catalysis and bringing the facts together in convenient and suggestive form in his book. I deem it a privilege to render his masterly work more accessible to English-speaking chemists.

The text and the unsigned footnotes represent Professor Sabatier's work as closely as I can make them. I have retained the characteristic italics. I have added a few notes which are signed by those responsible for them. In this connection I wish to thank my friends, among them Dr. Gibbs, Dr. Ittner, Dr. Adkins, and Dr. Richardson, for assistance, Professor Gomberg for verifying a number of Russian references, and Professor H. H. Lloyd for aid in proofreading.

To the chapter on the theory of catalysis, I have added an illuminating extension by Professor Bancroft, Chairman of the Committee on Catalysis of the National Research Council. In order to make the vast amount of detailed information in the book more readily available, I have prepared a subject index of some seven thousand entries and an author index of about eleven hundred names.

It is a pleasure to present a brief sketch of his life and abounding activities.

I have taken great pains to check the hundreds of references, but doubtless errors will be found. Corrections of any kind will be appreciated if sent me.

E. EMMET REID.

JOHNS HOPKINS UNIVERSITY, BALTIMORE, MD. August, 1921.

500141

V

• . . .

TABLE OF CONTENTS

[References are to Paragraphs]

CHAPTER I

CATALYSIS IN GENERAL

DEFINITION OF CATALYSIS HISTORICAL DIVERSITY IN CATALYSIS Homogeneous systems Heterogeneous systems Autocatalysis Negative Catalysis Stabilizers Reversal of Catalytic Reactions Reversible reactions, Limits	1 4 5 6 7 8 9 18 14
Velocity of Catalytic Reactions Influence of Temperature	23 24
Influence of Pressure	30
Influence of Mass of Catalyst	82
• •	
CHAPTER II	
ON CATALYSTS	
Solvents	36
DIVERSE MATERIALS CAN CAUSE CATALYSIS	41
Blements as Catalysts	42
Non-metals	43
Metals	50
Nickel, Conditions of Preparation	53
Copper Platinum, Various Forms used	59 61
	67
Colloidal Metals, Methods of Preparation	73
Water	73
Metallic Oxides	75
Influence of their Physical State	76
Mineral Acids	81
Bases	83
Fluorides, Chlorides, Bromides, Iodides	84
Cyanides	95
Inorganic Salts of Oxygen Acids	96
Various Compounds	104
DURATION OF THE ACTION OF CATALYSTS	111
Poisoning of Nickel	112
Poisoning of Platinum	116
Fouling of Catalysts	118
Regeneration of Catalysts	123
Mixture of Catalysts with Inert Materials	126

CONTENTS

CHAPTER III

MECHANISM OF CATALYSIS

Ideas of Berzelius	129
Physical Theory of Catalysis	131
Properties of Wood Charcoal	131
Heat of Imbibition	133
Absorption of Gas by finely divided Metals	135
Physical Interpretation of their Catalytic Rôle	138
Insufficiency of the Physical Theory	141
Chemical Theory of Catalysis	145
Reciprocal Catalysis	146
Induced Catalysis Auto-oxidations	149 150
Oxidation Catalysts	152
Catalysis with Isolatable Intermediate Compounds	156
Action of Iodine in Chlorination	156
Catalysis in the Lead Chamber	158
Action of Sulphuric Acid on Alcohols	159
Method of Squibb	161
Use of Copper in Oxidations	162
Action of Nickel on Carbon Monoxide	163
Catalysis without Isolatable Intermediate Compounds	164
Hydrogenation with Finely Divided Metals	165
Dehydration with Anhydrous Oxides	169
Decompositions of Acids	171
The Friedel and Crafts Reaction	173
The Action of Acids and Bases in Hydrolysis	175
Advantages of the Theory of Temporary Combinations	180
Theories of Catalysis by W. D. BANCBOFT	180a
CHAPTER IV	
isomerization — polymerization —	
DEPOLYMERIZATION - CONDENSATIONS BY ADDITION	N
§ 1. Isomerization	181
Changes of Geometrical Isomers	182
Changes of Optical Isomers	186
Migration of Double and Triple Bonds	190
Decyclizations	193
Cyclizations and Transformations of Rings	194
Migration of Atoms	199
§ 2. Polymerizations	209
Ethylene Hydrocarbons	210
Acetylene Hydrocarbons	212
Cyclie Hydrocarbons	216
Aldehydes	218
Aldolisation	
Polyaldehydes Passage into Esters	219
Passage into Fotors	219 222
I amage mto Dateis	219 222 225
Ketones	219 222 225 229
Ketones Nitriles and Amides	219 222 225 229 230
Ketones	219 222 225 229

CHAPTER VII

Hydration

Classification of Hydrations	305
Addition of Water	306
Ethylene Compounds	306
Acetylene Derivatives	308
Nitriles and Imides	311
Addition of Water with Decomposition, in Liquid Medium	818
Hydrolysis of Esters	313
Use of Acids	313
Use of Bases	318
Hydrolysis of Chlorine Derivatives	320 321
Ethers	322
Acetals Polysaccharides	323
Glucosides	327
Amides and their Analogs	331
Addition of Water with Decomposition, in Gaseous Medium	337
Hydrolysis of Esters	337
Ethers	338
Carbon Disulphide	339
Alcoholysis	340
	020
CHAPTER VIII	
TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	
HYDROGENATION	
Hydrogenation in Gaseous System, Generalities, Use of Nickel	342
Historical	342
Method of Sabatier and Senderens	343
Hydrogen Generator	346
Reaction Tube	347
Introduction of the Substance	350 355
Receiver for Collecting the Products	358
Hydrogenation over Nickel Duration of the Activity of the Metal	359
Choice of Reaction Temperature	361
RESULTS OF HYDROGENATION OVER NICKEL IN GASEOUS SYSTEM	366
Reduction without addition of hydrogen	367
Nitrous Oxide	368
Aromatic Alcohols	369
Phenols and Polyphenols above 250°	370
Furfuryl Alcohol	371
Carbon Disulphide at 500°	372
Reductions with Simultaneous Addition of Hydrogen	373
Oxides of Nitrogen	374
Aliphatic Nitro Derivatives	377
Aromatic Nitro Derivatives	378
Nitrous Esters	382
Oximes	383
Aliphatic Amides	386
Ethyl Aceto-acetate	387
Aromatic Aldehydes	
AIUMAMO AIUELLYUOS	388
Aromatic Ketones	389 389 391

CONTENTS	xi
Anhydrides of Dibasic Acids	392
Carbon Monoxide	393
Carbon Dioxide	395
Application to the Manufacture of Illuminating Gas	397
Aromatic Halogen Derivatives	403
Halogenated Aliphatic Acids	407
•	
CHAPTER IX	
HYDROGENATION (Continued)	
Hydrogenation in Gaseous System, Use of Nickel (Continued)	408
Addition of Hydrogen	408
1. Direct Addition of Hydrogen to Carbon	409
2. Addition to Hydrogen at Ethylene Double Bond	412
Hydrocarbons	413
Unsaturated Alcohols	416
Esters Ethers	417
Ethers Unsaturated Aldehydes	418 419
Unsaturated Ketones	420
Unsaturated Acids	422
3. Acetylene Triple Bond	423
4. Triple Bond between Carbon and Nitrogen	426
Aliphatic Nitriles	427
Aromatic Nitriles	428
Dicyanides	429
5. Quadruple Bond between Carbon and Nitrogen	430
Isocyanides	431
6. Double Bond between Carbon and Oxygen	432
Aliphatic Aldehydes	432
Aromatic Aldehydes	433
Pyromucic Aldehyde	434
Aliphatic Ketones	435 436
Cyclo-aliphatic Ketones	437
Diketones	438
Aromatic Ketones	441
Quinones	442
Ethylenic Oxides	443
7. The Aromatic Nucleus	444
Aromatic Hydrocarbons	446
Polycyclie Hydrocarbons	452
Aromatic Ketones	455
Phenols	456
Polyphenols	460
Phenolic Ethers	464
Aromatic Alcohols	465
Aromatic Amines	466 471
Aromatic Acids	471 472
8. Various Rings. Trimethylene Ring	472 472
Tetramethylene Ring	473
Pentamethylene Ring	474
Hexamethylene Ring	475
Ternenes	477

CONTENTS

Heptamethylene Ring	47
Octomethylene Ring	48
Naphthalene Nucleus	48
Anthracene Nucleus	48
Phenanthrene Nucleus	48
Pyrrol	48
Pyridine	48
Quinoline	48
Carbasol	490
Acridina	491
Acridine	492
9. Carbon Disulphide	
Hydrogenation with Decomposition	493
Hydrocarbons	493
Alcoholic or Phenolic Ethers	494
Phenyl Isocyanate	498
Amines	496
Diaso Compounds	497
Indol	497
CHAPTER X	
HYDROGENATION (Continued)	
81 Wednesday in Consum Control on Madela	498
§ 1. Hydrogenation in Gaseous System over Various Metals	499
Cobalt	
Ethylenic Hydrocarbons	500
Acetylene	501
Benzene and its Homologs	502
Aldehydes and Ketones	503
Oxides of Carbon	504
Iron	505
Ethylenic Hydrocarbons	506
Acetylene	506
Copper	507
Reduction of Carbon Dioxide	508
Nitro Derivatives	509
Nitrous Esters	513
Oximes	514
Ethylenic Compounds	515
Acetylene Compounds	518
Nitriles	521
Aldehydes and Ketones	522
Platinum	524
Combination of Carbon and Hydrogen	525
Ethylene Compounds	520
Acetylene Compounds	527
Hydrocyanic Acid	528
Nitro Derivatives	529
Aliphatic Aldehydes and Ketones	532
Aromatic Nucleus	534
Polymethylene Rings	535
Palladium	536
§ 2. Hydrogenation by Nascent Hydrogen in Gaseous System	537
Use of Alcohol Vapors	538
	539
Use of Formic Acid Vapors	540
Use of Carbon Menoxide and Water Vapor	UTU

CHAPTER XI

HYDROGENATION (Continued)	
Direct Hydrogenation of Liquids in Contact with Metal Catalyst	
Historical	54
General Conditions of the Reaction	54
§ 1. Method of Paal	54
Use of Colloidal Palladium	54
Reduction with Fixation of Hydrogen	54
Addition of Hydrogen	54
Application to Alkaloids	50
Use of Colloidal Platinum	5
§ 2. Method of Willstätter	56
Method of Operating	56
Use of Platinum Black	56
Nitro Derivatives	
Ethylene Double Bonds	
Acetylene Triple Bonds	56
Aldehydes and Ketones	
The Aromatic Nucleus	
Terpenes	
Complex Rings	
Use of Palladium Black	
Reduction of Carbonates to Formates	
Reduction of Acid Chlorides	57
Nitro Derivatives	57
Double and Triple Carbon Bonds	57
Cyclic Compounds	57
Cyclic Compounds	
Cyclic Compounds	
Cyclic Compounds	
Cyclic Compounds Use of other Metals of Platinum Group CHAPTER XII	
Use of other Metals of Platinum Group	
Use of other Metals of Platinum Group CHAPTER XII HYDROGENATION (Continued)	
Use of other Metals of Platinum Group CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal	58
Use of other Metals of Platinum Group CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.)	58
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief	58 58
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used	58 58 58
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel	58 58 58
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) § 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane	58 58 58 58 58
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds	58 58 58 58 58 58 58
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds Aldehydes and Ketones	58 58 58 58 58 58 58
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds Aldehydes and Ketones Aromatic Nucleus	58 58 58 58 58 58 58 58
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds Aldehydes and Ketones Aromatic Nucleus Terpenes	58 58 58 58 58 58 58 58
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) § 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds Aldehydes and Ketones Aromatic Nucleus Terpenes Various Rings	58 58 58 58 58 58 58 58 58
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds Aldehydes and Ketones Aromatic Nucleus Terpenes Various Rings Use of Iron	58 58 58 58 58 58 58 58 58 58 59 59
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds Aldehydes and Ketones Aromatic Nucleus Terpenes Various Rings Use of Iron Use of Copper	58 58 58 58 58 58 58 58 58 58 58 58
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds Aldehydes and Ketones Aromatic Nucleus Terpenes Various Rings Use of Iron Use of Copper Use of Other Metals	58 58 58 58 58 58 58 58 59 59 59
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds Aldehydes and Ketones Aromatic Nucleus Terpenes Various Rings Use of Iron Use of Copper Use of Other Metals 4. Hydrogenation of Liquids in Contact with Nickel under Low	58 58 58 58 58 58 58 58 59 59
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds Aldehydes and Ketones Aromatic Nucleus Terpenes Various Rings Use of Iron Use of Copper Use of Other Metals	58 58 58 58 58 58 58 58 59 59
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds Aldehydes and Ketones Aromatic Nucleus Terpenes Various Rings Use of Iron Use of Copper Use of Other Metals 4. Hydrogenation of Liquids in Contact with Nickel under Low Pressures Apparatus of Brochet	58 58 58 58 58 58 58 58 59 59 59 59
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds Aldehydes and Ketones Aromatic Nucleus Terpenes Various Rings Use of Iron Use of Copper Use of Other Metals 4. Hydrogenation of Liquids in Contact with Nickel under Low Pressures Apparatus of Brochet	58 58 58 58 58 58 58 58 59 59 59 59
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds Aldehydes and Ketones Aromatic Nucleus Terpenes Various Rings Use of Iron Use of Copper Use of Other Metals 4. Hydrogenation of Liquids in Contact with Nickel under Low Pressures Apparatus of Brochet Alleged Activity of Oxides	58 58 58 58 58 58 58 58 59 59 59 59
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds Aldehydes and Ketones Aromatic Nucleus Terpenes Various Rings Use of Iron Use of Copper Use of Other Metals 4. Hydrogenation of Liquids in Contact with Nickel under Low Pressures Apparatus of Brochet Alleged Activity of Oxides Method of Operating	58 58 58 58 58 58 58 58 59 59 59 59 59 59
CHAPTER XII HYDROGENATION (Continued) Direct Hydrogenation of Liquids in Contact with Metal Catalysts (Cont.) 3. Method of Ipatief Apparatus Used Use of Nickel Formation of Methane Ethylene Double Bonds Aldehydes and Ketones Aromatic Nucleus Terpenes Various Rings Use of Iron Use of Copper Use of Other Metals 4. Hydrogenation of Liquids in Contact with Nickel under Low Pressures Apparatus of Brochet Alleged Activity of Oxides	58 58 58 58 58 58 58 58 59 59 59 59 59

xiv	CONTENTS

Various Rings Use of Nascent Hydro	ogen in Liquid System in Contact with
	CHAPTER XIII
7	ARIOUS ELIMINATIONS
§ 1. Elimination of Ha	logens
	trogen
	7 06
§ 3. Elimination of Front	ee Carbon
Decarbonization of	Carbon Monoxide
	rbon Monoxide
	tals
§ 5. Elimination of Hy	drogen Sulphide
Mercaptans	
Formation of Thiot	nmonia
Action of Nickel on	Aliphatic Amines
Phenylation of Aron	natic Amines
	Phenylhydrasones
§ 7. Elimination of Ar	illine
	CHAPTER XIV
	DEHYDROGENATION
Historical	
Classification of De	hydrogenations
Classification of De 1. Dehydrogenation of	hydrogenations
Classification of De 1. Dehydrogenation of 2. Dehydrogenation of	hydrogenations f Hydrocarbons f Hydrocyclic Compounds
Classification of De § 1. Dehydrogenation of § 2. Dehydrogenation of Cyclohexane Compo	hydrogenations of Hydrocarbons f Hydrocyclic Compounds
Classification of De 1. Dehydrogenation of 2. Dehydrogenation of Cyclohexane Compo Hydrides of Naphth Terpenes	hydrogenations of Hydrocarbons of Hydrocyclic Compounds nunds salene, Anthracene, etc.
Classification of De 1. Dehydrogenation of 2. Dehydrogenation of Cyclohexane Componential Hydrides of Naphth Terpenes Piperidine	hydrogenations of Hydrocarbons of Hydrocyclic Compounds nunds salene, Anthracene, etc.
Classification of De 1. Dehydrogenation of 2. Dehydrogenation of Cyclohexane Componing Hydrides of Naphth Terpenes Piperidine Action of Palladium	hydrogenations of Hydrocarbons of Hydrocyclic Compounds nunds nalene, Anthracene, etc.
Classification of De \$1. Dehydrogenation of \$2. Dehydrogenation of Cyclohexane Composition of Hydrides of Naphth Terpenes	hydrogenations of Hydrocarbons of Hydrocyclic Compounds ounds nalene, Anthracene, etc.
Classification of De 1. Dehydrogenation of 2. Dehydrogenation of Cyclohexane Components Hydrides of Naphth Terpenes Piperidine Action of Palladium 3. Dehydrogenation of Mechanism of the Design of	hydrogenations of Hydrocarbons of Hydrocyclic Compounds ounds ualene, Anthracene, etc.
Classification of De 1. Dehydrogenation of 2. Dehydrogenation of Cyclohexane Components Hydrides of Naphth Terpenes Piperidine Action of Palladium 3. Dehydrogenation of Mechanism of the D Use of Copper Primary Alcohols, P	hydrogenations of Hydrocarbons of Hydrocyclic Compounds ounds salene, Anthracene, etc. of Alcohols Oecomposition of Alcohols reparation of Aldehydes
Classification of De 1. Dehydrogenation of 2. Dehydrogenation of Cyclohexane Composition Hydrides of Naphth Terpenes Piperidine Action of Palladium 3. Dehydrogenation of Mechanism of the D Use of Copper Primary Alcohols, P Secondary Alcohols,	hydrogenations of Hydrocarbons of Hydrocyclic Compounds sunds salene, Anthracene, etc. of Alcohols Decomposition of Alcohols reparation of Aldehydes Preparation of Ketones
Classification of De 1. Dehydrogenation of 2. Dehydrogenation of Cyclohexane Composition of Palladium Action of Palladium 3. Dehydrogenation of Mechanism of the Duse of Copper	hydrogenations of Hydrocarbons of Hydrocyclic Compounds ounds salene, Anthracene, etc. of Alcohols Decomposition of Alcohols reparation of Aldehydes Preparation of Ketones
Classification of De 1. Dehydrogenation of 2. Dehydrogenation of Cyclohexane Composition of Palladium 1. Priperidine 1. Action of Palladium 1. Dehydrogenation of Mechanism of the Duse of Copper 1. Primary Alcohols, P. Secondary Alcohols, Use of Nickel 1. Use of Cobalt 1	hydrogenations of Hydrocarbons of Hydrocyclic Compounds unds salene, Anthracene, etc. of Alcohols Decomposition of Alcohols reparation of Aldehydes Preparation of Ketones
Classification of De 1. Dehydrogenation of 2. Dehydrogenation of 2. Dehydrogenation of Cyclohexane Components of Naphth Terpenes Piperidine Action of Palladium 3. Dehydrogenation of Mechanism of the Duse of Copper Primary Alcohols, P. Secondary Alcohols, Use of Nickel Use of Cobalt Use of Iron	hydrogenations of Hydrocarbons of Hydrocyclic Compounds ounds alene, Anthracene, etc. of Alcohols decomposition of Alcohols reparation of Aldehydes Preparation of Ketones
Classification of De 1. Dehydrogenation of 2. Dehydrogenation of Cyclohexane Components Hydrides of Naphth Terpenes Piperidine Action of Palladium 3. Dehydrogenation of Mechanism of the D Use of Copper Primary Alcohols, P Secondary Alcohols, Use of Nickel Use of Cobalt Use of Iron Use of Palladium Use of Palladium Use of Palladium	hydrogenations of Hydrocarbons of Hydrocyclic Compounds ounds calene, Anthracene, etc. of Alcohols Decomposition of Alcohols reparation of Aldehydes Preparation of Ketones
Classification of De 1. Dehydrogenation of 2. Dehydrogenation of Cyclohexane Components Hydrides of Naphth Terpenes Piperidine Action of Palladium 3. Dehydrogenation of Mechanism of the Duse of Copper Primary Alcohols, P Secondary Alcohols, Use of Nickel Use of Cobalt Use of Iron Use of Palladium Use of Palladium Use of Palladium Use of Jinc	hydrogenations of Hydrocarbons of Hydrocyclic Compounds ounds calene, Anthracene, etc. of Alcohols decomposition of Alcohols reparation of Aldehydes Preparation of Ketones

CONTENTS	XV
Manganous Oxide Stannous Oxide Cadmium Oxide Other Oxides: their Classification Case of Methyl Alcohol Carbon \$4. Dehydrogenation of Polyalcohols \$5. Dehydrogenation of Amines Primary Amines, Return to Nitrile Secondary and Tertiary Amines \$6. Synthesis of Amines \$7. Ring Formation by Elimination of Hydrogen Use of Nickel Use of Aluminum Chloride Use of Anhydrous Oxides	672 673 674 675 676 680 681 682 683 684 684 685
CHAPTER XV	
DEHYDRATION	
Dehydration Catalysts 1. Dehydration of Alcohols Alone FORMATION OF ETHERS In Liquid Medium In Gaseous System Dehydration to Hydrocarbons Reaction in Liquid Medium Concentrated Mineral Acids Zinc Chloride Iodine Reaction in Gaseous System Elements Anhydrous Metallic Oxides Conditions which Regulate their Action Alumina Blue Oxide of Tungsten Thoria Metallic Salts Case of Benshydrol Catalytic Passage of an Alcohol to a Hydrocarbon Dehydration with Simultaneous Hydrogenation Dehydration of Polyalcohols Reaction in Gaseous System Ring Formation by the Dehydration of Polyalcohols	687 688 690 691 693 695 695 696 700 702 706 713 715 717 720 721 722 723
CHAPTER XVI	
DEHYDRATION (Continued)	ંડ
§ 2. Dehydration of Alcohols with Hydrocarbons § 3. Dehydration of Alcohols with Ammonia or Amines Reaction in Liquid System Reaction in Gaseous System Mixed Amines Alkyl-piperidines Pyrrol	728 729 729 731 738 741 742

CONTENTS

§ 4. Dehydration of Alcohols with Hydrogen Sulphide: Synthesi	
Mercaptans	
Comparison of the Activity of Various Oxides	• • • • •
§ 5. Dehydration of Alcohols with Acids: Esterification	• • • • •
Catalytic Esterification in Liquid Medium	• • • •
Use of Mineral Acids	• • • • •
Explanation of their Action	
The Case of Glycerine	
Use of Acetanhydride	• • • • •
Catalytic Esterification in Gaseous System	• • • • •
Mechanism of the Action of Oxides	• • • • •
Case of Bensoic Esters	
Use of Titania Laws of Esterification over Titania	• • • • •
Case of Formic Esters	
Esterification Rates	
Use of Berylia	
6. Dehydration of Alcohols with Aldehydes or Ketones	
Formation of Acetals	
Formation of Hydrocarbons	
CHAPTER XVII	
DEHYDRATION (Continued)	
7. Dehydration of Phenols Alone	
Preparation of Simple Phenol Ethers	
Diphenylene Oxides	• • • • •
Mixed Phenol Ethers	
§ 8. Dehydration of Phenols with Alcohols: Synthesis of A	lk y l
Phenol Ethers	
9. Dehydration of Phenols with Amines	• • • •
10. Dehydration of Phenols with Hydrogen Sulphide: Formati	ion
of Thiophenols	
11. Dehydration of Phenols with Aldehydes	· • • •
§ 12. Formation of Phenolic Glucosides	
§ 13. Dehydration of Aldehydes or Ketones	
Crotonisation of Aldehydes Alone	
Crotonization of Ketones Alone	• • • • •
Crotonisation of Aldehydes with Ketones	• • • • •
Crotonization in Gaseous System	• • • • •
Dehydration of a Single Molecule	 Calc
Condensation of Aldehydes or Ketones with Various Organic M	
cules	
14. Dehydration of Aldehydes or Ketones with Ammonia	• • • • •
15. Dehydration of Aldehydes with Hydrogen Sulphide	
§ 16. Dehydration of Amides	
Formation of Nitriles	
Transformation of Acid Chlorides into Nitriles	
§ 17. Dehydration of Oximes	
§ 18. Direct Sulphonation of Aromatic Compounds	
19. Condensations by the Elimination of Alcohol	• • • • •

CONTENTS	xvii
CONTINUE	XVII

CHAPTER XVIII

1)1	cc:c) M F	ו אנ אי	7°1 () R	TO I	- A(:	11132

Decomposition of Formic Acid	820
Dehydrogenation Catalysts	824
Dehydration Catalysts	825
Mixed Catalysts	826
Decomposition of Monobasic Organic Acids	829
Simple Elimination of Carbon Dioxide	831
Aliphatic Acids	831
Aromatic Acids	834
Simultaneous Elimination of Water and Carbon Dioxide	837
Preparation of Symmetrical Ketones	837
Use of Calcium Carbonate	839
Use of Alumina	840
Use of Zinc Oxide	841
Use of Cadmium Oxide	842
Use of the Oxides of Iron	843
Use of Thoria	
Use of Manganous Oxide	845
Use of Lithium Carbonate	846
Formation of Ketones in Liquid Medium	847
Preparation of Mixed Ketones	848
Preparation of Aldehydes	851
Decomposition of Dibasic Acids	855
Decomposition of Acid Anhydrides	857
CHAPTER XIX DECOMPOSITION OF THE ESTERS OF ORGANIC ACID	8
	_
§ 1. Esters of Monobasic Acids	858
General Mechanism of this Catalysis	859
Case of Alumina	860
Case of Thoria	861
Case of Titania	863
Case of Benzoic Esters	864 866
Formic Esters	
2. Decomposition of Esters with Ammonia	871
§ 3. Esters of Dibasic Acids	872
§ 3. Esters of Dibasic Acids	
§ 3. Esters of Dibasic Acids	
§ 3. Esters of Dibasic Acids	
\$3. Esters of Dibasic Acids CHAPTER XX ELIMINATION OF HALOGEN ACIDS OR SIMILAR MOLECULES	872
CHAPTER XX ELIMINATION OF HALOGEN ACIDS OR SIMILAR MOLECULES § 1. Separation of the Acid from a Single Molecule	872 876
CHAPTER XX ELIMINATION OF HALOGEN ACIDS OR SIMILAR MOLECULES § 1. Separation of the Acid from a Single Molecule Use of Anhydrous Metallic Chlorides	872 876 876
\$3. Esters of Dibasic Acids CHAPTER XX ELIMINATION OF HALOGEN ACIDS OR SIMILAR MOLECULES \$1. Separation of the Acid from a Single Molecule Use of Anhydrous Metallic Chlorides Mechanism of this Catalysis	872 876 876 878
CHAPTER XX ELIMINATION OF HALOGEN ACIDS OR SIMILAR MOLECULES § 1. Separation of the Acid from a Single Molecule Use of Anhydrous Metallic Chlorides Mechanism of this Catalysis Use of Oxides or Metals	872 876 876 878 881
CHAPTER XX ELIMINATION OF HALOGEN ACIDS OR SIMILAR MOLECULES § 1. Separation of the Acid from a Single Molecule Use of Anhydrous Metallic Chlorides Mechanism of this Catalysis Use of Oxides or Metals § 2. Molecular Condensations by the Elimination of a Halogen Acid	872 876 876 878 881 883
CHAPTER XX ELIMINATION OF HALOGEN ACIDS OR SIMILAR MOLECULES § 1. Separation of the Acid from a Single Molecule Use of Anhydrous Metallic Chlorides Mechanism of this Catalysis Use of Oxides or Metals § 2. Molecular Condensations by the Elimination of a Halogen Acid Alkylation of Aromatic Molecules	876 876 876 878 881 883 884
CHAPTER XX ELIMINATION OF HALOGEN ACIDS OR SIMILAR MOLECULES § 1. Separation of the Acid from a Single Molecule Use of Anhydrous Metallic Chlorides Mechanism of this Catalysis Use of Oxides or Metals § 2. Molecular Condensations by the Elimination of a Halogen Acid Alkylation of Aromatic Molecules Method of Operating	876 876 876 878 881 883 884 884
CHAPTER XX ELIMINATION OF HALOGEN ACIDS OR SIMILAR MOLECULES § 1. Separation of the Acid from a Single Molecule Use of Anhydrous Metallic Chlorides Mechanism of this Catalysis Use of Oxides or Metals § 2. Molecular Condensations by the Elimination of a Halogen Acid Alkylation of Aromatic Molecules	876 876 876 878 881 883 884

xviii CONTENTS

Synthesis of Ketones Method of Operating Results Obtained Formation of Amides Ring Formation Mechanism of the Reaction Chlorides that may be Substituted for Aluminum Chloride Formation of Aromatic Amines by Hofmann's Reaction Condensations in the Aliphatic Series § 3. Separation of Alkaline Chloride, Bromide or Iodide	891 892 893 895 896 898 899 901 902 904
CHAPTER XXI	
DECOMPOSITION AND CONDENSATION OF HYDROCARBONS	
Action of Heat on Hydrocarbons	905
Cracking	908
Case of Bensene	907
Case of Petroleum	908
Case of Solvent Naphtha	909
Action of Catalysts	910 911
Ethylene Hydrocarbons	912
Acetylene Hydrocarbons, Acetylene	913
First Kind of Reaction	914
Second Kind of Reaction	916 917
Cyclic Hydrocarbons	921
Terpenes	922
Reactions carried out in the Presence of Hydrogen	924
Case of Acetylene	925
Synthesis of Petroleums Theory of the Origin of Petroleum	926 928
Action of Anhydrous Aluminum Chlorids	929
Applications to the Treatment of Petroleum	932
Use of Finely Divided Metals	932
Use of Oxides	934
Use of Anhydrous Chlorides	935
· APPENDIX TO CHAPTERS XI AND XII	
HYDROGENATION OF LIQUID FATS	
Nature of Liquid Fats	027
Iodine Number	937 938
History of Hydrogenation	939
Catalysts	941
Nickel	941
Use of the Oxides and Salts of Nickel	943 946
Life of Catalysts	947
Neutralisation of Oils	948
Troubles with Moisture	949
Amount of Catalysts	951

CONTENTS			
Temperature	. 952		
Hydrogen	. 95		
Process of Bergius			
Volume of Hydrogen Required			
Apparatus			
Apparatus of Erdmann			
Apparatus of Schwoerer			
Apparatus of Schlinck			
Apparatus of Wilbuschewitch			
Apparatus of Ellis			
Apparatus of Kayser			
Apparatus of Woltman			
Results			
Physical Constants of Hardened Oils			

.

	ı
	ı
	ı

PERIODICALS CITED AND THEIR ABBREVIATIONS

Am. Chem. J. American Chemical Journal, Baltimore.

Annalen. Annalen der Chemie und Pharmacie (Liebig's), Leipzig.

Ann. Chim. Phys. Annales de chimie et de physique, Paris.

Arch. Pharm. Archiv der Pharmacie, Berlin.

Berichte Berichte der deutschen chemischen Gesellschaft, Berlin.

Bull. Soc. Chim. Bulletin de la Société chimique, Paris.

Caoutchoue et G. Caoutchoue et gutta-percha, Paris.

C. A. Chemical Abstracts, Columbus.

C. or Chem. Centr. Chemisches Centralblatt, Leipsig. Chem. News Chemical News (The), London.

Chem. Week. Chemisch Weekblad, Amsterdam.

Chem. Zeit. Chemiker Zeitung, Cöthen.

Compt. rend. Comptes Rendus des Séances de l'Academie des Sciences de Paris. Paris.

Dinglers Dinglers Polytechnischer Journal, Stuttgart.

Gas Light. Gas Lighting, London.

Gas. Chim. Ital. Gasetta chimica italiana, Palermo.

Gas Le Gas, Paris.

Jahresb. Jahresberichte über die Fortschritte der physischen Wissenschaften (von J. Berzelius), Tübingen.
J. Am. Chem Soc. Journal of the American Chemical Society, Easton.

J. Chem. Ind. Tokio Journal of Chemical Industry of Japan, Tokio.

J. Ind. Eng. Chem. Journal of Industrial and Engineering Chemistry, New York.

J. Chem. Soc. Journal of the Chemical Society, London.

J. Gas Light. Journal of Gas Lighting, London.

Jour. Off. Journal officiel de la République Française, Paris.

J. Pharm. Chim. Journal de Pharmacie et de Chimie, Paris.

J. Phys. Chem. Journal of Physical Chemistry, Ithaca.

J. prakt. Chem. Journal für praktische Chemie, Leipsig.

J. Russ. Phys. Chem. Soc. Journal of the Russian Physico-Chemical Society, Petrograd.

J. Soc. Chem. Ind. Journal of the Society of Chemical Industry, London.

Lincei Atti della Reale accademia dei Lincei, Rome.

Mat. grasses Matières grasses, Paris.

Monatsh. Monatshefte für Chemie, Vienna.

Nachr. Ges. der Wiss. Göttingen Nachrichten der königlichen Gesellschafft der Wissenschaften, Göttingen.

Phil. Mag. Philosophical Magazine, London.

Proc. Roy. Soc. Proceedings of the Royal Society, London.

Pogg. Ann. Annalen der Physik und Chemie (Poggendorf), Leipzig.

Quart. J. Science American Journal of Science, New Haven.

Rec. Trav. Chim. Pays-Bas Recueil des travaux chimiques des Pays-Bas, Leyden.

Rev. Mois. Revue du Mois, Paris.

Rev. gen de chim. pure et app. Revue générale de chimie pure et appliquée,

Rev. Sci. Revue Scientifique, Paris.

Sitz. Akad. Wien. Sitzungsberichte der mathematisch-naturwissenschaftlichen Klasse der kaiserlichen Akademie der Wissenschaften, Vienna.

PERIODICALS CITED AND THEIR ABBREVIATIONS xxii

Seif. Zeit. Seifensieder Zeitung.

Soc. Tech. Gas. Société technique de l'Industrie gasière, Paris.
Soc. Esp. Quim. Anales de la societad española de fisica y quimica, Madrid.
Trans. Far. Soc. Transactions of the Faraday Society, London.
Zeit. anorg. Chem. Zeitschrift für anorganische Chemie, Hamburg.

Zeit. f. Chem. Kritische Zeitschrift für Chemie, Physik und Mathematik

(Kekulé), Heidelberg and Göttingen.

Zeit. Elektroch. Zeitschrift für Elektrochemie, Halle.

Z. phys. Chem. Zeitschrift für physikalische Chemie, Leipsig.

INTRODUCTION

PAUL SABATIER

Paul Sabatier was born at Carcassonne Nov. 5, 1854. Admitted at the same time to the Polytechnic and the Normal School in 1874, he chose the latter from which he went out in 1877 receiving the highest grade in the competitive examination for agregation de physique.¹ After spending a year as Professor at the Lycée of Nîmes, he became, in October, 1878, assistant to Berthelot at the College de France. In July, 1880, he received the degree of Doctor of Science, his thesis being on Metallic Sulphides. After having been Maître de Conférence in physics in the Faculty of Sciences at Bordeaux for more than a year, he took charge, in January, 1882, of the course in physics in the Faculty of Sciences of Toulouse which he was never to leave. Taking charge of the chemistry course at the end of 1883, he was made Professor of Chemistry November 24, 1884, a position which he still occupies.

His chemical investigations are very numerous and touch various branches of that science: most of them have been published in the Comptes Rendus de l'Academie des Sciences, the Bulletin de la Société Chimique, and the Annales de Chimie et de Phisique.

His researches in physical chemistry stretch from 1879 to 1897 and comprise numerous thermochemical measurements (sulphides 1879–1881, chlorides 1889, chromates 1886, copper compounds 1896–1897, etc.), a thorough study of the velocity of transformation of metaphosphoric acid (1887–1889), studies on absorption spectra (1886 and 1894), on the partition of a base between two acids (1886–1887), etc.

In inorganic chemistry he has published numerous articles on metallic sulphides (1879–1880), the sulphides of boron and silicon (1880–1891), hydrogen disulphide (1886), the selenides of boron and silicon (1891), metallic chlorides (1881, 1894–1895), the chlorides (1881, 1888) and the bromide of copper (1896). A profound study of the oxides of nitrogen, which led to the characterization of metallic nitrides, was carried out (1897–1896) with the assistance of his pupil,

² The agregation is a competitive examination which is considered extremely difficult.

J. B. Senderens. He prepared the deep blue nitrosodisulphonic acid (1896-1897), defined the tetracupric salts (1897), and obtained the basic mixed argento-cupric salts (1897-1899) which formed the starting point for a whole series of analogous compounds which Mailhe prepared subsequently.

His investigations in organic chemistry (starting in 1897) are the most important and include the general method of catalytic hydrogenation in contact with finely divided metals, which was awarded the Nobel prize for chemistry in 1912. The experiments involved in this as well as in the inverse dehydrogenation, were carried out with the aid of his successive pupils, J. B. Senderens (1899–1905), Alfonse Mailhe (1906–1919), Marcel Murat (1912–1914), Léo Espil (1914) and Georges Gaudion (1918–1919).

The study of metallic oxides as catalysts led Sabatier with Mailhe to discover a whole series of methods of transforming alcohols and phenols into mercaptans, amines, ethers, esters, etc., and also transforming acids (1906–1914). At the same time he carried out, either with Mailhe or Murat, a large number of syntheses of hydrocarbons and alcohols of the cyclohexane series, etc. (1904–1915).

In agricultural chemistry, Sabatier has published about fifteen memoirs on various subjects as well as Lessons on Agricultural Chemistry.

The Academy of Sciences of Paris awarded him the Lacaze prize in 1897 and the Jecker prize in 1905 and elected him correspondent of the chemical section in 1901, then non-resident membre titulaire in April, 1913. Awarded the Nobel Prize in Chemistry in 1912, Sabatier received in 1915 the Davy Medal of the Royal Society of London of which he was elected a foreign member in 1918. He is also a foreign member of the Royal Institution, the Academy of Sciences of Amsterdam, the Academy of Sciences of Madrid, the Royal Society of Bohemia, etc.

Profoundly attached to Toulouse, where he belonged to various local academies, Sabatier refused to leave his University to occupy the chair at the Sorbonne left vacant in 1907 by the death of Moissan. Dean of the Faculty of Sciences since 1905, he has created the three technical Institutes of Agriculture, of Chemistry and of Electrotechnique which are thronged by a large number of students.

CATALYSIS IN ORGANIC CHEMISTRY

CHAPTER I

CATALYSIS IN GENERAL

1. By catalysis we designate the mechanism by virtue of which certain chemical reactions are caused, or accelerated, by substances which do not appear to take any part in the reactions.

A mixture of hydrogen and oxygen is stable at ordinary temperatures, but the introduction of a piece of *platinum black* causes immediate explosive combination; the platinum black is not visibly affected and can repeat the same effects indefinitely.

- 2. Hydrogen peroxide decomposes very slowly in cold water solution, a 30 volume solution requiring more than 240 hours at 17° for 50% decomposition, but the addition of 0.06 g. platinum black to 20 cc. of such a solution causes a vigorous evolution of oxygen and reduces the period of half decomposition to 8 seconds at 14°.¹ The platinum black, which does not seem to be altered, has by its presence enormously accelerated the reaction which normally takes place spontaneously but very slowly.
 - 3. Substances which provoke or accelerate reactions without themselves being altered are called *catalysts*.
 - 4. History of Catalysis. The first scientific observation of a catalytic transformation appears to be due to Kirchhof, who, in 1811, showed that mineral acids, in hot water solution, change starch into dextrine and sugar, without being themselves altered by the reaction.

A short time afterwards, in 1817, Sir Humphrey Davy * observed that a slightly heated platinum spiral introduced into a mixture of air and a combustible gas, hydrogen, carbon monoxide, or hydrocyanic acid, becomes incandescent and causes the slow oxidation of

- ¹ LEMOINE, Compt. rend., 162, 657 (1916).
- ² Kirchhof, Schweigger's Jour. 4, 108 (1812).
- ³ DAVY, H., Phil. Trans., 97, 45 (1817).

the gas. In 1820 Edmond Davy discovered that platinum black tan ignite alcohol with which it is wetted. Platinum sponge also possesses this power of provoking reactions without undergoing any appreciable change, and in 1831, Pelegrin Phillips, a vinegar manufacturer of Bristol, took out an English patent on the use of platinum sponge to oxidise by air the sulphur dioxide obtained by roasting pyrites, thus producing sulphur trioxide. This was the germ of the contact process for the manufacture of sulphuric acid, which required the labors of half a century to render it industrially practicable.

In his masterful Treatise on Chemistry, Berzelius discussed phenomena of this kind in which the presence of a material apparently having nothing to do with a reaction can yet cause that reaction to take place. Adopting a term which had been used in the seventeenth century by Libavius with a different meaning, he grouped these phenomena under the designation catalytic, from the Greek κάτα, down, and λύω loose, I unloose.

- 5. Diversity in Catalysis. The reactions in which catalysis is observed have multiplied with the advance of chemistry. They are extremely varied but can be divided into two distinct groups.
- 6. First we have catalysis in a homogeneous system, that is, where there is an intimate mixing of the various constituents, or at least between one of them and the catalyst that causes or accelerates the reaction. This is the case with the soluble ferments which are not considered in this treatise; it is also the case with water vapor in gaseous mixtures; with iodine, sulphur and various metal chlorides employed to aid chlorinations; with mineral acids in aldolization or crotonization as well as in the formation or saponification of esters; with alkalies in saponification; with ferrous or manganous salts in oxidations; with zinc chloride in the dehydration of alcohol; with mercurous sulphate in the sulphonation of aromatic compounds; with anhydrous ether in the preparation of organo-magnesium complexes; and even doubtless, in the Friedel and Crafts reaction with aluminum chloride which is partially soluble in the liquids used.
- 7. The second group is that of heterogeneous systems in which, for example, a solid catalyst is brought into contact with gaseous or liquid systems capable of reacting. It acts only by its surface, if it is compact and remains so during the reaction; by all its mass if it is

⁴ DAVY, E., Schweigger's Jour. 34, 91 (1822); 38, 321 (1823).

⁵ English patent 6,069 of 1831.

⁶ Berzellus, Traité de Chemie, I, 110 (1845).

⁷ Lebavius, Alchemia, Lib. II, vol. I, chapters XXXIX and XL, Frankfort, 1611.

porous, its surface then being extremely large as compared with its weight. The influence of the almost indefinite extension of the surface in the finely divided state is such that we are tempted to think of the catalytic activity of a material as belonging exclusively to that state (130).

8. Autocatalysis. Ostwald has designated by this term those reactions in which the products of the reactions accelerate the reactions.

Thus hydrogen and oxygen, rigorously dried, do not combine even at 1000°, but if the combination is once started, the water vapor so formed greatly favors the reaction, rendering it excessively rapid and explosive.

The decomposition of hydrogen selenide, of arsine and of stibine to are cases of autocatalysis, since the selenium, arsenic, and antimony set free accelerate the reactions when once they are started.

Pure nitric acid acts only slowly on many pure metals, silver, copper, bismuth, cadmium, and mercury, but when once started, the reaction accelerates itself because *nitrous fumes* are produced which facilitate the attack so that the reaction may become violent.¹¹

We find further examples of autocatalysis in the spontaneous changes which certain organic nitro compounds undergo, e. g. powders with nitrocellulose as a base, such as powder B; 12 these changes produce acid vapors which accelerate the decomposition.

- 9. Negative Catalysts. Certain materials, when present in a chemical system, exercise an unfavorable or retarding influence; such are negative catalysts, the presence of which increases rather than decreases the chemical friction and may sometimes even paralyze the normal play of affinities.
- 10. For the present, it is convenient to place in this class substances capable of altering positive catalysts so as to diminish their efficient action.

As early as 1824, Turner 12 observed that traces of various substances suppressed the catalytic activity of finely divided platinum and mentioned as such ammonium sulphide, carbon disulphide, and hydrogen sulphide.

- * Bodenstein, Zeit. physik. Chem., 29, 428 (1899).
- COHEN, Zeit. physik. Chem., 20, 303 (1896).
- ¹⁰ STOCK and GUTTMANN, Berichte, 37, 901 (1904). Bodenstein, Ibid, p. 1361.
- ¹¹ VELEY, Jour. Soc. Chem. Ind., 10, 204 (1891).
- 12 The French cannon powder which was used during the World War. It is pure nitrocellulose gelatinized by a mixture of 2 parts ether to 1 part alcohol.
 - 18 TURNER, Pogg. Ann., 2, 210 (1824).

In the manufacture of sulphuric acid by the contact process the presence of vapors of mercury, phosphorus, and particularly arsenic in the gas is sufficient to impair rapidly and destroy ultimately the catalytic action of the platinized asbestos.

In the use of finely divided nickel as a catalyst for direct hydrogenation, traces of chlorine, bromine, iodine, or sulphur compounds in the metal, in the hydrogen, or in the substance to be treated, suffice to prevent the reaction completely and somehow act as veritable poisons for the mineral ferment.¹⁴

Many other substances, without being toxic to the nickel, which they do not seem to injure, can retard the hydrogenation by their presence, e. g. glycerine, various organic acids, etc. Examples will be given in Chapter II (112 et seq.). In hydrogenations with nickel, the presence of small amounts of carbon monoxide in the hydrogen exercises a marked retarding influence.¹⁵ 16

11. Negative catalysts, which by their presence, stabilize a chemical system and render its transformation more difficult, have been less studied than positive, but numerous examples may be given. It has long been known that hydrogen peroxide keeps better when slightly acid. The addition of a few hundredths of one per cent of sulphuric or hydrochloric acid to a 30 volume hydrogen peroxide considerably augments its stability. Thus at 65°, pure hydrogen peroxide required 3.2 hours for 50 per cent decomposition but this was increased to 35 hours by the addition of 0.026 molecule of hydrochloric acid.¹⁷

The spontaneous oxidation of chloroform to carbonyl chloride is hindered by the presence of a little alcohol.

Hydrocyanic acid is stabilized by traces of hydrochloric or sulphuric acid. 18

In the oxidation of phenols by hydrogen peroxide in the presence of ferric chloride as catalyst, the reaction is retarded by the presence of mineral acids and even more by acetic, oxalic, and citric acids.¹⁹

The formation of the organo-magnesium halides in the Grignard reaction is retarded by the presence of anisol, ethyl acetate, chloro-form or carbon disulphide (303).

- 14 SABATIER, Berichte, 44, 1984 (1911).
- 15 MAXTED, Chem. News, 117, 73 (1918).
- ¹⁶ Numerous quantitative experiments made by the translator in the Laboratory of Colgate and Company showed that catalytic nickel for hydrogenation is more injured, in use, by carbon monoxide than by any other catalyzer poison that is apt to be present.—E. E. R.
 - 17 LEMOINE, Compt. rend., 161, 47 (1915).
 - 18 Limbig, Annalen, 18, 70 (1836).
 - 19 Colin and Sénéchal, Compt. rend., 153, 76 (1911).

In the abstraction of halogens in the Wurtz or Fittig synthesis of hydrocarbons, benzene and petroleum ether exercise an unfavorable influence (605).

In the very complex reaction of the vulcanization of rubber, in which a large number of substances have a beneficial effect (104 and 107), phenyl-hydrazine is a very marked negative catalyst.²⁰

12. Water which so often acts as a positive catalyst, can sometimes retard or even prevent reactions.

Moist hydrogen reduces nickel oxide less rapidly than dry.21

The decomposition of oxalic acid by hot concentrated sulphuric acid is impeded by the addition of very small amounts of water. The time of decomposition, under the same conditions of heating, is more than trebled by the addition of 0.05% of water, while 1% of sulphur trioxide renders the reaction tumultuous.²²

The presence of a little water retards the decomposition of diazoacetic ester in alcoholic solution.²⁸

Moisture retards the fixation of oxygen in the direct oxidation of unsaturated organic compounds in the presence of metallic catalysts.²⁴

The presence of traces of water hinders the attack on metallic aluminum by fatty acids and by methyl, butyl, amyl, and benzyl alcohols as well as by various monophenols, ordinary phenol, the cresoles and α - and β -naphthols.²⁵

13. In chemical systems in which autocatalysis takes place (8), the presence of substances which form stable compounds with the catalysts engendered during the reaction, hinders their effect. Hence such substances are stabilizers, or negative catalysts.

In the action of nitric acid on metals, various oxidising agents, hydrogen peroxide, potassium permanganate, and chloric acid are negative catalysts because they hinder the accumulation of nitrous fumes by oxidising them to nitric acid and thus preventing their action as positive catalysts.

With regard to powders having organic nitrates as bases (powder B, nitrogylcerine, etc.), all substances, such as amyl alcohol and diphenylamine, which are capable of fixing, either as salts or as esters, the acid products engendered by the slow spontaneous denitrification of such powders and which hasten their decomposition, are stabilizers.

- ²⁰ Peacher, Jour. Soc. Chem. Ind., 36, 424 (1917).
- ²¹ SABATUR and ESPIL, Compt. rend., 158, 668 (1914).
- 22 Bredig and Franker, Berichte, 39, 1756 (1906).
- ²⁸ Millar, Zeit. physik. Chem., 85, 129 (1913). Braune, Ibid., p. 170. Snethlage, Ibid., p. 211.
 - 24 FOKIN, Zeit. anorg. Chem., 22, 1451 (1909).
 - 25 SELIGMAN and WILLIAMS, J. Soc. Chem. Ind., 27, 159 (1918).

14. Inversion of Reactions. According to circumstances, catalysts are frequently able to work in inverse directions.

We have seen above (2) that platinum black thrown into hydrogen peroxide, induces its rapid decomposition with separation of oxygen. Inversely, platinum black serves to oxidise many substances, for example, alcohol which it transforms into aldehyde and acetic acid (244). It is now an oxidation catalyst and now a deoxidation catalyst.

15. At about 350°, hydrogen and iodine vapor combine rapidly in contact with platinum sponge,²⁶ and at the same temperature and with the same catalyst, hydrogen iodide is dissociated.²⁷

Finely divided metals such as nickel reduced from the oxide, readily add hydrogen to hydrogenizable substances at 180°; benzene is thus transformed into cyclohexane (446). On the contrary, the inverse effect is produced when cyclohexane vapor is passed over nickel at 300°; hydrogen is eliminated and benzene is regenerated (641).

Reduced copper which is capable of hydrogenating aldehydes to alcohols at 180° (522), dehydrogenates alcohols at 250° to produce aldehydes (653).

The direct hydrogenation of nitriles over nickel at 180° readily furnishes primary amines (426); but inversely, nickel causes the decomposition of the amines at 350° into the nitriles and hydrogen (681).

Platinum, nickel, and copper are thus catalysts of hydrogenation or of dehydrogenation as the case may be.

- 16. Phenol vapor passed over thoria at 450°, is regularly dehydrated to form phenyl oxide (786); but the same catalyst at the same temperature can bring about the splitting of phenyl oxide by water to regenerate phenol.²⁸ Hence thoria is at the same time a catalyst for hydration and for dehydration.
- 17. It is the same way with strong mineral acids, such as sulphuric and hydrochloric, which are equally capable of bringing about the addition of water as in the saponification of esters (313), or its elimination as in esterification (749).
- 18. Soluble ferments, such as *emulsine*, which are in reality true catalysts, acting in homogeneous system, easily decompose glucosides by hydration and are also capable of synthesizing glucosides by dehydration. Thus *galactose* treated with emulsine in concen-

²⁶ CORENWINDER, Ann. Chim. Phys. (3), 34, 77 (1852).

²⁷ HAUTEFEUILLE, Compt. rend., 64, 608 (1867).

²⁸ SABATTER, and ESPIL, Bull. Soc. Chim., (4), 15, 228 (1914).

trated solution condenses by dehydration into galactobiose; the latter, on the contrary, in dilute solution, is hydrated by the emulsine to regenerate the galactose.²⁹

- 19. Reversible Reactions. In any reaction in which catalysts are able to activate the transformation in the two opposite directions, there results an equilibrium, the same limit being reached from either end. The catalyst only modifies the velocity of the opposing reactions without essentially changing their character; consequently in reversible reactions, the location of the limit is not, in general, changed by the intervention of the catalyst, though the catalyst enormously shortens the time required to reach that limit.
- 20. Lemoine has verified this for hydriodic acid which immediately reaches its limit of decomposition, 19% at 350°, in the presence of platinum sponge. Without a catalyst, at the same temperature, under 2 atmospheres pressure, the limit was 18.6% but was not reached till after 250 to 300 hours.³⁰
- 21. Berthelot arrived at the same conclusions with the esterification of alcohols by acetic acid. For equivalent amounts of ethyl alcohol and acetic acid, the limit of 66.6% esterification is not attained at room temperature till after the lapse of several years of contact: on the contrary, in the presence of traces of hydrochloric or sulphuric acids, the identical limit is reached in a few hours.
- 22. An immediate consequence of the foregoing is that, in reversible reactions, the location of the limit is independent of the nature of the catalyst. This has been verified for the condensation of acetaldehyde. Whatever causes its polymerization into paraldehyde (hydrochloric acid, sulphur dioxide, oxalic acid, or zinc sulphate, etc.) always transforms the same proportion.³¹
- 23. Velocity of Catalyzed Reactions. The presence of a catalyst greatly influences the velocity of reactions. It is in order to examine the effect of:
 - 1. Temperature,
 - 2. Pressure,
 - 3. Quantity of catalyst.
- 24. Temperature. Temperature plays a capital rôle in many catalytic reactions, just as it does in most chemical changes. They do not take place except above a certain temperature; the direct hydrogenation of benzene in the presence of nickel hardly takes place
 - 29 BOURQUELOT and AUBRY, Compt. rend., 163, 60 (1916).
 - ⁸⁰ LEMOINE, Ann. Chim. Phys. (5), 12, 145 (1877).
 - ⁸¹ Turbaba, Zeit. physik. Chem., 38, 505 (1901).

at all below 70°, while that of ethylene begins as low as 30° (413), and that of acetylene goes on at room temperature (423).

The decomposition of alcohol into ethylene and water by blue oxide of tungsten commences only at about 250° (709); the dehydration of phenol to phenyl oxide by thoria requires a temperature above 400° (786).

- 25. Elevation of the temperature also increases greatly the velocity of reactions: in fact it is found that, in a large number of cases, this velocity is doubled when the temperature is raised 10°. Reactions in which catalysts intervene do not escape the general rule and are greatly accelerated by elevation of temperature which is consequently favorable, so long as it does not greatly change the mechanism of the reaction which, however, frequently happens. Thus catalytic hydrogenation is frequently replaced, above a certain temperature, by its reverse, catalytic dehydrogenation.
- 26. For example in the hydrogenation of benzene over nickel, the velocity of the formation of cyclohexane increases rapidly from 70°, where it is very slow, up to 180–200°, the most favorable temperature. From there on it decreases as 300° is approached, at which this reaction no longer takes place, cyclohexane being, on the contrary, decomposed into benzene and hydrogen or even into benzene and methane according to the equation:

$$3C_4H_{12} - 2C_4H_6 + 6CH_4$$

this latter reaction becoming more important as the temperature is raised.**

- 27. In the hydrogenation of acetylene which takes place without complications at room temperature (423), elevation of temperature tends to introduce, by the side of the transformation into ethane, the condensation of acetylene into more complex molecules even to the formation of solid carbonaceous deposits (924).
- 28. In the dehydration of primary alcohols by contact with anhydrous oxides, elevation of temperature tends to introduce or to accelerate the reaction of dehydrogenation whereby aldehydes or compounds produced from them are formed (709).
- 29. Thus, by a judicious choice of reaction temperature, it is frequently possible to obtain, at will, various degrees of combination. For example, in the hydrogenation of anthracene over nickel, at 180°, perhydroanthracene, C₁₄H₂₄, is obtained along with the dodecahydro-; at 200° the octohydro- is prepared and at 260°, the tetrahydro-.²⁸

³² SABATTER, and SENDERENS, Ann. Chim. Phys. (8), 4, 334 (1905).

³⁸ GODCHOT, Ann. Chim. Phys. (8) 12, 468 (1907).

30. Pressure. Increase of pressure can scarcely have any considerable effect except in gaseous systems or in heterogeneous systems having a gaseous phase. In such cases, it can be foreseen that it will have a beneficial effect in those cases in which the number of molecules is diminished in the reaction.³⁴ This is the case in the hydrogenation of compounds containing an ethylene bond and practical use is made of it in the hydrogenation of liquid fats (956).

Likewise in the direct hydrogenation of phenol by nickel, in the liquid system around 150°, the formation of cyclohexanol is extremely slow in hydrogen at ordinary pressure, but, on the contrary, is rapid and complete under 15 atmospheres.³⁵

- 31. On the contrary, molecular decompositions such as the dehydrogenation of alcohols into aldehydes or ketones, in contact with finely divided copper, are favored by a lowering of the pressure, which diminishes also the reverse reaction (653).
- 32. Quantity of Catalyst. We must at once distinguish between the two cases, whether the catalyst acts in homogeneous or heterogeneous systems.

In homogeneous systems, in which the catalyst remains in intimate mixture with the components of the reaction, it acts by its mass and its action increases with its concentration.

In the manufacture of sulphuric acid by the lead chamber process, in which oxides of nitrogen serve as the catalyst, the velocity is proportional to their concentration up to a certain limit.

In the inversion of sugar solutions by mineral acids (324), and in the saponification of esters by the same agents (313), the active agents in the catalysis are the free hydrogen ions arising from the electrolytic dissociation of the acids and the velocity of the reaction is proportional to the concentration of these ions.

In the catalytic decomposition of hydrogen peroxide by small amounts of alkali, the rapidity of the decomposition is nearly proportional to the concentration of the alkali.²⁶

- 33. It is the same way with certain solid catalysts, iodine in the chlorination or organic compounds (278), and anhydrous aluminum chloride in the Friedel and Crafts reaction (883), which do not act till they have been dissolved in the liquids of the system to be transformed and then are comparable to liquid catalysts, with activity proportional to their concentration.
 - 34. Heterogeneous systems are much more frequently met with:
 - ⁸⁴ DARZENS, Bull. Soc. Chim. (4), 15, 588 (1914).
 - ⁸⁵ Brochet, *Ibid.* (4), 15, 554 (1914).
 - ³⁶ LEMOINE, Compt. rend., 161, 47 (1915).

the catalyst in such is a solid phase in a liquid or gaseous medium and exercises its useful power only on its surface. The action, at first sight, depends on the extent of the surface, or at least on the mass of an extremely thin layer. A layer of silver 0.0002 mm. thick, deposited on glass, causes a very rapid decomposition of hydrogen peroxide.²⁷

35. Solid catalysts are more active the greater their surface, and, for the same weight, the finer their grains; but there is, by no means, a rigorous proportionality between the activity and the extent of surface.

In liquids, convection currents which bring the material to be transformed into more or less perfect contact with the catalysts, have an important influence on the rate of the reaction, but one difficult to estimate. If the mixture is kept perfectly homogeneous, the active surface of a given catalyst, made up of grains of the same size, should be proportional to the number of grains, that is to say, to the total mass, but should increase very rapidly as the grains become smaller.

For a solid catalyst acting in a gaseous system, the incessant and very rapid movement of the gas particles is sufficient to assure the homogeneity of the system. The activity of the catalyst, if it is in a very thin layer, is proportional to the area of this layer. If the layer is thick, not only the surface particles act but also those within, the effect of the interior particles being more important, in proportion as the grains which compose the catalytic material are lighter and less agglomerated. With a solid in a fine powder, which is readily penetrated by the gas, the useful surface is extremely large as compared with the exterior surface of the layer. The state of division of a solid catalyst is a matter of prime importance. The catalytic power of nickel in sheets or even in thin foil is quite minute and of no practical value, while it is highly developed in the finely divided nickel which is obtained by reducing nickel oxide by hydrogen, below red heat, and particularly so when the oxide obtained by dehydration of nickel hydroxide is itself finely divided.

From this point of view, there are great differences in various catalysts according to the conditions of their preparation (see Chapter II).

⁸⁷ LEMOINE, *Ibid.*, 155, 15 (1912).

CHAPTER II

ON CATALYSTS

- 36. As chemistry has developed, the number of catalytic phenomena has increased enormously and it has been recognized that the rôle of catalyst is played, not by a few bodies only but by a multitude of substances of every sort.
- 37. Solvents. The definition proposed by Ostwald, "A catalyst is a substance which, without appearing in the final product, influences the velocity of a reaction," leads us to consider an infinite number of substances as catalysts. Solvents, whatever their nature, are catalysts since they do not appear in the equation of the reaction which they cause to take place.

In the absence of a liquid which dissolves them and thus realizes the contact which is indispensable to combination, solid substances which have no appreciable vapor pressure in the cold, are incapable of reacting with each other.

Dry crystals of oxalic acid and chromic anhydride can be mixed cold without any chemical change, but the addition of water which establishes perfect contact between the two substances, immediately starts the oxidation of the oxalic acid at the expense of the chromic anhydride. The water may be recovered completely and unchanged by the reaction. It acts as a catalyst.

38. The nature of the solvent can change greatly the velocity of reactions which take place in it, and furthermore, the influence which it exercises is absolutely special in each case.

Water is a true catalyst in the decomposition of hydrogen peroxide.¹

In the fixation of hydrogen, by colloidal palladium, upon the acetylene triple bond, the solvent has an important influence of its own.

The combination of triethyl-amine with ethyl iodide to form tetraethyl-ammonium iodide at 100°, is 203 times as rapid in ethyl alcohol, 718 times in acetophenone, and 742 times in benzyl alcohol, as it is in hexane.

- ¹ Lemoine, Compt. rend., 155, 9 (1912).
- ² Zal'Kind and Pischikov, Jour. Russian Phys. Chem. Soc., 46, 1527 (1914), C. A., 9, 2067.
 - ³ MENSCHUTKIN, Zeit. phys. Chem., 1, 611 (1887); 6, 41 (1890).

- 39. In reversible reactions, the *limit* will not be altered by a change of solvent if this does not react in any way with either the reactants or the products: otherwise the limit will be modified. For example, in reactions between electrolytes, brought about in alcohol or in water, electrolytic dissociation is of great influence in case water is the solvent.
- 40. Solvents are not commonly classed with true catalysts as this designation is usually reserved for those substances which act in small concentration and of which a small quantity is able to cause large quantities of other materials to react.

DIVERSE SUBSTANCES CAN ACT AS CATALYSTS

41. The number of substances capable of acting as catalysts, is already very large and continues to increase with the progress of chemistry.

We find in this class the most varied materials: elements, oxides, mineral acids, bases, metallic chlorides, bromides, iodides, fluorides and oxygen salts, ammonia and its derivatives, and diverse organic compounds. But, particularly for solids, the catalytic activity can vary greatly according to their origin, either if they can exist in distinct molecular forms, or, more frequently, if they present themselves in different states of sub-division (32).

ELEMENTS AS CATALYSTS

- 42. Elements which are of themselves true catalysts, maintaining themselves unchanged during the course of the reactions which they provoke, are quite numerous and it is convenient to consider along with them those which pass immediately into compounds which act as catalysts. This is the case with chlorine, bromine, iodine, tellurium, sulphur, and phosphorus among the non-metals and tin, antimony, and thallium among the metals.
- 43. Chlorine and Bromine. These probably act by the immediate formation of the hydro-acids, to transform aldehydes into the polymeric paraldehydes.
- 44. Iodine. Iodine acts in the same way in the same reactions. It is frequently employed in chlorinations, and acts then by transforming itself into the *trichloride* which is the real factor in the catalysis. It permits the direct sulphuration of aromatic amines with the elimination of hydrogen sulphide (296). It can aid in causing

the condensation of aromatic amines with naphthols (790). It serves also to facilitate the reaction in the preparation of the organomagnesium halides of the Grignard reagent, when it is desired to prepare these from chlorides or bromides (302).

- 45. Sulphur and Tellurium. Employed as carriers in chlorination, they certainly act in consequence of the initial formation of an equivalent amount of the chlorides. Tellurium has been proposed as an agent in direct oxidation (251).
- 46. Phosphorus. Red phosphorus has been mentioned as a catalyst for the dehydration of alcohols above 200° (699). The chief factor in this catalysis appears to us to be the small quantity of acids of phosphorus which exist in the phosphorus or which are produced from it by the oxidising effect of the alcohol.
- 47. Antimony, Tin and Thallium. Their use in chlorination is based on the primary formation of their perchlorides.
- 48. Carbon. All the porous forms of carbon have been employed as catalysts.

The carbonaceous mass obtained by calcining blood with potassium carbonate is a good catalyst for chlorination.

Animal charcoal is a mediocre catalyst for the dehydration of alcohols (699), but is efficient in the preparation of carbonyl chloride from carbon monoxide and chlorine (282).

Coke may serve as an oxidation catalyst (258).

Wood charcoal, or baker's charcoal possesses considerable absorbing power for many gases, the consequence of which is frequently the production of special reactions. Carbon saturated with oxygen can produce oxidations: ethyl alcohol is changed to acetic acid. Ethylenic hydrocarbons are partially burned.

Carbon saturated with chlorine enables us to chlorinate sulphur dioxide in the cold as well as hydrogen.6

Baker's charcoal catalyzes the decomposition of primary alcohols above 380°, giving, at the same time, aldehydes and ethylene hydrocarbons (679). It is frequently employed for the preparation of carbonyl chloride (282).

49. The porosity of the carbon has a great influence. Thus in the case of 30 volume hydrogen peroxide of which the half decomposition at 17° required 240 hours, the addition of 5% of cocoanut charcoal (in pieces 1 to 2 mm. in size) reduced this time to 15.4 hours, while the same weight of charcoal from the black alder lowered it

⁴ DAMOISEAU, Compt. rend., 83, 60 (1876).

⁵ CALVERT, Ibid., 64, 1246 (1867).

[•] MELSENS, Ibid., 76, 92 (1873).

only to 212 hours. Sugar charcoal falls between these two as an activator.

- 50. Sodium brings about the isomerization of unsaturated hydrocarbons, e. g., diethylallene into diethylallylene (192). It polymerizes isoprene (213) as well as acetonitrile (231).
- 51. Magnesium. Magnesium powder has been mentioned as very active in decomposing hydrocarbons at 600° (918).

Aluminum. The same property has been claimed for aluminum which has been proposed as a chlorination catalyst also because it changes immediately to the chloride. Aluminum turnings are only a mediocre catalyst for oxidation (255).

52. Manganese. Powdered manganese is a poor catalyst for oxidations (255) but is an excellent aid to bromination (292).

Zinc turnings, at 100°, can cause the condensation of acetaldehyde into aldol or into crotonic aldehyde (219). The same metal acts as a dehydrogenating agent on alcohols at 600-50°, temperatures at which the metal is melted, a condition unfavorable to catalytic action (670).

- 53. Nickel. Employed in the state of extremely fine division, as is obtained by the reduction of the oxides by hydrogen or carbon monoxide, nickel is a marvelous catalyst, the manifold activity of which has been established by the investigations of Sabatier and Senderens, beginning in 1879. It is specially suitable for the direct hydrogenation of volatile organic compounds, but it is equally capable of producing dehydrogenations and decompositions whether they are followed by molecular condensations or not. Chapters VIII, IX and XII are devoted to catalytic reactions effected by nickel.
- 54. The metal in sheet or even in thin foil possesses only slight activity. Catalytic nickel should be prepared by reducing the oxide, and as the metal so produced is readily oxidised and frequently pyrophoric, it is generally best to carry out the reduction in the same tube in which the catalysis is to be effected. However this is not absolutely necessary, if the precaution is taken to cool the reduced metal perfectly in the current of hydrogen, or better still in a current of pure nitrogen. The metal so prepared can be preserved in a well-
 - ⁷ LEMOINE, *Ibid.*, 162, 725 (1916).
- ³ When freshly prepared highly active nickel is exposed freely to the air, a rapid heating takes place that considerably impairs its catalytic activity. The change which takes place in the nickel is brought about and augmented by the heat produced by the catalytic oxidation of the hydrogen occluded and surrounding the nickel when it comes in contact with an excess of oxygen from the air. Similar oxidation of hydrogen is well known in the presence of catalytic palladium or platinum. In the case of catalytic nickel, however, the heat thus

stoppered bottle for quite a long time without considerable alteration.

55. The activity of the reduced nickel varies greatly according to the nature of the oxide and the manner of reduction. The metal is more active, the greater its surface; and the lighter the oxide and the lower the reduction temperature, the greater is this surface.

Nickel reduced at a bright red is no longer pyrophoric and possesses a considerably reduced catalytic power.

On the contrary, that which comes from the hydroxide precipitated from the nitrate, dried and reduced around 250°, has an excessive activity along with maximum alterability. It can be compared to a spirited horse, delicate, difficult to control, and incapable of sustained work.

Applied to phenol, it passes by cyclohexanol and produces cyclohexane to a large extent. It tends to produce molecular dislocations in bodies submitted to catalysis.

56. An excellent quality of nickel is obtained by dissolving the commercial cubes in pure nitric acid (free from hydrochloric), calcining the nitrate at a dull red and reducing at about 300° the oxide thus obtained. Such a nickel can do all kinds of work and maintains its activity for a long time.

It has been stated that nickel prepared above 350° is incapable of hydrogenating the aromatic nucleus, but Sabatier and Espil have shown that this ability is still possessed by a nickel prepared at 700° even when it is kept at this temperature for several hours, but not by nickel prepared by reduction above 750° or heated for some time at 750°.1°

57. Cobalt. Finely divided cobalt, such as is obtained by the reduction of the oxide by hydrogen, can be employed as a catalyst for the same purposes as nickel, but is less useful as it is less active,

generated in the presence of an excess of oxygen, or air, produces an oxidation of the catalyzer to an extent that lessens or destroys its activity. A number of experiments were made in which freshly prepared nickel catalyzer still in the presence of hydrogen was subjected to the action of a Geryk pump which exhausted practically all of the excess hydrogen gas. In different experiments the catalyzer was then, while cold, allowed slowly to come in contact with carbon dioxide, nitrogen, and air. The catalyzers so formed were active and retained their activity reasonably well. In case air was admitted to the vacuum vessel containing the catalyzer, it was introduced very slowly so that any oxidation would be so slight as not to increase the temperature sufficiently to produce cumulative oxidation.— M. H. Ittner.

DARZENS, Compt. rend., 139, 869 (1904); BRUNEL, Ann. Chim. Phys. (8),
 6, 205 (1903).

¹⁰ SABATIER and Epsil, Bull. Soc. Chim. (4), 15, 779 (1914).

and as the reduction of its oxide requires a higher temperature, in fact above 400°.

58. Iron. Reduced iron can replace nickel in quite a large number of cases, but disadvantages, like those mentioned for cobalt, are more serious, the oxides being still more difficult to reduce. Between 400° and 450°, it is necessary to prolong the action of the hydrogen for six or seven hours to obtain complete reduction. Furthermore, the metal reduced at this high temperature is no longer pyrophoric and retains only mediocre activity. However, pulverized iron is a useful catalyst for decompositions accomplished at a low red heat (932).

Iron has been mentioned as a chlorination catalyst, but in that case it serves only to form iron chloride which is the real catalyst.

59. Copper. Copper, reduced from its oxide by hydrogen, constitutes, on account of its ease of preparation, the low temperature at which the oxide can be reduced, below 180°, and the regularity of its action, a valuable catalyst for certain reactions, but it is not capable of effecting all kinds. Its activity also varies considerably according to the method of production. The black oxide of copper, prepared by roasting the metal or by calcining the nitrate at a bright red, furnishes by reduction, with incandescence, a clear red, very compact metal with low catalytic power. By reducing with a slow current of hydrogen (to avoid incandescence) at about 200°, the tetracupric hydroxide — such as is precipitated from boiling cupric salt solutions by alkalies — a very light violet colored metal is obtained with much greater catalytic activity. The very fine copper powder which is commercially prepared for imitation gilding, can frequently be used: it is only necessary to free it from grease by washing with ether or ligroine.

This latter has been used to facilitate several of the reactions of aromatic diazonium salts in which nitrogen is eliminated (606). It is efficient in causing the production of phenyl oxide by the action of brombenzene on sodium phenylate (904).

Copper in spirals, or in gauze, has been employed, with advantage, in the catalytic oxidation of alcohols, ethers, hydrocarbons, and amines (254).

- 60. Silver. Silver powder is an excellent oxidation catalyst (253). Inversely, it causes the rapid decomposition of hydrogen peroxide, transforming itself into the oxide Ag₄O₈ which continues the catalysis.¹¹
 - 61. Platinum. Platinum is one of the longest known catalysts.
 - ¹¹ BERTHELOT, Bull. Soc. Chim. (2), 34, 135 (1880).

Not oxidisable in the air at any temperature, it is a powerful catalyst for oxidation or for hydrogenation, especially when it is finely divided and presents a large surface. This is the condition realized in platinum sponge, a porous material obtained by calcining ammonium chlorplatinate, and even better in platinum black and in colloidal platinum, which can be mixed intimately with liquids submitted to catalysis (67).

62. Platinum black can be prepared either by reducing acid solutions of platinic chloride ¹² by zinc, or better by magnesium, or by treating the platinum chloride with alcohol and alkalies, ¹³ or by reducing the platinum salt with sodium formate, ¹⁴ or with sodium tartrate, or even with glucose in alkaline solution, or by glycerine and potash. ¹⁵

An excellent method is that of Loew: 35 cc. formalin is added to 25 g. platinum chloride dissolved in 30 cc. water and then, little by little, while cooling 25 g. caustic soda dissolved in its own weight of water. After twelve hours it is filtered off and washed. A spongy mass is thus obtained which is dried in the cold over sulphuric acid.¹⁶

Platinum black always retains traces of substances with which it has been in contact during its preparation. Blacks prepared in alkaline solution are more active than those from acid solution.

63. According to Lemoine the grains of platinum black, of which the diameter is about 0.1 mm., are much more active than those of the sponge for the same area. With a specimen of hydrogen peroxide which, without catalyst, required ten days for half decomposition, this time was reduced by platinum black to 0.00013 hour and with the same surface of the sponge only to 0.2 hour. The black possesses a specific activity which is, without doubt, due to less molecular condensation and which disappears when it is heated to 400 to 500°.17

This weakening by heating is progressive. Thus platinum black is not sensibly altered as a hydrogenation catalyst when heated below 300° and still retains its power to transform limonene into menthane by the fixation of 2H₂. If it is heated to 430°, it is considerably weakened and can add only H₂ to the external double bond, giving carvomenthene. Heated to 500°, it loses all activity.¹⁸

- 12 BOETTGER, J. Prakt. Chem. (2), 2, 137 (1870).
- 18 Zeise, Pogg. Ann., 9, 632 (1827).
- ¹⁴ DOEREREINER, Ibid., 28, 181 (1833).
- ¹⁵ ZDRAWKOWITCH, Bull. Soc. Chim. (2), 25, 198 (1876).

¹⁶ Loww, Berichte, 23, 289 (1890). Improved directions for this important preparation are given by Willstätter and Waldschmidt-Leftz in Berichte, 54, 121 (1921).—E. E. R.

¹⁷ LEMOINE, Compt. rend., 162, 657 (1916).

¹⁸ VAVON, Ibid., 158, 409 (1914).

Compact platinum in foil or wire has a certain activity, at least, if it has been previously heated above 50°. A heated platinum spiral introduced into a mixture of alcohol vapor and air or oxygen, causes the formation of aldehyde and the incandescence which results from the heat liberated in the oxidation, maintains itself indefinitely so long as the mixture is renewed: this is the *lamp without flame*.¹⁹

64. Rhodium, Ruthenium, Iridium, and Osmium. Employed in the form of the pulverulent black, or as sponge, these metals act in the same manner as platinum, at least as regards reactions of oxidation or of decomposition, but they are less active in hydrogenation (580).

Rhodium or iridium black decomposes, in the cold, formic acid into hydrogen and carbon dioxide (822). In contact with alcohol and caustic soda, hydrogen is evolved with the formation of sodium acetate.²⁰

- 65. Palladium. Palladium exhibits the property of absorbing very large quantities of hydrogen, even up to 930 times its own volume.²¹ Palladium thus saturated with hydrogen can effect a large number of hydrogenations. But the metal can serve also as a temporary support for hydrogen, that is to say, as a hydrogenation catalyst, in the form of sponge or black (573), and can be employed as a catalyst for dehydrogenation (669), for decomposition (624), or for polymerization (212).
- 66. Gold. Gold, when finely divided, has catalytic properties resembling those of silver.
- 67. Colloidal metals. The catalytic activity of metals, being in direct relation to the extent of their surfaces, consequently to the minuteness of their particles, should reach its maximum in the colloidal state. As the chemical alterability of the metals is also intensified by their extreme subdivision, it would hardly be expected that any could be practically used in this state except those not oxidisable in the cold, such as platinum, palladium, gold and silver.
- 68. Bredig ²² has described a simple method for preparing colloidal metals: an electric arc is made to play between two wires of the metal under pure water. A sort of nebulosity is observed which becomes darker and darker till it is soon so opaque that the spark
 - 19 HOFMANN, Annalen, 145, 358 (1868).
 - ²⁰ Sainte-Claire-Deville and Debray, Compt. rend., 78, 1782 (1874).
- ²¹ Graham, Phil. Mag., (4), 32, 401 and 503 (1866); 36, 63 (1868). Proc. Roy. Soc., 15, 223, 502 (1867); 16, 429 (1868); 17, 212 and 500 (1869). Compt. rend., 63, 471 (1866) and 68, 101 (1869).
- ²² Bredig, Zeit. physik. Chem., 31, 258 (1899); 37, 1, 323 (1901); Berichte, 37, 798 (1904); Zeit. Elektroch., 14, 51 (1908).

can not be seen. Solutions thus obtained can be preserved for a long time and contain 0.09 to 0.02 g. gold per liter and a less amount of palladium or platinum: the number of particles in such a solution may reach as high as a billion per cubic millimeter.

69. Unfortunately such solutions are unstable in the presence of various substances. The presence of suitable organic materials gives them stability and Paal has found that egg albumen has this effect. He dissolves 15 parts of caustic soda in 500 parts of water, adds 100 parts egg albumen and warms on the water bath till solution is nearly complete. It is acidulated with sulphuric acid and the precipitate filtered off. The solution is neutralized with soda, evaporated on the water bath to a small volume and again acidulated with sulphuric acid.

The filtered solution is dialyzed to separate the sodium sulphate. The liquid remaining in the dialyzer is treated warm with baryta water which precipitates the remaining sulphate ions. The filtered solution is evaporated on the water bath and several volumes of alcohol added, which precipitates white flakes which Paal has named lysalbinic acid. When dry, this is a white powder, soluble in water and nearly insoluble in alcohol: its weight is about one-fourth that of the albumen.

One gram of the above product is dissolved in 30 cc. water and made alkaline with a slight excess of soda, 2 g. platinum chloride dissolved in a little water is added and then a slight excess of hydrazine hydrate. The solution turns dark and a gas is evolved: after five hours it is dialyzed to eliminate electrolytes, carefully evaporated on the water bath and dried in vacuum. Brilliant black scales are obtained which dissolve in water to form a black opaque solution: this is colloidal platinum.

Colloidal palladium is prepared in an analogous manner.²³ Solutions of these are very stable and can even be heated for a long time without change.

- 70. In this way colloidal solutions can be prepared of silver, gold, copper, osmium, and iridium, all decomposing hydrogen peroxide with extreme energy. Traces of osmium produce this effect.²⁴
- 71. Skita prepared a colloidal palladium hydroxide, for use as a hydrogenation catalyst, by heating to boiling a solution of palladium chloride, PdCl_a, with soda and a little gum arabic. The solution is
- ²⁸ Paal, Berichte, 35, 2195 (1902). Paal and Amberger, Ibid., 37, 126 (1904) and 38, 1398 (1905). Kelber and Schwartz, Ibid., 45, 1946 (1912). Seita and Meyer, Ibid., 45, 3579 (1912).
- ²⁴ Paal and American, Berichte, 40, 2201 (1907). Paal, Biehler and Steyer, *Ibid.*, 50, 722 (1917).

dialyzed till neither silver nitrate nor baryta water gives a test outside. The solution, evaporated to dryness in a vacuum, gives brown scales of colloidal palladium hydroxide, insoluble in cold water but soluble in water containing traces of acid or alkali.

Another method of preparing colloidal palladium, given by the same author, is to pass a current of hydrogen through a warm solution of palladous chloride and gum arabic.

A colloidal platinum hydroxide, analogous to that of palladium, is obtained by treating a boiling solution of potassium chlorplatinate with the theoretical amount of decinormal soda and adding gum arabic. The brown solution by dialysis, and evaporation in vacuum, gives a black solid, insoluble in water but made soluble by a trace of alkali.

The solutions so obtained can be neutralized, dialyzed and evaporated in vacuum: the black scales so obtained dissolve readily in water and can be employed for hydrogenations in acid media (561). The solutions are not coagulated by boiling with acetic acid, nor by heating with water under pressure.

In another process, called the *germ method*, the same chemist adds to a solution of platinum chloride, PtCl₄, containing gum arabic, a trace of a previously prepared colloidal platinum in solution, and submits the liquid to the action of compressed hydrogen, by which means a colloidal solution of the metal is obtained.²⁵

72. Among colloidal metals, the maximum activity for oxidations belongs to platinum, osmium being only slightly active: ²⁶ for hydrogenations, silver and osmium are much inferior to platinum and particularly to palladium; gold and copper produce no effect.²⁷

OXIDES AS CATALYSTS

73. Water. Water appears frequently as a positive catalyst: quite a large number of reactions are not readily carried out except in the presence of traces of moisture. Oxidations are generally more difficult to realize by means of oxygen rigorously dried.²⁸ A mixture of absolutely dry carbon monoxide and oxygen can not be made to explode. A flame of carbon monoxide is extinguished in perfectly dry air.²⁹ Carbon and even phosphorus refuse to burn in perfectly

²⁵ SKITA, Berichte, 45, 3312 (1912).

²⁶ PAAL, Berichte, 49, 548 (1916).

²⁷ PAAL and GERUM, Berichte, 40, 2209 (1907).

²⁸ DIXON, Proc. Roy. Soc., 37, 56 (1884).

²⁹ Trauba, Berichte, 18, 1890 (1885).

dry oxygen.⁵⁰ Hydrogen and oxygen thoroughly dried do not combine up to 1000°. Ammonia and hydrogen chloride when rigorously freed from moisture do not form any solid ammonium chloride and, conversely, thoroughly dry ammonium chloride can be volatilized without decomposition and the density of its vapor is then normal.⁵¹

A trace of moisture is sufficient to cause the transformation of vitreous arsenic trioxide into its octahedral isomer (porcelain like). Absolutely dry fluorine does not attack glass (Moissan).

This beneficial catalytic effect of water is quite exceptional in organic reactions, but we may mention that in the catalytic oxidation of methyl alcohol vapors by a platinum spiral, the presence of water favors the production of formaldehyde. With absolute methyl alcohol, incandescence is not produced unless the spiral has an initial temperature of at least 400°, while with 20% of water in the alcohol, 175° is sufficient.³⁸

- 74. Sulphur Dioxide. Small amounts of this gas are sufficient to cause the polymerization of acetaldehyde into paraldehyde or metaldehyde (482).
- 75. Anhydrous Metallic Oxides. Manganese dioxide rapidly decomposes hydrogen peroxide, without itself being altered. The same is true of the yellow oxide of lead in alkaline solution. Cuprous oxide is an active catalyst for the decomposition of diazonium salts (606).

The studies that have been made in commercializing the contact process for sulphuric acid, discovered in 1831 (4), have shown that various finely divided metallic oxides may be substituted for the platinum. As early as 1852, Wöhler and Mahla suggested for this purpose, oxides of iron, chromium and copper; and Pétrie, Plattner, and Reich advised the use of pulverized silica.³⁴ In 1854, Tornthwaite proposed manganese oxide.

The application of anhydrous metallic oxides to the catalytic oxidation of volatile organic compounds was proposed anew in 1906 by Sabatier and Mailhe, who mentioned specially the oxides of copper, nickel, cobalt, chromium, manganese and uranium (260). Matignon and Trannoy made the same suggestion (260).

Several anhydrous metallic oxides, particularly alumina, thoria, blue oxide of tungsten, titania and zirconia, etc., are endowed with

- 30 Baker, J. Chem. Soc., 47, 349 (1886).
- ⁸¹ BAKER, *Ibid.*, 65, 611 (1894).
- ³² Winkler, J. pr. Chem. (2), 31, 247 (1885).
- ²⁸ Trillat, Bull. Soc. Chim., (3), 29, 35 (1903).
- ²⁴ Silica gel has been found by Patrick to be an excellent catalyst for the oxidation of nitric oxide by oxygen.—E. E. R.

important catalytic activity towards alcohols, which they can decompose into unsaturated hydrocarbons (701). They can catalyze the synthesis of thiols (743), amines (732), ethers or phenol ethers (786 and 789) and esters (762). These oxides and manganese oxide, employed as catalysts with acids produce symmetrical ketones (837), mixed ketones (847), aldehydes (851) and decompose esters (858). They can also bring about the isomerization or polymerization of unsaturated hydrocarbons (211).

76. The catalytic power of these various oxides is very variable, according to the method of preparation.

Catalysis being a matter of surface, the amorphous oxides prepared from precipitated hydroxides, dehydrated at low temperatures, are much more active than crystallized oxides or those that have been sintered together by calcination at a red heat.

These latter possess, for equal mass, a much smaller surface and are frequently, without doubt, in an advanced stage of molecular condensation. This is particularly true of the oxides of the metals of small atomic weight, aluminum, iron, silicon, chromium, etc. The action of acids has long shown such differences.

77. Amorphous alumina, obtained by dehydrating the hydroxide below 400°, dissolves readily in mineral acids and is an active catalyst for alcohols, while crystallized alumina and amorphous alumina calcined at a bright red, are insoluble in acids and have almost no catalytic power for alcohols.

Analogous differences are observed with the different varieties of silica, though, for the decomposition of hydrogen peroxide, silica calcined at red heat is more active than the dried silica.³⁵

Ferric oxide prepared by dehydrating the precipitated hydroxide below 350°, is a much more powerful catalyst for alcohols than that obtained at a red heat.³⁶

It is the same with regard to hydrogen peroxide of which the former decomposes 50% in 10 seconds, while the latter requires 1550 seconds.

78. Furthermore, the very nature of the catalyst is modified by these changes of constitution of the oxides.

The sesquioxide of chromium, prepared by dehydrating the blue precipitated hydroxide, gives with ethyl alcohol 4.2 cc. gas per minute containing 91% of ethylene, while, after calcination at 500°, the same oxide furnishes only 2.8 cc. gas with 40% ethylene. The oxide pre-

⁸⁵ LEMOINE, Compt. rend., 162, 702 (1916).

²⁶ Sabatier and Mailhe, Ann. Chim. Phys., (8), 20, 313 (1910).

pared by the explosion of ammonium bichromate and, consequently formed with incandescence, gives 1.2 cc. gas, with 38% ethylene.²⁷

The crystallized oxide gives no gas at all at 350°, and 400° must be reached to obtain 2 cc. which is then nearly pure hydrogen. The catalytic function is modified at the same time that it is weakened.³⁸

Analogous variations have been observed with *silica* and *alumina*, both in the intensity and in the direction of the decomposition, ³⁰ and a relation has been noted between the catalytic activity of alumina and its solubility in acids. ⁴⁰

- 79. Thoria, on the contrary, does not present this inconvenience and its activity is not sensibly diminished when it is calcined at a red heat: it appears that such a heavy molecule can not suffer important polymolecular condensations.
- 80. Nickel oxide and especially nickel suboxide, which results from the incomplete reduction of the monoxide, have been regarded by some chemists as the best catalysts for carrying out the hydrogenation of organic compounds in a liquid medium. At least as active as reduced nickel, they have the advantage of being less alterable and consequently of retaining their catalytic activity longer (584). The researches of Sabatier and Espil have indeed established the existence of a suboxide, apparently Ni₄O, which is the first step in the reduction of the monoxide, but they have shown that this suboxide, while it is being formed, is partially reduced to the metal and it is this latter which is the sole factor in the hydrogenations that have been attributed to the oxide.⁴¹

The same reservations should be applied to the *oxide* of *osmium*, which has been proposed as a hydrogenation catalyst (583) and which, doubtless, serves only as a source of finely divided osmium.

MINERAL ACIDS

81. Strong mineral acids frequently act as catalysts in chemical reactions.

Hydrochloric and sulphuric acids, employed in small amounts, bring about the rapid esterification of alcohols by organic acids (749). Hydrochloric acid shows itself also efficacious for the production of acetals from alcohols (782) as well as of similar compounds from

- ⁸⁷ Lemoine, Compt. rend., 162, 702 (1916).
- 88 SABATIER and MAILHE, Ann. Chim. Phys., (8), 20, 339 (1910).
- 89 SENDERENS, Bull. Soc. Chim., (4), 3, 823 (1908).
- 40 IPATIEF, Berichte, 37, 2986 (1904).
- 41 SABATIER and ESPIL, Compt. rend., 158, 668 (1914) and 159, 140 (1914).
- 42 NORMANN and Schick, Arch. Pharm., 252, 208 (1914), C. A., 8, 3129.

glucose with alcohols and thiols.⁴⁸ It also causes catalytic dehydrations in the condensation of ketones (795) and in analogous reactions.

Sulphuric acid behaves similarly in the crotonization of aldehydes and in similar condensations.

These two acids intervene in a similar manner in the acetylation of amines, e. g. of urea. Acetanhydride, without catalyst, gives a yield of only 19.3%, but 73.3% with one molecule of hydrochloric acid, and 61% with one molecule of sulphuric acid.⁴⁴

82. But these acids more frequently accomplish the reverse catalysis in causing hydrolysis, or decomposition by addition of water, and this aptitude they have in common with all strong soluble mineral acids, because it is in consequence of their ionization and should be considered as due to the hydrogen ions which they furnish. Their hydrolytic activity is proportional to their electrolytic dissociation.

We have cases of this decomposition by the addition of water, in the various catalytic effects of acids in the saponification of esters and fats (314), the hydrolysis of amides (331), of anilides, of certain aromatic sulphonic acids, of acetals, in the inversion of sucrose, and, in a more general manner, in the decomposition of polysaccharides such as starch and dextrine.

Hydrochloric acid is a very active polymerizing catalyst for aldehydes, whether it produces a simple aldolization with conservation of the aldehyde function (219), or a cyclization into molecules more or less condensed such as paraldehyde (222).

Sulphuric acid, in small amounts, can likewise cause the change of acetaldehyde into paraldehyde and also the polymerization of ethylene hydrocarbons (210).

Hydriodic acid, in its capacity of a strong acid, can effect hydrolyses, as do the above acids. We may mention also its use in facilitating the preparation of the mixed organo-magnesium halides from chlorides in the Grignard reaction (302).

Nitrous acid catalyzes the transformation of oleïc acid into its isomer, elaïdic acid (186).

INORGANIC BASES

- 83. The alkalies, and alkaline earths, caustic potash and soda, baryta and lime, frequently act as catalysts. In inorganic chemistry they cause the rapid decomposition of hydrogen peroxide and hydrogen persulphides.
 - 48 EMIL FISCHER, Berichte, 26, 2400 (1893) and 27, 615 (1894).
 - 44 Böeseken, Rec. Trav. Chim. Pays-Bas, 29, 330 (1910).
 - 48 CRAFTS, Berichte, 34, 1350 (1901).

In water solution, these strong bases, being highly ionized, hydrolyze esters rapidly. Saponification, when carried out in the presence of excess of alkali, appears, at first sight, to be simply the consequence of the formation of the alkali salt of the acid of the ester, but, in reality, the phenomenon consists of two successive phases, first the hydrolysis which liberates the acid and then the neutralization of the acid to form the salt.

Solutions of *lime* bring about rapid aldolization of aldehydes (221).

A mixture of formaldehyde and acetaldehyde, on long contact with milk of lime, engenders a tetraprimary erythrol along with formic acid.⁴⁶

Solid caustic potash causes the aldolization of acetaldehyde and alcoholic potash, the polymerization of isobutyric aldehyde (224).

Caustic alkalies frequently produce isomerizations (185).

FLUORIDES, CHLORIDES, BROMIDES, AND IODIDES

- 84. Boron Fluoride. Among fluorides, that of boron merits special mention. It produces polymerizations of hydrocarbons: one part of it is sufficient to polymerize 160 parts of oil of turpentine.⁴⁷
- 85. Iodine Chloride. The trichloride ICl_s, the immediate product of the action of excess of chlorine on iodine, is a valuable agent in the direct chlorination of organic compounds by gaseous chlorine (278).
- 86. Barium Chloride. The anhydrous salt readily causes the decomposition of alkyl chlorides into hydrochloric acid and the ethylene hydrocarbons (876).
- 87. Aluminum Chloride. The anhydrous chloride is a catalyst of immense value. It can be employed as an agent in direct chlorination or bromination (284 and 293).

It causes the direct fixation on benzene, of oxygen (263), of sulphur (296), and of sulphur dioxide (297).

It can bring about the decomposition of alkyl chlorides (877) and of thiophenol (297).

In the acetylation of urea it is a much more active catalyst than hydrochloric acid.⁴⁸

Anhydrous aluminum chloride is the basis of a very important

- 46 Tollens and Wigand, Annalen, 265, 317 (1891).
- 47 BERTHELOT, Ann. Chim. Phys., (3), 38, 41 (1853).
- 48 Börseken, Rec. Trav. Chim. Pays-Bas, 29, 330 (1910).

general method for the condensation of organic compounds, which we owe to Friedel and Crafts,⁴⁹ and of which the principal applications and methods of operation will be set forth in Chapter XX.

It acts powerfully on hydrocarbons to cause decompositions as well as molecular condensations (Chapter XXI).

88. Ferric Chloride. Anhydrous ferric chloride can be substituted for aluminum chloride in many of its catalytic reactions. It gives good results as agent of direct chlorination or bromination (285) and even of iodination (295).

It can serve as catalyst in the production of acetals (781 and 783).

It can replace aluminum chloride in the Friedel and Crafts synthesis (899) as well as in analogous condensations (902).

89. Zinc Chloride. This chloride, having a strong affinity for water, is frequently employed as a dehydrating agent. The reactions which it produces are frequently considered as not catalytic, but a closer examination classes them as such, since they are generally produced by small amounts of the salt, smaller than would be required for a chemical reaction.

Thus zinc chloride is a well defined catalyst in the acetylation of glycerine by acetanhydride (761), in the crotonization of aldehydes (795), and in the formation of substituted indols by the decomposition of phenylhydrazones (633). Its rôle is less easy to define and to distinguish from that of an ordinary chemical reagent in quite a number of reactions, such as the condensation of benzaldehyde with nitromethane,⁵⁰ with chloral hydrate,⁵¹ with ethyl orthoformate,⁵² or with phthalic anhydride,⁵³ or of phenols or polyphenols with aromatic amines,⁵⁴ or with fatty acids.⁵⁵

Anhydrous zinc chloride can replace aluminum chloride in the Friedel and Crafts synthesis (899), and can also produce polymerizations (211).

Chlorides of Cobalt, Nickel, Cadmium, and Lead. These decompose alkyl chlorides after the manner of barium chloride (876).

- 90. Stannic Chloride. In certain condensations of organic molecules as of aliphatic aldehydes with phenols, 56 its rôle as a catalyst is difficult to define, as has been said of zinc chloride, or in the
 - 49 FRIEDEL and CRAFTS, Ann. Chim. Phys. (6), 1, 489 (1884).
 - ⁵⁰ Priebs, Annalen, 225, 321 (1884).
 - 51 BOESSNECK, Berichte, 19, 367 (1886).
 - 52 Fischer and Körner, Berichte, 17, 98 (1884).
 - 58 Fischer, Annalen, 206, 86 (1881).
 - ⁵⁴ Calm, Berichte, 16, 2786 (1883).
 - 55 Goldzweig and Kaiser, J. prakt. Chem., (2), 43, 91 (1891).
 - ⁵⁶ FABINYI, Berichte, 11, 283 (1878).

formation of phthaleïnes from phenols and phthalic anhydride,⁵⁷ but it is well established in the addition of acid chlorides to ethylene hydrocarbons (241).

Chlorides of Antimony, Molybdenum, Thallium and Uranium. These can be used as chlorination catalysts (286).

- 91. Cuprous Chloride, Bromide, and Iodide. These cause the decomposition of diazonium salts with the hydracids into the corresponding aromatic halogen compounds, with the elimination of nitrogen (the Sandmeyer reaction) (606). They can bring about the decomposition of phenylhydrazine (611) as well as the production of indols by the decomposition of the phenylhydrazones (633). Cuprous chloride causes the scission of chlorinated hydrocarbons (879). Cuprous iodide has been employed with success in the phenylation of primary aromatic amines (901).
- 92. Mercuric Chloride. This accelerates the isomerization of isobutyl bromide (200) and permits acetaldehyde to be prepared by the hydration of acetylene (309).
- 93. Aluminum Bromide. This is advantageously employed as catalyst in bromination. It causes rapid transformation of propyl bromide into the isomeric isopropyl bromide (199).
- 94. Potassium Iodide. Organic chlorine derivatives usually react with less facility than the corresponding iodides. Their action can be greatly facilitated by the addition of 10% potassium iodide, which apparently permits the progressive transformation of the chloride into the more reactive iodide.⁵⁸
- 95. Potassium Cyanide. It acts as an efficient catalyst of aldolization (220) and even of polymerization in the strict sense (230).

The double cyanide of potassium and copper has been employed as oxidation catalyst (268).

INORGANIC SALTS OF OXYGEN ACIDS

96. A large number of these salts can act as catalysts in organic reactions.

Salts formed from weak acids or from weak bases or ammonia, readily separated by dissociation, usually show effects which could be produced by their constituents separately.

- 97. Alkaline Carbonates. These may be used advantageously in place of caustic potash in reactions of aldolization or of analogous condensations (219 and 236).
 - ⁵⁷ BAEYER, Annalen, 202, 154 (1880).
 - ⁵⁸ Wohl, Berichte, 39, 1951 (1906).

Potassium Bisulphate. This salt can act as free sulphuric acid, either in esterification or in the direct production of acetals, or in condensations effected with elimination of water such as that of dimethyl aniline with benzaldehyde.⁵⁹

Ammonium Sulphate, Nitrate, and Chloride. These can act as the free acids in esterification, or in analogous reactions, such as the production of acetals (783).

98. Barium and Calcium Carbonates. These are equivalent to the free oxides.

Calcium Sulphate. Either as the hydrate, or dehydrated below 400°, it possesses a certain activity for dehydrating alcohols into the ethylene hydrocarbons (718).

99. Aluminum Sulphate and Phosphate. These are dehydration catalysts analogous to free alumina (718).

Silicates. Clay and kaolin, hydrated silicates of aluminum, catalyze the dehydration of alcohols as does alumina (717).

Broken glass, which is a mixed silicate of variable composition, has properties which vary with this composition. In the decomposition of formic acid around 300°, Jena glass yields mainly carbon dioxide and hydrogen, while the ordinary white glass gives water and carbon monoxide, approaching pure silica (828).

Pumice, in spite of its porous structure, is only slightly active as a catalyst and approaches silica in its action.

- 100. Ferrous and Manganous Salts. In the presence of water, these are active oxidation catalysts (264). Thus the presence of various manganous salts aids the oxidation of oxalic acid solutions.
- 101. Magnesium Sulphate. This is an excellent catalyst for the dehydration of glycerine into acrolein (725).
- 102. Mercuric Sulphate. This can cause the hydration of acetylene hydrocarbons into ketones (309), and the oxidation of organic compounds by fuming sulphuric acids (272). Its presence determines the nature of the isomers produced in the direct sulphonation of aromatic molecules (816). It can also determine isomerizations (195).
- 103. Copper Sulphate. In Deacon's process, it is copper sulphate that catalyzes the oxidation of hydrochloric acid by air at 430° with the production of chlorine. It can, although with disadvantage, replace mercuric sulphate in the oxidation of organic compounds by fuming sulphuric acid (272).
 - 59 WALLACH and Wüsten, Berichte, 16, 149 (1883).
 - 60 Jorissen and Reicher, Zeit. physik. Chem., 31, 142 (1900).

VARIOUS COMPOUNDS

104. Ammonia. The presence of ammonia favors the polymerization of cyanamide (233).

Amines. Aliphatic primary and secondary amines are of use as catalysts in the complex reactions in the vulcanization of rubber. Piperidine has been suggested for the purpose.⁶¹ Nitrosodimethylaniline has been recommended in the ratio of 0.3 to 0.5 part to 100 parts caoutchout and 10 parts sulphur at 140°.⁶²

Alkyl Halides and Esters. A small quantity of an alkyl iodide, especially methyl or ethyl iodide, greatly facilitates the preparation of the organo-magnesium compounds in the Grignard reaction, particularly when chlorides are used (302).

Acetaldehyde, heated to 100° with ethyl iodide, condenses to paraldehyde. 63

Ethyl oxalate, by its presence, favors the reduction of ethylene bromide to ethyl bromide by the alloy of sodium and zinc.⁶⁴

Ethyl nitrite, in alcohol solution, causes the transformation of thiourea into ammonium isosulphocyanate.

Ethers. Ethyl ether, as well as amyl ether, and anisol, C_eH_s.O.CH_s, plays an important rôle as catalyst in the formation of the organo-magnesium complexes in the Grignard reaction (300).

105. Aldehydes. Acetaldehyde provokes the hydration of cyanogen to oxamide (311).

106. Organic Acids. Acetic acid can sometimes act, after the fashion of mineral acids, to cause combinations with elimination of water, as in the production of acetals (780). Its catalytic rôle can be disputed in the condensation of benzaldehyde with malonic acid. 65

Isoprene heated with acetic acid is transformed into artificial rubber (215).

Oxalic acid acts like hydrochloric or phosphoric acid in the polymerization of aldehydes.

107. Alkaline Acetates. Sodium acetate is a quite active dehydration catalyst. It produces the crotonization of aldehydes (795) as well as their simple polymerization. It is employed as a catalyst to aid in the esterification of alcohols by acetanhydride.

Quite a large number of organic condensations, which take place

- 61 BAYER & Co., German Patent, 265,221 (1912), C. 1913, (2), 1444.
- 62 PRACHEY, English Patent, 4,263 of 1914.
- 68 LIEBEN, Annalen, Suppl., 1, 114 (1861).
- 64 Michael, Am. Chem. J., 25, 419 (1901).
- 65 CLAISEN and CRISMER, Annalen, 218, 155 (1883).

with elimination of water, have as their basis the use of sodium acetate, but it is usually employed in such large proportions that its catalytic rôle is masked. This is the case in the condensation of phthalid with phthalic anhydride to form diphthalid.66

Likewise potassium acetate permits the condensation of acetic acid with phthalic anhydride to form phthalylacetic acid. 67

It is under the same conditions — that is, employed in large quantity — that sodium acetate causes acetanhydride to act on benzaldehyde to form cinnamic acid in Perkin's synthesis.**

- 108. Nitroso Compounds. The nitroso derivatives of methylaniline, dimethylaniline, and diphenylamine are accelerators in the vulcanization of caoutchouc. The same property belongs to nitrosophenol and nitrosonaphthol but not to the isomeric nitrosoamines.
- 109. Alkyl Cyanides. Methyl and ethyl cyanides are active catalysts in the reaction of sodium with alkyl iodides, or with similar compounds (605).
- 110. Fibrine. It may be recalled that fibrine catalytically decomposes hydrogen peroxide very rapidly.

DURATION OF THE ACTION OF CATALYSTS

111. It would seem, by definition, that the action of catalysts should be prolonged indefinitely, and this perpetuity would be assured to them if they did not suffer any alteration in the course of the reactions which they effect. If any change does take place, as is most frequently the case with solid catalysts acting in gaseous or liquid media, an alteration of the surface, even slight, brings on progressive diminution of activity which may go as far as total suppression.

In hydrogenations carried on by nickel in gaseous systems, using pure and sufficiently volatile substances and thoroughly purified hydrogen, at a carefully regulated temperature, the action can be continued by the same metal a very long time without appreciable weakening. Sabatier and Senderens were able to effect the transformation of benzene into cyclohexane for more than a month with the same nickel, the operation being interrupted every evening and resumed in the morning. The slight oxidation which the metal suffered over night, in the cold tube, caused no inconvenience because the oxide was again reduced by the hydrogen at the beginning of the next run.⁷⁰

- 66 GRAEBE and GUYE, Annalen, 233, 241 (1886).
- 67 GABRIEL and NEUMANN, Berichte, 26, 925 (1893).
- 68 PERKIN, J. Chem. Soc., 31, 388 (1877).
- 69 PRACHEY, J. Soc. Chem. Ind., 36, 424 (1917).
- 70 SABATTER and SENDERENS, Ann. Chim. Phys., (8), 4, 334 (1905).

112. Poisoning of Catalysts. On the contrary, traces of chlorine, bromine, iodine and sulphur in the system are frequently sufficient to suppress the activity of the nickel entirely. It appears to be poisoned. Benzene which is not absolutely free from thiophene can not be hydrogenated. An infinitely small amount of bromine in phenol renders it incapable of being changed into cyclohexanol.⁷¹ Chlorine or bromine derivatives of benzene have never been hydrogenated since the first portions of these compounds alter the metal immediately in an irremediable manner.

113. But the conditions under which this poisoning of the metal take place are quite complex. The presence of free halogens or halogen acids in the hydrogen is much less harmful than the presence of combined halogen in the vapors submitted to hydrogenation. This has been observed by Sabatier and Espil in the hydrogenation of benzene.¹²

In an apparatus in which the hydrogenation of benzene was progressing regularly over nickel at 180°, the benzene was replaced by benzene containing 0.5% iodine. The hydrogenation continued for several hours with an excellent yield. The escaping hydrogen, after the condensation of the cyclohexane, disengaged abundant fumes of hydriodic acid showing that the iodine had been hydrogenated by the catalyst. The operation was interrupted after 130 g. of cyclohexane had been collected and it was found that the nickel had combined with iodine in the first half only of the tube. This half was incapable of carrying on the hydrogenation but the other half was unhurt. The poisoning of the metal by the iodine had taken place only slowly and step by step; the hydriodic acid had had, on its own account, no harmful effect and had not converted into the iodide the metal the surface of which was covered with an unstable hydride which produced the hydrogenation (167). Doubtless the fixation of the hydrogen on the iodine and the benzene in contact with the nickel is much more rapid than the reaction of the nickel with the iodine or with the hydriodic acid. As in the direct hydrogenation of unsaturated hydrocarbons (422), the metal protects itself, by its own action, against the permanent alteration which would render it inactive.

114. Similar results have been obtained, by the same authors, in hydrogenating benzene with hydrogen containing hydrogen chloride, but if traces of brombenzene or chlorbenzene are added to the benzene, the production of cyclohexane ceases almost immediately and the nickel is incapable of regaining its activity.

⁷¹ SABATIER and MAILHE, Compt. rend., 153, 160 (1911).

⁷² SABATHER and ESPIL, Bull. Soc. Chim., (4), 15, 778 (1914).

It is plain that free chlorine or bromine in the hydrogen, unlike iodine, would produce a definite poisoning of the metal since they would offer the possibility of direct substitution in the benzene which iodine does not do.

Sabatier and Espil have likewise been able, for several hours, to transform into cyclohexane benzene containing 10% of carbon disulphide, but traces of thiophene added to the benzene stopped the reaction at once.

115. The use in the oil industry (937 et seq.) of nickel as hydrogenation catalyst suspended in the liquid, has led to the determination of the greater or less toxicity of a number of substances which may be present in small amounts in the oils to be treated.

The soaps formed from the various metals or oxides are, from this point of view, very dissimilar: while those of nickel, thorium, cerium, aluminum, and calcium are absolutely without harmful effect, those of potassium, barium, zinc, cadmium, lead, and uranium are harmful.

The nickel salts of organic monobasic acids, as well as of lactic, oxalic, and succinic acids, are without effect. The same can be said of the free fatty acids such as acetic and stearic, but oxystearic, malic, tartaric and citric acids are true poisons for the nickel catalyst. Toxicity is also shown by calcium hydroxide, potash, boric acid, ammonium molybdate, as well as by sulphur, selenium, red phosphorus, glycerine, lecithine, morphine, strychnine, amygdaline, and cyanides. Tin and aluminum in powder are without action, but iron, lead, and zinc are harmful.⁷⁸

116. With a platinum catalyst, the extreme toxicity of compounds of sulphur,⁷⁴ phosphorus and arsenic and of cyanides, etc., has long been known. The activity of colloidal platinum is diminished or destroyed by a large number of materials. Their toxicity has been measured by means of the velocities of decomposition of hydrogen peroxide and it has been suggested to designate by the term toxicity, the dilution (in liters per gram-molecule) at which the velocity of decomposition in contact with 0.000,01 gram-atom of platinum, is reduced one-half.⁷⁵

Among the violent poisons, hydrocyanic acid stands at the head with toxicity 21,000,000, followed by iodine with 7,000,000, mercuric chloride with 2,500,000, sodium hyposulphite, carbon disulphide, carbon monoxide, and phosphorus. Among the moderate poisons, are placed aniline with toxicity, 30,000, bromine with 23,000, hydrochloric

⁷⁸ SEIICHIDENO, J. Chem. Ind., Tokyo, 21, 898 (1918).

⁷⁴ TURNER, Pogg. Ann., 2, 210 (1824).

⁷⁵ Bradig and Ikana, Zeit. phys. Chem., 37, 1 (1901).

acid with 3,100, oxalic acid, amyl nitrite, arsenious acid, and ammonium chloride. Among the feeble poisons, are found phosphorus acid, 900, sodium nitrite, and hydrofluoric acid, while potassium chlorate, alcohol, ether and pinene have no toxicity and formic acid, hydrazine, and dilute nitric acid are rather favorable. These toxicity coefficients would certainly be very different if measured with platinum black or sponge.⁷⁶

117. Platinum black is very sensitive to the poisons enumerated for colloidal platinum. Traces of potassium cyanide are sufficient to take from the metal all power to hydrogenate the aromatic nucleus, and also to weaken greatly the hydrogenation of ethylene bonds.

Contrary to what has been said about colloidal platinum, the hydrogenation velocity of pinene is diminished if it is dissolved in alcohol or in any substance capable of furnishing alcohol e. g. ether or ethyl acetate. The fatty acids have little action, except formic, which has a marked toxic effect.⁷⁸

118. The Fouling of Catalysts. Other causes of alteration can come in to bring on the decline of catalysts. It happens quite frequently that, along with the principal reaction, there are side reactions which become more important at elevated temperatures and which give rise to highly condensed substances which are only slightly volatile, carbonaceous or tarry. Such substances are slowly deposited on the active surfaces where they hinder the contact with the gas, rendering the useful reaction slow.

In hydrogenations, or decompositions by finely divided metals, the more active the metals, the more rapid are formations of this sort. The most fiery catalysts are the most rapidly enfeebled.

The decline of a catalyst, either from poisoning or fouling, is indicated by the diminishing of the yields in the reaction which it catalyzes.

When a fatigued nickel catalyst is dissolved in dilute hydrochloric acid, fetid hydrocarbons are evolved with the hydrogen and brown carbonaceous or viscous materials are deposited.

119. It can be seen that an analogous enfeeblement will take place when the reaction produces a material which is only slightly volatile at the temperature of the tube and which impregnates the metal more or less rapidly thus opposing its regular activity. This takes place in the hydrogenation of aniline in the presence of nickel at 190°, since

⁷⁶ See comprehensive article by BANCROFT J. Phys. Chem., 21, 767 (1917).

⁷⁷ Madinavettia, Soc. Espan. Phys. Chim., 11, 328 (1913).

¹⁸ BÖESEKEN, VAN DER WEIDE and Mom, Rev. Trav. Chim. Pays-Bas, 35, 260 (1916).

there is produced, in addition to the cyclohexyl amine boiling at 134°, two other amines which are only slightly volatile, dicyclohexyl amine and cyclohexyl aniline, which boiling above 250°, are carried off with difficulty by the hydrogen and remain partly in the liquid form in contact with the metal.

120. It is to avoid analogous effects that it is necessary to watch that the metal is never wetted by an excessive flow of the liquid which is being used or in consequence of an accidental lowering of the temperature of the tube. In the preparation of cyclohexanol or its homologs by the hydrogenation of phenol or the cresols, the reaction is carried on at a temperature only a little above the boiling points of the liquids and it happens sometimes that the nickel is wetted by the liquid. The catalyst immediately becomes nearly inactive, because the surface is, without doubt, altered permanently by contact with the liquid phenol or cresol.

121. Catalytic hydrogenation by finely divided metals is, to a certain extent, comparable to the action of the figured ferments. As with these, there are three periods, an initial period in which the catalyst adapts itself to its function, a period of normal activity and a period of decline, ending in the death of the ferment.

The first period is a variable state and is usually of short duration: it corresponds, without doubt, to the superficial modification which the metal undergoes when the atmosphere of pure hydrogen which surrounded it, is replaced by a mixture of the vapors with hydrogen.

The second period, that of normal functioning, is usually very long and would be indefinite unless something is passed in or is produced which can alter the surface of the metal. Such substances may enter with the hydrogen or with the substance to be hydrogenated or may be produced in the reaction.

- 122. Catalytic oxides, although less sensitive than the metals to chemical alterations of their surfaces, may, nevertheless, suffer from this cause notable diminution of activity even to complete suppression of their function. In many cases they are so fouled that they are weakened or paralyzed.
- 123. Regeneration of altered Catalysts. In so far as the alteration of metallic catalysts is due simply to fouling by deposits of carbon or of tarry substances, calcination in a current of air is suf-

re "Figured ferments" is an obsolete expression for "organized ferments," meaning ferments in which cells can be found with the microscope, as in the yeasts; in contradistinction to such ferments as saliva, etc. The cells were spoken of as "figures," hence "figured ferments."—H. S. Jennings.

ficient to burn off these substances, converting the metal (nickel, iron, copper) into the oxide which a new reduction, carried out at a suitable temperature, will reconvert to the metal. These operations can be carried out in turn in the tube itself in which the catalysis takes place.

This procedure is not suitable for platinum black, which by being heated to redness loses nearly all of its catalytic activity (63).

It does not serve well for the greater part of the metal oxides which are greatly diminished in activity by heating to a high temperature; but it does serve well for *thoria* which has been fouled by long use (708).

124. Metallic catalysts poisoned by vapors of chlorine, bromine, iodine, sulphur, etc., are difficult to revivify except by dissolving in a suitable acid and working over completely.

Calcination does not remove chlorine from slightly chlorinated nickel. The action of hydrogen reduces the chloride to the metallic state below 400°, but the resulting metal is in a peculiar fibrous state and is incapable of reducing benzene to cyclohexane. Even after oxidation and a second reduction it is a poor catalyst.

125. It can be slowly restored to complete activity by employing it for some time in the reduction of nitrobenzene to aniline, work which poisoned nickel is still capable of doing. The aniline which is produced contains increasing amounts of cyclohexyl amine. After some hours of this treatment the power of the metal to produce cyclohexane from benzene is completely restored. On the contrary, poisoning by bromine or iodine seems to resist this treatment.⁸⁰

MIXTURE OF CATALYSTS WITH INERT MATERIALS

126. The desire to increase the active surface of solid catalysts had led to disseminating them over inert porous materials such as pumice, asbestos, infusorial earth, and various metal salts. This practice has appeared specially advantageous for expensive catalysts such as platinum and palladium. Thus in the manufacture of sulphuric acid by the contact process, the catalytic masses are either platinized asbestos, or anhydrous magnesium sulphate impregnated with platinum (about 14 g. metal per kilogram of sulphate).

Nickeled pumice which has been employed by certain chemists in place of nickel powder for hydrogenations, is readily prepared by incorporating the crushed pumice in a thick paste of precipitated

⁸⁰ SABATTER and ESPIL, Bull. Soc. Chim., (4), 15, 779 (1914).

nickel hydroxide, drying in the oven, and finally reducing in the tube that is to be used for the hydrogenations.⁸¹

127. In the case of catalytic metals which have to be carried to a red heat (932), the use of inert siliceous carriers may have serious consequences owing to the formation of silicates which may suppress the activity of the metal. In such cases it is best to use carriers free from silica, such as magnesia, alumina, natural bauxite, lime or carbonate of calcium, etc., either by employing these substances in powders intimately mixed with the oxides, the reduction of which is to furnish the metals, or by previously sticking together these mixtures in little lumps with the aid of non-siliceous materials (Sabatier and Mailhe).

128. In certain cases the use of inert supports for solid catalysts can lead to serious trouble. When the catalyst is to be heated on a furnace, it is disposed in a thin layer in the tube. By a fear entirely unjustified, in view of the great velocity of diffusion of hot gases, some have doubted the sufficiency of the contact between the gas, circulating too freely in the upper part of the tube, and the catalyst. Guided by this thought, the whole height of the tube has been filled with the pumice impregnated with the catalyst. But these conditions are not favorable, since the temperature varies much from bottom to top of the tube. On the contrary, filling the tube entirely with the catalytic mass presents no inconvenience when the tube is heated all around as, for example, by an electric resistance wound around it.

⁸¹ BRUNEL, Ann. Chim. Phys., (8), 6, 205 (1905).

CHAPTER III

THE MECHANISM OF CATALYSIS

129. The extreme diversity of catalytic reactions makes it evident that difficulties will be encountered in giving an explanation that will fit all cases.

Berzelius, who was the first to define catalytic phenomena and to give them this name (4) did not really furnish any explanation for them and found only vague terms with which to characterize the catalytic force which he regarded as the cause of reactions of this kind. "It is evident," said he, "that the catalytic force acts principally by means of the polarity of the atoms which it augments, diminishes or changes. In other words, the catalytic force manifests itself by the excitation of electrical relations which, up to the present, have escaped our investigation." And he adds: "From all that precedes, it follows necessarily that the sources of power (light, heat, electricity) contain the cause of the activity of matter, which, without their influence, would be inert and in a state of unalterable and eternal repose."

To the mind of Berzelius, catalytic forces are then of the order of the sources of power "different effects of one first cause which, under definite circumstances, pass from one modification into another." But their nature remains no less mysterious: the calorific phenomena, sometimes intense, which frequently accompany catalyses, may be the consequences rather than the determining cause.

130. In a great number of catalyses, such as are realized by platinum black and by finely divided metals prepared by reduction of oxides, the porous state seems, at least at first sight, to be the determining cause of the catalytic activity and this thought is the basis of the explanation that has been given of the mechanism of catalysis and which, accepted readily by many chemists, has been usually elaborated in treatises.

- ¹ Benzellus, Traité de Chemie, 2nd Ed., Paris, 1845, I, 112.
- ² Berzelius, loc. cit., 36.

PHYSICAL THEORY OF CATALYSIS

131. Porous materials, whose surfaces are very large as compared with their masses, enjoy the property of absorbing gases with more or less energy. A case of the absorption of gases by solids, that has been much studied, is that of wood charcoal.

When 1.57 g. coconut charcoal, corresponding to 1 cc. of compact carbon, has been heated to redness and cooled under mercury, it absorbs in the cold (at 15° and 760 mm.) quite various volumes of gases, all the way from 2 cc. for argon to 178 cc. for ammonia. These volumes increase nearly proportionally with pressure and decrease greatly when the temperature is raised.

The volume mentioned above for ammonia shows that this gas, if compressed to a volume equal to the total volume of the charcoal would require a pressure of 178 atmospheres, and as this gas is liquefied at 15° under 5.5 atmospheres, it is necessary to assume that the ammonia exists in the pores of the charcoal in the liquid condition, in which it would occupy a volume of about 0.2 cc. (from the known density of liquid ammonia).

The absorption of the gas by the carbon liberates much heat and this amount of heat is even larger than that obtained by the lique-faction of the gas. Thus the amounts of heat per cubic centimeter of gas are: *

	Absorption by	
	carbon	Liquefaction
Sulphur dioxide	0.61 to 0.47 cal.	0.26 cal.
Ammonia	0.45 to 0.33 cal.	0.20 cal.

For ammonia, the heat of absorption is little different from the heat of solution in water and is much larger than the heat of solution in the case of sulphur dioxide.

For hydrogen, the heat of absorption by carbon is six times the heat of liquefaction (Dewar).

132. To explain these singular phenomena, it is assumed that the enormous attraction of the surface of the cavities of the wood charcoal causes the accumulation of the gases in the cavities, at pressures which are not very great for the permanent gases (argon, hydrogen, nitrogen), however, exceeding 35 atmospheres for oxygen, but which are very high for the easily liquefiable gases, generally much greater

^{*} Favre and Silbermann, Ann. Chim. Phys., (3), 37, 465 (1853). Regnault, Ibid., (4), 24, 247 (1871).

⁴ Le Chatelier, Leçons sur le Carbone, Paris, 1908, p. 133.

than the pressures required for liquefaction: this liquefaction would be actually accompanied by a strong compression of the thin layer of liquid produced on the carbon walls. This compression would be responsible for the excess of the heat of absorption over that of liquefaction.

133. An analogous evolution of heat has been observed when any liquid whatever is absorbed by a solid having a very large surface, such as a fine powder, and is called heat of *imbibition*.

Powdered quartz, with grains averaging 0.005 mm. diameter, disengages per gram, when wetted:

With	water	14	calories
With	benzene	4	"

Calculating the surface of the grains, the heat of wetting by water appears to be 0.00105 cal. for 1 sq. cm. of quartz at 7°.

It has been shown likewise, that the wetting by water of 1 g. starch evolves 22 calories, 1 g. wood charcoal, 7 calories, 1 g. alumina, 2 calories.

134. The absorption of gases in the pores of the carbon is equivalent to compressing the gases to a greater or less pressure. Simultaneously there is the liberation of considerable heat by the absorption. It is imagined that the heat and pressure cause reactions to take place. Hydrogen and chlorine may unite in the cold when they meet each other thus in the pores of the carbon, and it is the same way with carbon monoxide and chlorine and with hydrogen sulphide and oxygen.

The oxygen which is absorbed combines little by little with the carbon in the cold to give carbon dioxide. When the gases are pumped out of wood charcoal, which has been exposed to air, scarcely anything is obtained except nitrogen and carbon dioxide.

It would seem then that porous carbon should be a universal catalyst for all gas reactions, lowering the reaction temperatures greatly. However, except for the formation of carbonyl chloride (282), carbon is a mediocre catalyst and of little use, doubtless because gaseous interchanges do not take place rapidly enough in it.

135. Various powdered substances have greater or less power of absorbing gases, but generally, especially for oxides and salts, this power is not great.

Finely divided metals are, in certain cases, able to absorb considerable amounts of gases, but this aptitude is always specific and limited to a small number of gases. In the case of charcoal, the amounts of various gases absorbed are roughly in proportion to their

ease of liquefaction, while with metals the absorption is markedly characterized by a sort of selective affinity.

136. It is one of the most difficultly liquefiable gases, hydrogen, that is absorbed the most readily by metallic powders. The maximum of such absorption is shown by palladium, which, in the form of sponge, can absorb 680 to 850 times its own volume of hydrogen, whatever be the pressure of the gas, provided the pressure be not too low: for all of the hydrogen is given up in a vacuum, even in the cold.⁵

At 20°, platinum black absorbs 110 volumes of hydrogen, whatever the pressure, provided it is more than 200 mm., and here, likewise, the hydrogen is given up in a vacuum.

Reduced cobalt can absorb 153 volumes of hydrogen, finely divided gold, 46, reduced iron or reduced nickel, up to 19, and reduced copper, only 4.7

137. The precious metals have an analogous, though less energetic affinity for oxygen. Thus platinum black absorbs up to 100 volumes of oxygen in the cold and here again this amount is not increased by additional pressure and all of the gas is given up in a vacuum.

Finely divided gold and silver can also take up greater or less amounts of oxygen.8

138. The activity of these finely divided metals, as hydrogenation or oxidation catalysts, would then be due to their power to absorb hydrogen or oxygen along with the vapor which is to be transformed. The compression and local heating thus produced would cause the reaction to take place which without this help would have required a much higher temperature, frequently a temperature so high that the products would not be stable.

The dehydrations of alcohols which are effected by contact with alumina, would result from the condensation of the alcohol vapors in the pores of the alumina, this condensation producing effects comparable to superheating the vapors.

139. The powdered or porous state would be a sufficient condition to produce such effects, since a body containing an infinite number of very small cavities, offers the possibility of realizing simultaneously

- MOND, RAMSAY, and SHIELDS, Phil. Trans. Roy. Soc., 186, 657 (1896).
 Proc. Roy. Soc., 62, 50 and 290 (1897). DEWAR, Chem. News, 76, 274 (1897).
 - 6 Mond, Rambay and Shields, Phil. Trans. Roy. Soc., 186, 675 (1896).
 - Moissan, Traité de Chimie Mineral, I, 13.
- ⁸ NEUMANN, Monatsh., 13, 40 (1892). Mond, Ramsay and Shields, Proc. Roy. Soc., 62, 50 (1897) and Zeit. phys. Chem., 25, 657 (1898). Ramsay and Shields, Phil. Trans. Roy. Soc., 186, 657 (1896). Engler and Wöchler, Zeit. anorg. Chem., 29, 1 (1901).

all possible temperatures and all possible pressures thus causing a great number of reactions by condensation and heating. To this local pressure, there is added also, in the case of metals, the effect of immediate contact with a good conductor and, consequently, electrical influences which might aid. 10

140. A reaction which, without the aid of the catalyst, would take place at an infinitely slow rate, at the temperature of the experiment, would thus receive, on account of the pressure of the catalyst, an immense acceleration and go to completion in a relatively short time.

Catalysis would then be, as Ostwald ¹¹ has defined it, only the acceleration of a chemical phenomenon which otherwise would take place slowly. The presence of the catalyst in the system suppresses the chemical friction which slows up the reaction to the point of stopping it entirely. Its rôle would then be similar to that of oil in clockwork, the movement of which it accelerates, though the forces which produce the movement are not increased.

141. This physical explanation, applicable to all porous catalysts, meets with objections numerous and difficult to get rid of.

Right at the start, the cause which determines the condensation of gases and vapors in the pores of a solid remains mysterious and inexplicable; this physical attraction of solids for gaseous substances presents no visible relation to the properties of the gases. The absorption by wood charcoal is indeed greater for gases which are readily liquefied, but it is just the other way with platinum and various metal powders where the gas that is most absorbed is hydrogen which is very difficult to liquefy.

The same theory is difficult to apply to the case where hydrogen is taken up with the aid of platinum black or nickel held in suspension in a liquid medium (Chapters XI and XII), and even more difficult where the catalyst is colloidal platinum or palladium: for it is difficult to see how high local pressures and temperatures could be developed in such cases.

142. Furthermore, a purely physical conception of the causes of the reaction does not take account of the specificity of catalysts and of the remarkable diversity of the effects produced.

At the same temperature, 300°, the vapors of an alcohol, isobutyl, for example, decompose:

in the presence of copper, into aldehyde and hydrogen, exclusively; in the presence of alumina, into isobutylene and water, exclusively;

- Duclaux, Compt. rend., 152, 1176 (1911).
- 10 VAN'T HOFF, Leçons de Chim. Phys., 1898, 3, 216.
- 11 OSTWALD, Rev. Sci., 1902 (1), 640.

in the presence of *uranium oxide*, both ways, giving at the same time the aldehyde and isobutylene.

Manganous oxide gives the same decomposition as copper, only slowly.

If we assume that the metallic characteristic of conductivity accounts for the fundamental difference between copper and alumina; we can not explain the differences between alumina, and the oxides of manganese and uranium, if the physical condensation in the pores of the catalyst is the sole cause of catalysis.

The action of the catalytic oxide can not be entirely like an elevation of temperature, since the direction of the reaction is intimately connected, not with the physical state of the oxide, but with its chemical nature.

143. The decomposition of formic acid furnishes a no less striking example of the specificity of catalysts (821). Finely divided metals and likewise zinc oxide, decompose this acid into hydrogen and carbon dioxide exclusively, but at the same temperature, titanium oxide gives carbon monoxide and water exclusively, while certain oxides, as thoria, bring about a mixed reaction, more or less complicated by the production of formaldehyde and even of methyl alcohol.

Yet from the physical point of view there does not appear to be any great difference between the oxides of zinc, titanium, and thorium.

144. Furthermore, this explanation of catalysis can not possibly apply to the effects of liquid catalysts in homogeneous systems and it is hard to imagine that there are fundamental differences between the various kinds of catalysis.

CHEMICAL THEORY OF CATALYSIS

145. An entirely general explanation of catalytic phenomena can be based on the idea of the temporary formation of unstable chemical compounds which, serving as intermediate steps in the reaction, determine its direction or increase its velocity.

In order to arrive at a clearer idea of the catalytic mechanism, a special case can be first considered which can be classed as catalytic and which can be designated by the name reciprocal catalysis.

146. Reciprocal Catalysis. Suppose two distinct chemical systems capable of reacting independently, each on its own account: however, each one of them, if left to itself, remains in false equilibrium or, at least, reacts with extreme slowness. But if these two systems

are mixed, they mutually catalyze each other and the two reactions proceed simultaneously very rapidly in correlative proportions.¹²

147. An example is furnished by hydrogen peroxide, opposed by chromic acid, H₂CrO₄. The hydrogen peroxide tends to decompose into water and oxygen, but in the cold, this spontaneous decomposition is very slow and would require more than a year.

The chromic acid solution, acidified with sulphuric acid, is also stable in the cold, but, if heated it decomposes with evolution of oxygen. On heating, we would have:

$$3 \text{ H}_2\text{O}_2 = 3\text{H}_2\text{O} + 3 \text{ O}$$

and $2 \text{ H}_2\text{CrO}_4 + 3 \text{ H}_3\text{SO}_4 = \text{Cr}_4(\text{SO}_4)_4 + 5 \text{ H}_3\text{O} + 3 \text{ O}$.

But if the two solutions are mixed cold, in the exact proportions represented by the formulae above, there is immediate decomposition, simultaneous and complete, of both the hydrogen peroxide and the chromic acid, and this decomposition, manifested by a brisk effervescence of oxygen, takes place in such a manner that the amount of oxygen coming from the hydrogen peroxide is exactly the same as that from the chromic acid.

This proportionality indicates the cause of the reaction, which is apparently the production of an unstable combination of hydrogen peroxide and chromic acid in the proportion 3 H_2O_2 : 2 H_2CrO_4 . As soon as this compound is formed, it decomposes, with liberation of oxygen, leaving water and chromic oxide which dissolves in the sulphuric acid present.

This fugitive combination, the temporary formation of which destroys the false equilibrum of the two systems, really exists: for it appears as an intense blue coloration, when the two liquids are mixed, and can even be isolated. If a dilute solution of hydrogen peroxide is poured into a slight excess of chromic acid: in place of a stormy effervescence a blue solution is obtained. When this is shaken with ether, the dark blue unstable compound passes into the ether. The evaporation of the ether at —20°, leaves a dark blue oil, which, on warming to room temperature, decomposes into chromic oxide, water, and oxygen. We have in succession, 18

$$2H_2CrO_4 + 3 H_2O_2 = 4 H_2O + H_2Cr_2O_{10}$$

 $H_2Cr_2O_{10} = Cr_2O_3 + H_2O + 3 O_2.$

148. Another example of reciprocal catalysis is offered by an acid solution of potassium permanganate opposed by hydrogen peroxide.

¹² SABATIER, Rev. gén. de Chimie pure et app., 17, 185 (1914).

¹⁸ Moissan, Traité de Chimie Min., I, 275 (1904).

The permanganate which is itself an energetic oxidising agent, reduces the hydrogen peroxide immediately, and is itself reduced. Here again there is exact equality between the amounts of oxygen coming from the two reacting substances.

A solution of potassium permanganate, acidified with sulphuric acid, is stable in the cold, but when heated there is a slow reaction:

$$2~{\rm KMnO_4} + 3~{\rm H_2SO_4} - 2~{\rm MnSO_4} + {\rm K_2SO_4} + 3~{\rm H_2O} + 5~{\rm O}.$$

Likewise the hydrogen peroxide alone would give very slowly in the cold:

$$5 \text{ H}_{\bullet}\text{O}_{\bullet} - 5 \text{ H}_{\bullet}\text{O} + 5 \text{ O}.$$

On mixing the two solutions there is immediately a vigorous effervescence, liberating 10 O. The reaction is quantitative and is used practically for the estimation of hydrogen peroxide by titrating with standard potassium permanganate solution. As in the case of chromic acid, this proportionality indicates the formation of an unstable compound, the decomposition of which disengages 5 O₂; but in this case it is difficult to detect. According to Berthelot, the permanganate acts on hydrogen peroxide to substitute hydroxyl groups for the hydrogen atoms, furnishing a sort of hydrogen tetroxide:

which is very unstable and soon decomposes into water and 3 O. When the solutions are mixed at -12° , the permanganate is decolorized without the evolution of oxygen, but the colorless tetroxide, stable at -12° , decomposes on warming, liberating the oxygen. Potassium and caesium tetroxide, which are known, are the alkaline salts of this hydrogen tetroxide.¹⁴

Thus in reciprocal catalysis the simultaneous and correlated reactions of two systems, which apart only tend to react, are determined by the production of an unstable combination which serves as a common intermediate product for the two reactions. This intermediate compound is sometimes visible as in the case of the hydrogen peroxide-chromic acid and sometimes difficult to perceive as in the case of the hydrogen peroxide-permanganate mixture.

149. Induced Catalysis. Suppose a chemical system which tends to react but which remains in false equilibrum or undergoes change infinitely slowly. But if another system which is reacting rapidly in an analogous manner be associated with the first, the first system is drawn into the reaction, without the second seeming to take any

¹⁴ BERTHELOT, Ann. Chim. Phys. (5), 21, 176 (1880) and (7), 22, 433 (1901).

part in the reaction of the first, except, so to speak, setting it an example. This may be called *induced catalysis*, and, as in the case of reciprocal catalysis, there is found to be a proportionality between the two reactions.

Frequent examples of reactions of this sort are found among oxidations by oxygen gas and are called *auto-oxidations*.

150. Auto-oxidations. A large number of substances directly oxidisable by oxygen, or by air, stimulate by their own oxidation that of substances which, without this circumstance, would not be directly oxidisable.

Thus palladium hydride when allowed to oxidise spontaneously in water solution, causes intense oxidations; indigo is decolorized and potassium iodide is oxidised into potassium hydroxide and iodine; ammonia goes into nitric acid, benzene into phenol, and toluene into benzoic acid. Carbon monoxide is oxidised to the dioxide, an oxidation which ozone and hydrogen peroxide are incapable of accomplishing.¹⁵

Ethyl alcohol, exposed to the simultaneous action of sunlight and air, is not appreciably changed, but in the presence of xylene, which is oxidised, the alcohol goes into acetic acid: under the same conditions, amyl alcohol gives valeric acid, and mannite yields mannose.¹⁶

Oxidations of the same nature accompany the spontaneous oxidation of phosphorus in moist air, of turpentine, of aqueous solutions of pyrogallol, of alkaline sulphites, of ferrous hydroxide, of ammoniacal cuprous salts, of benzaldehyde, etc. Such substances are called auto-oxidisers, and experiment has shown that in every case they render active, that is to say, able to oxidise substances otherwise not attacked, exactly the same amount of oxygen as they use up in their own oxidation.¹⁷

151. The cause of the phenomenon appears to be that the auto-oxidiser takes up oxygen to form a sort of peroxide which is then destroyed in the oxidation of the associated substance.

The auto-oxidiser, A, alone would give:

$$A + \underbrace{0 - 0}_{\text{oxygen}} = A \underbrace{0}_{0}.$$

- ¹⁸ Hoppe-Seyler, Berichte, 12, 1551 (1879); 16, 1917 (1883); 20, R795 (1887); Baumann, Ibid., 16, 2146 (1883); 17, 283 (1884). Remsen and Keiser, Am. Chem. Jour., 4, 154 (1883); 5, 424 (1884). Leeds, Chem. News, 48, 25 (1883).
 - 16 CIAMICIAN and SILBER, Berichte, 46, 3894 (1912).
- ¹⁷ Engler and Wild, Berichte, 30, 1669 (1897). Engler, Rev. gén de Chim. pure et app., 6, 288 (1903).

Then in contact with the oxidisable substance, B:

$$A \underbrace{\overset{O}{\overset{}_{\circ}}}_{\circ} + B = \underbrace{A:O}_{\text{stable}} + \underbrace{B:O}_{\text{stable}}.$$

The temporary formation of the combination, A , is the determining cause in the oxidation of the substance B, which would not otherwise have taken place.

In the absence of B, the second reaction would have taken place with the aid of a second molecule of A, thus:

$$A = 2(A:0)$$
.

Whenever this latter reaction is sufficiently slow, the unstable peroxide can be prepared, by the action of oxygen on the auto-oxidiser alone, and may be kept for a time. Thus turpentine shaken with a large volume of air, forms a peroxide which, later on in the absence of air, can decolorize indigo, cause guaiac tincture to turn blue, or liberate iodine from potassium iodide.

The auto-oxidiser, A, is not a catalyst, since it oxidises in proportion to its own mass, and since it does not emerge unchanged from the reaction which it has caused.

152. Oxidation Catalysts. Let us suppose that in the case of the auto-oxidiser, A, opposed by the oxidisable substance, B, that the latter can be oxidised not only at the expense of the unstable

peroxide, $A < \begin{pmatrix} 0 \\ \cdot \\ 0 \end{pmatrix}$, but also by reducing the stable oxide, A:O, we will

then have the succession of reactions:

$$A + O_2 = A O$$

$$A O + B = AO + BO$$

$$AO + B = BO + A$$
regenerated

Thus the auto-oxidiser would be entirely regenerated and could again serve as a carrier of the free oxygen to the oxidisable substance. A limited amount of A could serve to oxidise an unlimited amount of B: A would then be an oxidation catalyst.

153. This condition is realized by cerium salts with glucose in alkaline solution. A cerium salt, dissolved in the presence of potassium carbonate, is a colorless auto-oxidiser. We have:

$$\frac{\text{Ce(OH)}_{s} + \text{O}_{s} + \text{Ce(OH)}_{s} - \text{Ce(OH)}_{s} \cdot \text{O} \cdot \text{O} \cdot \text{Ce(OH)}_{s}}{\text{unstable peroxide}}$$

Water reacts with this compound:

$$\begin{array}{c} \text{Ce(OH)}_3.\text{O.O.Ce(OH)}_3 + \text{H}_2\text{O} = \\ & \underbrace{\text{Ce(OH)}_4}_{\text{outle hydroxide}} + \underbrace{\text{Ce(OH)}_3.\text{O.OH.}}_{\text{blood red}}. \end{array}$$

The blood-red peroxide, when brought into contact with an oxidisable substance, such as potassium arsenite, oxidises it, returning to the state of the stable yellow ceric hydrate. There has been no catalysis. But if glucose is added, the ceric hydrate oxidises the glucose, being itself reduced to cerous hydroxide which can recommence the cycle of reactions. This is catalysis.¹⁸

It is in this manner that small amounts of manganous salts can cause the direct oxidation of unlimited quantities of pyrogallol or hydroquinone.¹⁹

154. Platinum and Related Metals. The activity of platinum and related metals can be explained by a similar mechanism (243). In contact with oxygen, a sort of unstable peroxide is produced on

the surface of the metal, comparable to the A. of the auto-

oxidisers. With an oxidisable substance, B, there is production of BO and AO, but the unstable AO oxidises another molecule of B to form BO and free A. Under these conditions the platinum would serve to render the oxygen *atomic*, and since the platinum is regenerated in the course of the reaction, the cycle can be repeated indefinitely.

The result is that the use of the platinum not only serves to lower the otherwise high temperature required by certain oxidations (e. g. of hydrogen or carbon monoxide) but also to realize other oxidations which can not be accomplished by molecular oxygen at any temperature whatsoever, for example, the liberation of iodine from potassium iodide, which is effected in the cold by aerated ²⁰ platinum black, or

¹⁸ Job, Ann. Chim. Phys., (7), 20, 207 (1900). Compt. rend., 134, 1052 (1902); 136, 45 (1903).

¹⁹ BESTEAND, Bull. Soc. Chim., (3), 17, 578 and 619 (1897). VILLERS, Ibid., (3), 17, 675 (1897).

²⁰ English and Wöhler, Z. anorg. Chem., 29, 1 (1901).

the production of nitric acid from ammonia, by hot platinum sponge.

These fixations and liberations of oxygen take place at the surface of the metal and, for that reason, the catalytic power is proportional to the extent of that surface: it is immeasurably greater for platinum sponge, and especially for the black, than for the metal in foil or wire.

155. General Explanation of Catalysis. The idea of a temporary unstable combination has served in explaining readily the mechanism of reciprocal catalyses (146), of induced catalyses (150), and also of catalyses in the strict sense of the term, such as direct oxidations (152). This notion can be generalized and applied to all sorts of catalyses.

The formation and decomposition of intermediate compounds furnished by the catalysts usually correspond to a diminution of the free energy of the system and this diminution by steps is frequently much easier than the immediate direct diminution, somewhat as the use of a staircase facilitates a descent. Ordinarily these successive step-downs take place quite rapidly, though rapidity is not a necessary condition of catalysis.

These intermediate compounds can be isolated in a sufficiently large number of cases for us to generalize the idea and assume their formation in cases in which we can not prove their existence.

156. Catalyses in which the Intermediate Compounds can be Isolated. Berthelot has pointed out well defined examples in the decomposition of hydrogen peroxide by alkalies and by silver oxide. We will cite some other examples belonging to very different types.

Chlorination of Organic Compounds. In order to facilitate the direct chlorination of a liquid organic compound, iodine is dissolved in it. The chlorine unites with it to form iodine trichloride, ICl_s, which could be isolated if the iodine were alone, but which, finding itself in contact with the organic substance, gives up chlorine to it returning to the lower state of iodine monochloride which the free chlorine transforms into the trichloride, this process being repeated again and again, thus:

$$ICl_s + MH = HCl + MCl + ICl$$

 $ICl + Cl_s = ICl_s$.

It can be proved that the chlorination is proportional to the weight of the iodine trichloride. When the operation is carried on with a continuous current of chlorine, the trichloride is constantly regenerated and we have catalysis (278).

157. The mechanism is doubtless the same for all of the anhydrous

metal chlorides which are used as chlorine carriers in direct chlorination (283). The intermediate products are easy to perceive in the case of the chlorides of antimony, thallium, molybdenum, etc., where several different degrees of chlorination are known of which the highest are formed by direct action of chlorine, and which give up chlorine to the organic substance, returning to the lower stages which again take up chlorine.

It is harder to see in the case of aluminum chloride, for which, by analogy, we must also assume a higher chloride, possibly due to the supplementary valencies of the chlorine atoms.²¹

- 158. Manufacture of Sulphuric Acid. The manufacture of sulphuric acid in the lead chamber process employs, as catalyst, nitric oxide which intimately mixed with the reacting gases (sulphur dioxide, oxygen of the air, and water vapor) serves to render rapid the reaction which would otherwise take place slowly. The production of an intermediate product is doubted by no one although there is not entire agreement as to the true nature of such compound.
- 159. Action of Sulphuric Acid on Alcohol. The mechanism of the action of concentrated sulphuric acid on alcohol is well known and is designated by the name of Williamson's reaction.²² The first reaction is the production of ethyl sulphuric acid:

$$CH_2CH_2OH + H_2SO_4 - H_2O + CH_2CH_2 \cdot O \cdot SO_2H$$
.

The latter, at 140°, reacts with a second molecule of alcohol to form ether, regenerating sulphuric acid:

$$CH_{3}CH_{2}.O.SO_{3}H + CH_{3}CH_{2}OH - H_{2}SO_{4} + (CH_{3}CH_{2})_{2}O.$$

The sulphuric acid can again form ethyl sulphuric acid and so on indefinitely, since the temperature is high enough to cause the elimination of the water along with the ether. Theoretically the action should continue indefinitely: it is a well defined case of catalysis. But a portion of the sulphuric acid is reduced to *sulphur dioxide* gradually diminishing the amount of the acid.

If the mixture is heated higher, towards 160-170°, the ethyl sulphuric acid is rapidly decomposed into sulphuric acid and ethylene:

$$CH_2CH_2 \cdot O \cdot SO_2H - H_2SO_4 + CH_2 : CH_2$$

The regenerated sulphuric acid can repeat the reaction on the alcohol and hence is a catalyst for the formation of ethylene from

²¹ It is possible to consider this a case of the FRIEDEL and CRAFTS reaction, the aluminum chloride combining with the hydrocarbon to form an intermediate complex which reacts readily with Cl-Cl as it does with ClR.—E. E. R.

²² WILLIAMSON, J. Chem. Soc., 4, 106, 229 and 350 (1852).

unlimited amounts of alcohol and can continue this function so long as it is not too much diminished by reduction to sulphur dioxide. This reduction is more serious in this case as the reaction temperature is higher.

160. Hydrogen Peroxide. In the catalytic decomposition of hydrogen peroxide by alkalies and alkaline earths, unstable intermediate compounds are plainly formed and can be isolated.²⁸

The intermediate steps are equally visible in many catalyses brought about in gaseous and liquid media by solid catalysts.

161. Squibb's Method. A fine example is the method of Squibb for the preparation of acetone 24 (837).

If acetic acid vapors are passed over calcium carbonate heated to 400°, calcium acetate is produced with the liberation of carbon dioxide. If the acid is discontinued and the temperature is raised to 500°, the calcium acetate is decomposed, regenerating the carbonate and liberating acetone:

At
$$400^{\circ}$$
 2 CH₃CO₂H + CaCO₃ = CO₂ + H₂O + (CH₅CO₂)₂Ca
At 500° (CH₅CO₂)₂Ca = CaCO₃ + CH₃. CO. CH₃.

If the acetic acid is passed over the calcium carbonate at 500°, it is evident that the first reaction will tend to take place with the formation of calcium acetate, but this would decompose immediately to form acetone: the calcium carbonate would then be a catalyst (839), the reaction being:

$$2 \text{ CH}_{2}\text{CO}_{2}\text{H} = \text{CO}_{2} + \text{H}_{2}\text{O} + \text{CH}_{3} \cdot \text{CO} \cdot \text{CH}_{3}$$

162. Catalytic Oxidation by Copper. If a current of oxygen is passed over copper heated to 250°, a layer of oxide is formed: if the vapors of an organic compound, such as an aliphatic hydrocarbon, are passed over the copper so oxidised, at the same temperature, they are immediately oxidised with the production of water, carbon dioxide, etc., and with regeneration of metallic copper. If the hydrocarbon vapors and the oxygen are sent together over the copper at the same temperature, there is production of the oxide and immediate reduction of the oxide by the hydrocarbon; the copper functions as a catalyst. The total heat of oxidation may be great enough to carry the metal, on the surface of which it is taking place, to incandescence.²⁵ It is easy to see that copper oxide is the intermediate step.

²⁸ Schöne, Annalen, 192, 257 (1878) and 193, 241 (1878). Berthelot, Ann. Chim. Phys., (5), 21, 153 (1880).

²⁴ SQUIBB, J. Amer. Chem. Soc., 17, 187 (1895) and 18, 231 (1896). CONROY, J. Soc. Chem. Ind., 21, 302 (1902). Rev. gén. Sc., 13, 563 (1902).

²⁵ Sabatter and Mailhe, Compt. rend., 142, 1394 (1905).

163. Action of Nickel on Carbon Monoxide. Another example of the same kind is furnished by the destruction of carbon monoxide by nickel at 300°.

Carbon monoxide acting on reduced nickel around 100°, produces nickel carbonyl, Ni(CO)₄. This warmed to about 150° decomposes completely into carbon monoxide and nickel, while from 250° to 300°, it decomposes entirely differently, into nickel, carbon, and carbon dioxide:

$$Ni(CO)_4 = Ni + 2C + 2CO_2$$
.

If carbon monoxide is passed over nickel at 150°, there appears to be no action since the nickel carbonyl that is formed is decomposed immediately, in place, into carbon monoxide and carbon. If the operation is carried on at 300°, there should still be the production of nickel carbonyl but it is at once decomposed into carbon dioxide, carbon, and nickel. The regenerated nickel can carry on the transformation of carbon monoxide into carbon and carbon dioxide indefinitely.

164. Catalyses in which the Intermediate Compounds can not be Isolated. In the cases given above, the intermediate products which serve as stepping-stones for the reaction can be readily observed and even isolated as well defined chemical compounds, but in more numerous cases, these intermediate steps are difficult to perceive and it is only by analogies that we can surmise their nature with more or less uncertainty.

165. Hydrogenation by Finely Divided Metals. The catalytic rôle of finely divided metals, nickel, copper, platinum, etc., in direct hydrogenation is easily explained by the assumption of unstable hydrides on their surfaces.²⁶ Such condensation of hydrogen actually takes place to a certain extent, as we have seen above (136), and particularly with palladium, a really definite combination takes place in the cold. This has only a feeble dissociation pressure and has been assigned the formula, Pd₂H₂, by Dewar.²⁷

²⁶ According to Willstätter and Waldschmidt-Leitz (Berichte, 54, 120 (1921)) oxygen must be present for hydrogenation to take place. They assume that the platinum combines with the oxygen first to form a sort of peroxide which then unites with the hydrogen:

$$Pt + 0_1 \rightarrow Pt \quad \text{and} \quad Pt \quad + H_2 \Rightarrow Pt \quad \cdot \\ 0 \quad 0 \quad H \quad 0$$

This peroxide hydride is the active intermediate compound, passing its hydrogen on to the substance to be hydrogenated and taking up more.—E. E. R.

²⁷ DEWAR, Chem. News, 76, 274 (1897).

The hydrogen thus combined with palladium is able to produce many reactions which free hydrogen can not. It combines directly in the cold and in the dark with chlorine and with iodine as well as with oxygen.²⁸ It reduces chlorates to chlorides, nitrates to nitrites, ferric salts to ferrous, mercuric to mercurous, potassium ferricyanide to ferrocyanide, indigo blue to indigo white, sulphur dioxide to hydrogen sulphide, and arsenic trioxide to arsenic.²⁹ It transforms benzoyl chloride into benzaldehyde and nitrobenzene into aniline.³⁰

166. Hydrogen occluded by platinum produces analogous effects.³¹ Thus when the vapors of nitrobenzene are directed onto *platinum black* previously charged with hydrogen, all the hydrogen which is present is utilized in the production of aniline. If at this moment, more hydrogen is introduced, a new fixation takes place followed by a further reduction of nitrobenzene.

If the hydrogen and nitrobenzene vapors arrive simultaneously, there will be continuous reduction of the latter; the platinum is a hydrogenation catalyst.

The catalysis appears to be a consequence of the occlusion of the hydrogen, that is to say, of the formation of a sort of combination of the hydrogen and the metal and the use of platinum as a catalyst is advantageous since the interchange of gases is rapid with it.

Palladium, although it absorbs much more hydrogen, is usually inferior to platinum, probably because the hydrogen is not given up rapidly enough to the molecules to be hydrogenated.

167. Copper, iron, cobalt, and especially nickel, reduced from their oxides are still more advantageous, although they can retain only small amounts of free hydrogen, probably because the formation and decomposition of the hydrogen addition products are much more rapid.

With nickel, the process goes on as if there were formed, on the surface, an actual unstable hydride capable of liberating hydrogen in the atomic condition and consequently more active than the original molecular hydrogen. The facts lead even to the idea that there are two stages in the fixation of hydrogen such as Ni—H and Ni—H, the latter more active combination being formed by metal reduced from the oxide below 300° and capable of more kinds of work. The former, less active combination, would be produced by nickel reduced above 700°, or made from the chloride and able to hydrogenate ethy-

²⁸ BOWTTOER, Berichte, 6, 1396 (1873).

²⁹ GLADSTONE and TRIBE, Chem. News, 37, 68 (1878).

³⁰ Kolbe and Saytzeff, J. prakt. Chem., (2), 4, 418 (1871).

⁸¹ GLADSTONE and TRIBE, loc. cit. Cooke, Chem. News, 58, 103 (1888).

lenic compounds, nitriles, and nitro bodies but not the aromatic nucleus.

The catalytic hydrogenation of an ethylene hydrocarbon would be represented by:

$$H_2 + Ni_2 - Ni_2H_2$$
.
 $Ni_2H_2 + C_2H_4 - C_2H_8 + Ni_2$.

The regenerated nickel would continue indefinitely to produce this effect so long as the hydrogen and ethylene continued to arrive simultaneously.

168. If finely divided metals with free hydrogen give quickly formed and readily decomposable unstable hydrides, they should also be able to take hydrogen from substances which hold it only feebly and should be *dehydrogenation* catalysts. In general, experiment has verified this prediction (651).

169. Dehydration by Anhydrous Oxides. The dehydration of alcohols by certain oxides, as alumina, thoria, etc., can be interpreted readily by a close analogy to Williamson's reaction.

These oxides can be regarded as the anhydrides of metallic hydroxides capable of exercising the acid function, whether exclusively acid as with silicic or titanic acid, or either acid or basic (hydroxides of aluminum, thorium, chromium, etc.). Thus with alumina, the alcohol vapor would give an unstable aluminate which in contact with alcohol would decompose to give ether, or at a higher temperature would immediately decompose evolving ethylene; the regenerated alumina would be able to carry on this reaction indefinitely:

$$\begin{array}{ll} & Al_2O_3 + 2C_nH_{2n+1}.OH = H_2O + Al_2O_2(OC_nH_{2n+1})_2 \\ \text{Then} & 2C_nH_{2n+1}.OH + Al_2O_2(OC_nH_{2n+1})_2 = 2\underbrace{(C_nH_{2n+1})_2O}_{\text{ether}} + Al_2O_2(OH)_3 \\ \text{and} & Al_2O_2(OH)_2 = H_2O + Al_2O_2 \\ \text{or} & Al_2O_2(OC_nH_{2n+1})_2 = 2C_nH_{2n} + Al_2O_2(OH)_2 \\ \hline & \text{hydrocarbon} \end{array}$$

which would be immediately followed by the dehydration of the alumina.

Such alcoholates can be isolated in various ways, for example, aluminum ethylate, which is decomposed cleanly into ethylene and alumina.³²

In the case of methyl alcohol, only the first sort of reaction is possible, but in most other cases the other takes place exclusively.**

170. It would be the same way with thoria which would furnish with alcohol vapors, a sort of thorium alcoholate which the heat de-

⁸² GLADSTONE and TRIBE, Jour. Chem. Soc., 41, 5 (1882).

²³ SABATIER and MAILHE, Ann. Chim. Phys., (8), 20, 349 (1910).

composes into an ethylene hydrocarbon and thoria, which is capable of reproducing the same effect indefinitely. If this is the case, this sort of *ester* would be capable of reacting chemically with various substances with which it is brought into contact and experiments have bountifully confirmed the predictions made by Sabatier and Mailhe on this point.³⁴

In contact with thoria, alcohol vapors react directly with hydrogen sulphide to give *mercaptans* (743), with ammonia to form *amines* (732), with phenols to produce mixed ethers (789), and with aliphatic acids to yield *esters* (762).

171. Decomposition of Acids. In the decomposition of aliphatic acids by anhydrous oxides it is frequently easy to perceive the intermediate compound which serves as a stepping stone in the reaction; namely, the salt formed by the acid and the oxide. It appears undecomposed at temperatures lower than those used in the catalysis, as is the case with lime and zinc oxide (841). At a higher temperature the salt is immediately decomposed to form the ketone.

This intermediate formation ceases to be apparent when the acid is passed over the oxide at a higher temperature, because the formation of the salt is then balanced by its rapid destruction. For certain oxides, as *thoria* and *titania*, it can not even be perceived since, doubtless, the formation does not take place at a lower temperature than the decomposition, but the analogy is so close that we can not fail to assume similar mechanisms with all of the oxides.

172. In the decomposition of formic acid by metals or oxides (821), the intermediate compounds would be formed either from the hydrogen (passing over the metals), or from the carbon dioxide (fixed by the zinc oxide), or from the formic acid itself giving with the oxide a formate the decomposition of which would vary according to its nature. The molecule of this acid is a structure with little stability, tending to decompose in the two directions, into $CO + H_2O$ or into $CO_2 + H_2$; the affinity of the catalyst giving a transient compound, decides the direction.

173. The Friedel and Crafts Reaction. The catalytic activity of anhydrous aluminum chloride in the Friedel and Crafts reaction (884) can be explained by the production of a temporary combination between the chloride and the organic material. Thus with aromatic hydrocarbons, we would have:

$$C_6R_8H + AlCl_8 = HCl + AlCl_8$$

84 SABATIER and MAILHE, Compt. rend., 150, 823 (1910).

The latter compound would react immediately on the halogen derivative present and we would have:

$$Al \left\langle \begin{array}{c} Cl_2 \\ C_6R_5 \end{array} \right. + R'Cl = \underbrace{AlCl_3 + R'.C_6R_5.}_{\text{regenerated}}$$

The regenerated aluminum chloride would react again with the hydrocarbon and the same reactions would be repeated. It is then a catalyst and a small amount of the salt should effect the transformation of an unlimited amount of the mixture. This is in fact what takes place in some cases where the aluminum chloride can condense a hundred times its own weight of benzene with other molecules.

174. Practically, it is often necessary to employ large amounts of the aluminum chloride, sometimes even several times the weight of the aromatic hydrocarbon. For this reason some chemists have questioned the catalytic rôle of the chloride. It is, however, not to be doubted, as the necessity of sometimes using such large amounts of the catalyst is due either to the tardiness of the reaction in some cases and the desire to hasten it by providing for the formation of a large amount of the required intermediate compound or, in other cases, to the fact that the aluminum chloride forms stable combinations with some of the reactants which withdraw a portion of it from the reaction. The reality of the formation of addition products of the aluminum chloride with the organic compounds has been established by Gustavson who has been able to isolate an addition product with benzene, an orange colored oil, AlCl₂.3C₆H₆, decomposable by water, 35 and in the case of the mixture of benzene and ethyl chloride, AlCl_a. (C₂H₄)₂.3C₆H₆, which heat dissociates into benzene and

Al , which is stable and serves as catalyst for the trans-(C₂H₄Cl)₂ formation of the mixture.³⁶

175. Action of Acids and Bases in Hydrolysis. In the decompositions by addition of water, or hydrolyses, such as the saponification of esters by strong mineral acids (313), or by strong bases (318), the inversion of cane sugar, the decomposition of glucosides (327), or of acetals and, inversely, in the production of esters in presence of small amounts of mineral acids (749), the active factors of the catalysis appear to be the ions resulting from the electrolytic dissociation of the acid or base. The activity of the catalyst is closely

³⁵ Gustavson, Berichte, 11, 2151 (1878).

³⁶ Gustavson, Compt. rend., 136, 1065 (1903); 140, 940 (1905).

⁸⁷ VAN'T HOFF, Lecons Chim. Phys., 1898, III, 140.

connected with the amount of this dissociation and the velocity is proportional to the number of free ions in the solution.

176. In saponifications catalyzed by soluble bases, the active factors are the hydroxyl ions resulting from the electrolytic dissociation of the base and we are justified in believing that the attack on the molecule of ester, ROA, derived from the oxy-acid AOH, is the work of the OH ions derived from the base. Thus with caustic potash we would have:

$$\frac{\text{ROA}}{\text{ester}} + \left\{ \begin{array}{c} \overline{\text{OH}} \\ + \\ K \end{array} \right\} - \frac{\text{ROH}}{\text{alcohol}} + \left\{ \begin{array}{c} \overline{\text{OA}} \\ + \\ K \end{array} \right\}$$

The ionized salt, AOK, is formed in the solution, but as the corresponding organic acid, AOH, is only slightly dissociated into ions, water hydrolyzes the salt to give:

The acid, AOH, is thus liberated and the ions of the original caustic potash are free to recommence their catalytic action.

In the saponification of esters by acids it is the hydrogen ions that cause the effect. Thus with hydrochloric acid, we have:

$$\frac{\text{ROA}}{\text{ester}} + \left\{ \frac{\text{H}}{\text{Cl}} \right\} = \frac{\text{ROH}}{\text{alcohol}} + \left\{ \frac{\text{A}}{\text{Cl}} \right\}$$

But there is immediate reaction with water to give:

$$\begin{Bmatrix} A \\ - \\ Cl \end{Bmatrix} + H_2O - AOH + \begin{Bmatrix} H \\ - \\ Cl \end{Bmatrix}.$$

The regenerated ions of the initial molecule of hydrochloric acid can repeat the reaction and so indefinitely. Esterification is brought about according to the same mechanism but in the inverse direction.

178. The velocity of a hydrolysis of this sort is proportional to the number of ions that are active in producing it. With the strong acids at such dilutions that they may be regarded as completely dissociated, the effect will be independent of the nature of the acid and proportional to the concentration only. This has been verified for hydrochloric, hydrobromic, hydriodic, nitric and chloric acids.** It

38 OSTWALD, J. prakt. Chem., (2), 28, 449 (1883).

is the same way with strong soluble bases, potassium, sodium, barium, and calcium hydroxides in sufficiently dilute solutions.³⁹

179. Catalysis in general appears to be the result of purely chemical phenomena accomplished by the aid of the catalyst which gives, with one of the elements of the primitive system, a temporary unstable combination, the decomposition of which, or the reaction of which, with one of the other reactants, determines the transformation of the system, the catalyst being regenerated in its original condition and able to repeat the reaction indefinitely.

180. Ostwald has criticised the conception of the formation of intermediate compounds because it does not rest on a sufficiently exact knowledge of the reactions and because it would be further necessary to prove that the succession of reactions assumed requires less time than the direct reaction, and adds that no theory is of value in the absence of exact measurements.

To tell the truth, we do not know much more as to the true nature of the absorption of gases and vapors by porous catalysts or even by wood charcoal; this absorption, or occlusion, which is determined by a sort of selective affinity between the gas and the solid is a real solution penetrating to a certain depth in the solid and similar to the temporary combination which we have assumed, the differentiation of chemical and physical phenomena being always uncertain.

The theory of catalysis by means of intermediate compounds still contains many obscurities and has the fault of leaning frequently on the assumption of hypothetical intermediate products which we have not yet been able to isolate, but it is the only hypothesis that is able to explain catalysis in homogeneous solution and has the merit of applying to all cases.

As far as I am concerned, this idea of temporary unstable intermediate compounds has been the beacon light that has guided all my work on catalysis; its light may, perhaps, be dimmed by the glare of lights, as yet unsuspected, which will arise in the better explored field of chemical knowledge. Actually, such as it is, in spite of its imperfections and gaps, the theory appears to us good because it is fertile and permits, in a useful way, to foresee reactions.

²⁰ Reicher, Annalen, 228, 275 (1885). Ostwald, J. prakt. Chem., (2), 35, 112 (1887). Arrhenius, Zeit. phys. Chem., 1, 110 (1887). Bugarszky, Ibid., 8, 418 (1891).

⁴⁰ SABATIER, Berichte, 44, 2001 (1911).

THEORIES OF CONTACT CATALYSIS

By WILDER D. BANCROFT

180a. For purposes of discussion the theories of contact catalysis may be grouped under three headings:—

- 1. Stoichiometric theory.
- 2. Adsorption theory.
- 3. Radiation theory.

The stoichiometric theory is the one most commonly held because it involves nothing new or strange. According to this theory, one or more of the reacting substances forms with the catalytic agent a definite compound which then reacts in such a way as to give the final products. In the catalysis of hydrogen peroxide by mercury, the intermediate formation of mercuric peroxide 41 can be detected by the eye, because there is an intermittent building-up of a film which then breaks down, only to grow again. The formation of graphite is usually preceded by the formation of a carbide. The conversion of acetic acid into acetone 42 by passing the vapor over heated barium carbonate presumably involves the intermediate formation of barium acetate. In the catalytic oxidation of carbon monoxide it is usually believed that there is an alternate oxidation and reduction of the oxides which act as catalytic agents. Hydrogen peroxide is said to oxidize cobaltic oxide to peroxide and to be decomposed catalytically by cobaltic oxide.48 Nickel peroxide reacts quantitatively with hydrogen peroxide; but the resulting oxide is not converted back into peroxide by hydrogen peroxide and consequently does not decompose it catalytically.

180b. While there are undoubtedly many cases of contact catalysis which come under this general head, it does not follow that this is the only type. It seems improbable that it would be so difficult to make carbon tetrachloride if the chlorine, which is absorbed by carbon and thereby made active,⁴⁴ were present as a definite compound of carbon and chlorine. Oxygen absorbed by charcoal will oxidize ethyl alcohol to acetic acid ⁴⁵ and ethylene to carbon dioxide and water, reactions which certainly are not characteristic of any known

⁴¹ Bredig and von Antropoff, Zeit. Elektrochemie, 12, 585 (1906); von Antropoff, Jour. prakt. Chem., (2), 77, 273 (1908).

⁴² Squibb, Jour. Am. Chem. Soc., 17, 187 (1895).

⁴⁸ BAYLEY, Phil. Mag., (5), 7, 126 (1879).

⁴⁴ DAMOISEAU, Compt. rend., 73, 60 (1876).

⁴⁵ CALVERT, Jour. Chem. Soc., 20, 293 (1867).

oxide of carbon. It is very important that we should decide in each particular case whether a definite intermediate compound is formed and, if so, what compound. Only in this way can we escape from the haziness which handicaps so much of the work on catalysis. For instance, it seems obvious to account for the hydrogenating power of pulverulent nickel by postulating the formation of an unstable hydride; but the recent work of Professor Taylor of Princeton shows that no hydride is formed. It is easy to account for the different action of nickel, thoria, and titania on ethyl acetate by postulating the formation of intermediate compounds; but there is no experimental evidence that these hypothetical compounds would break down in the desired way if formed. To this day people are not agreed as to what intermediate compound is formed in the Deacon chlorine process.

180c. The absorption theory does not postulate the intermediate formation of definite chemical compounds. The assumption is that the absorption of the substances to be catalyzed makes them more active chemically. This may occur in different ways. Since the reaction velocity is a function of the concentration, it was natural to ascribe the catalysis of oxyhydrogen gas by platinum to the increased concentration at the surface of the metal. This seems to have been disproved by the recent experiments in which oxyhydrogen gas is reported to be quite stable in presence of an alkaline solution when under a pressure of three thousand atmospheres. This explanation will not suffice to account for the cases in which the same substance decomposes in one way in presence of one catalytic agent and in another way in presence of another. On the other hand, the increase in concentration must have an effect in some cases and it seems probable that this could be found most easily if one studies a reaction which takes place at a measurable rate in the absence of a catalytic agent, say ester formation, and if one takes an extremely non-specific absorbent, such as Patrick's silica gel.

180d. Langmuir ⁴⁶ considers that an adsorbed gas is held chemically by the unsaturated valences at the surface of the solid, thus forming a new type of compound which I have called indefinite compounds because they are not of the ordinary type and because no definite formulas can be written for them. In the case of the adsorption of argon by charcoal, for instance, we should have to write C_xAr_y where x varies with the mass of the charcoal and y with its surface as well as with the pressure and temperature. Chemical reactions may take place either between adjacent atoms

⁴⁶ Jour. Am. Chem. Soc., 37, 1139 (1915); 38, 1145 (1916).

on the surface or when gas molecules strike molecules or atoms on the surface. So far as the catalytic part is concerned this is much the same as the view of Debus.⁴⁷ "If now a piece of platinum is placed in peroxide of hydrogen, the molecules of the latter will place themselves in such a position on the surface of the platinum that one oxygen atom of the peroxide is turned towards the platinum and as near to it as possible. The peroxide is polarized. But this has the effect also of bringing the oxygen atoms of different molecules of peroxide in such close proximity on the surface of the metal that they can combine to form common oxygen, the decomposition of the peroxide into water and oxygen and the development of energy being the consequence. The action of the platinum places the molecules of the peroxide in the position of reaction towards each other."

180e. Langmuir has contemplated the possibility of a reaction between two adsorbed molecules and between one adsorbed and one free molecule. The second case is one in which a more effective collision is produced. This is a perfectly legitimate hypothesis. According to the kinetic theory the reaction velocity is proportional to the number of collisions between possibly reacting molecules: but it does not follow at all that two molecules react every time they collide. If a large number of collisions is necessary on an average before a pair of molecules react, anything which would make these collisions more helpful might increase the reaction velocity enormously. The first question is then whether there is any evidence of ineffective collisions. This matter has been studied by Strutt 48 who comes to the conclusion that a molecule of ozone reacts every time it strikes a molecule of silver oxide; but that a molecule of active nitrogen collides with a molecule of copper oxide five hundred times on an average before they react, while two molecules of ozone at 100° collide on an average 6 x 1011 times before they react. Without insisting on the absolute accuracy of these figures there is evidently plenty of margin for an increase in reaction velocity with ozone at 100° if one could produce more effective collisions. Langmuir 49 finds that, at a pressure of not over 5 bars, and at 2770° K. 15% of all oxygen molecules striking a tungsten filament react with it to form tungstic oxide, WO. This coefficient increases at higher temperatures and at 3300° K about 50% of all the oxygen molecules which strike the filament react with it to form tungstic oxide.

⁴⁷ Jour. Chem. Soc., 53, 327 (1888); Cf. Hüfner, Jour. prakt. chem., (2), 10, 385 (1874).

⁴⁸ Proc. Roy. Soc., 87, A, 302 (1912).

⁴⁹ Jour. Am. Chem. Soc., 35, 105 (1913); 38, 2270 (1916).

180f. It is possible that a catalytic agent may cause one molecule to strike another amidships instead of head-on and may thereby increase the effectiveness of the collisions. It is not impossible that part, at least, of the effect of solvents on reaction velocity may be due to some such thing as this. If we adopt the views of Debus and Langmuir on oriented adsorption, all sorts of things become possible. If ethyl acetate, for instance, attaches itself to one adsorbent by the methyl group, to another by the ethyl group, and to a third by the carboxyl group, it might very well be that bombardment of the captive molecule by free ones might lead to very different reaction products in the three cases. Such a suggestion is of very little value, however, unless it can be made definite. We do not know as vet whether ethyl acetate is actually adsorbed in one way by nickel, in another way by thoria, and in a third way by titania, nor do we know whether the difference in the manner of adsorption, assuming it to occur, is of such a nature as to account for the differences in the reaction products.

180g. It is possible not to make an assumption as to the precise way in which adsorption takes place and merely to consider the surface of the solid as acting like a solvent. If the chemical potential of a possible reaction product is lowered in any way, there is an increased tendency for that reaction product to form. 50 If one treats a substance with a dehydrating agent, the tendency to split off water is increased. If a substance like alcohol can react in two different ways, we should expect a given catalytic agent to accelerate the reaction producing the reaction products which are adsorbed the most strongly by that catalytic agent.⁵¹ This appears to happen in the simpler cases. Ipatief 52 states that the decomposition of alcohol into ethylene and water in presence of heated alumina is due to the taking up of water by the alumina. That alumina takes up water very strongly was shown by Johnson,58 who found that up to a certain point alumina adsorbs water vapor as completely as does phosphorus pentoxide. Sabatier attributed the decomposition of alcohol into acetaldehyde and hydrogen in presence of pulverulent nickel to the tendency to formation of a nickel hydride. Both he and Ipatief assume the formation of definite compounds; but the argument is just as strong in case we postulate that the catalytic agent adsorbs the reaction products strongly instead of combining with them. An

⁵⁰ MILLER, Jour. Phys. Chem., 1, 636 (1897).

⁸¹ BANCROFT, Jour. Phys. Chem., 21, 591 (1917).

⁵² Berichte, 37, 2986 (1904).

⁵³ Jour. Am. Chem. Soc., 34, 911 (1912).

excess of the adsorbed reaction product should cut down the rate of reaction and that is the case. When working at high pressures, the first stage in the dehydration of alcohol in presence of heated alumina is the production of ether. When an equimolecular mixture of ether and water is passed over alumina at 400°, practically no ethylene is formed.⁵⁴ Engelder ⁵⁵ showed that presence of water vapor decreased very markedly the rate of decomposition of ethyl alcohol by alumina. Titania causes alcohol to split both into acetaldehyde and hydrogen and into ethylene and water. Engelder showed that addition of hydrogen to the alcohol vapor increased the relative yield of ethylene and addition of water vapor increased the relative yield of acetaldehyde, though the difference was not as marked as one might have wished. A somewhat similar result appears to have been obtained unconsciously by Berthelot 56 fifty years ago. He heated formic acid at 260° without any specified catalytic agent and found that when only a third of the formic acid is decomposed the reaction appears to be

$$HCO_2H - CO + H_2O_2$$

If all the formic acid is decomposed, the reaction is approximately

$$2 \text{ HCO}_2\text{H} = \text{CO} + \text{H}_2\text{O} + \text{CO}_2 + \text{H}_2.$$

This unexpected result can only be true in case the reaction

$$HCO_2H = CO_2 + H_2$$

predominates during the latter part of the decomposition and this can happen only in case the original decomposition products check the initial reaction and thus permit the second reaction to come to the fore. The experiments by Berthelot should be repeated so as to make sure that they are right and that the suggested explanation is the true one.

180h. While this seems very satisfactory, there are certain points which must not be overlooked. When making ethylene at Edgewood Arsenal during the war, it was found advisable to have a large amount of steam present with the alcohol vapor in order to make temperature regulation easier. This undoubtedly decreased the rate of decomposition of the alcohol; but that difficulty was overcome by working at a higher temperature. I find it very difficult to see how alumina can dehydrate alcohol in presence of a large amount of water vapor if the reason the alumina acts is because of its strong adsorption of water vapor. In spite of the fact that the theory of the selective

⁵⁴ IPATIEF, Berichte, 37, 2996 (1904).

⁵⁵ Jour. Phys. Chem., 21, 676 (1917).

⁵⁶ Ann. Chim. Phys., (4), 18, 42 (1869).

adsorption of the reaction products undoubtedly contains a great amount of truth, it must be admitted that, as now formulated, it is not the final word. It must be modified before it can be considered as satisfactory. If it breaks down temporarily in the simple case of the decomposition of alcohol, it is not surprising that we cannot as yet predict the decompositions of the esters by means of it.

180i. The whole problem of catalysis has been put in a general but vague form by Baly and Krulla and Baly and Rice ⁵⁷ who consider that we have a partial conversion of one or more reacting substances into active forms through opening up fields of force by the rupture of normal valence or of contra-valences. The trouble with this is that it is as yet too vague to be of much value as a working hypothesis, though it makes an admirable starting-point. Methods must be devised for showing in each particular case what particular valences or contra-valences are ruptured as a preliminary step in the reaction.

180j. The radiation theory postulates that the catalytic agent emits radiations which convert one or more of the reacting substances into active modifications. Miss Woker 58 has given a sketch of the earlier speculations as to radiation. The only one which has survived is that of Barendrecht,59 and his calculations have been criticized severely by Henri. 60 Krüger 61 has attempted to account for a number of phenomena in homogeneous solutions by postulating infra-red radiation. This idea has been developed by W. C. McC. Lewis 62 and applied to the change of reaction velocity with the temperature and to contact catalysis. More recently, Perrin es has put forward similar views without making any reference to the work of others. Lewis believes that the catalytic agents emit infra-red rays which activate the reacting substance. This would seem to make it possible for a catalytic agent to act at a distance; but this difficulty can be avoided by assuming that the intensity of the infrared radiation is so low that it is effective only when the distances are molecular. An interesting case comes up in homogeneous solu-

⁵⁷ Jour. Chem. Soc., 101, 1469, 1475 (1912).

⁵⁸ Die Katalyse, p. 60 (1910).

⁵⁰ Zeit. phys. Chem., 49, 456 (1904); 54, 367 (1906); Proc. Kon. Akad. Wet. Amsterdam, 22, 29 (1919).

⁶⁰ Zeit. phys. Chem., 51, 19 (1905).

⁶¹ Zeit. Elektrochemie, 17, 453 (1911).

⁶² Jour. Chem. Soc., 105, 2330 (1914); 107, 233 (1915); 109, 55, 67, 796 (1916); 111, 457, 1036 (1917); 113, 471 (1918); 115, 182 (1919); System of Physical Chemistry, 3, 138 (1919).

⁶³ PERRIN, Ann. Physique, (9), 11, 5 (1919).

tions. Methyl acetate has a strong absorption band between 5 µ and 11 u the hydrogen ion is supposed to emit wave-lengths over the range 1.1-11 u. and hydrogen ion catalyzes methyl acetate solutions. Professor Rideal of the University of Illinois has shown that infra-red radiations corresponding to the absorption band of methyl acetate do accelerate the reaction between methyl acetate and water: but this would happen on any hypothesis. It has not been shown that the catalytic action of the infra-red rays supposed to be emitted by hydrogen ion corresponds quantitatively with the catalytic action of the hydrogen ion. This might be a difficult thing to establish to the satisfaction of the doubters; but there is a test which would probably be accepted as crucial by everybody. Heated nickel decomposes ethyl acetate into propane and carbon dioxide; heated thoria converts it into acetone, ethyl alcohol, ethylene and carbon dioxide; while heated titania changes it into acetic acid and ethylene. 44 If somebody would produce these three sets of reactions separately by means of infra-red radiations with no catalytic agent present, the radiation theory would have a standing which it does not have at present. Since alumina is very permeable to infra-red radiations and ferrous oxide is not,65 the latter should be a very efficient catalytic agent according to the radiation theory. This has not been tested so far as I know. Tyndall 66 states that gum arabic is practically impermeable to infra-red radiations. If this is true, gum arabic should catalyze the hydrolysis of methyl acetate enormously if the radiation theory is sound.

180k. This brief sketch of the theories of contact catalysis shows how unsatisfactory our present knowledge is. This is due to the inaccurate and incomplete way in which the single reactions have been studied. We do not know which cases involve definite intermediate compounds and which do not. When we are agreed that definite intermediate compounds are formed, we do not agree as to their nature. We talk about breaking normal valences or contra-valences; but we do not specify which valences or which contra-valences. When ethyl alcohol is decomposed by pulverulent nickel into acetal-dehyde and hydrogen, does molecular hydrogen split off or do the two hydrogens come off separately? If the latter happens does the first hydrogen come from the hydroxyl group or not? When ethyl alcohol is decomposed by alumina into ethylene and water, does water, hydrogen, or hydroxyl come off first? It can hardly be water because it is possible to stop the reaction at the intermediate stage of ether,

⁶⁴ SABATTER and MAILHE, Compt. rend., 152, 669 (1911).

⁶⁵ ZSIGMONDY, Dingler's Polytech. Jour., (6), 37, 17, 68, 108; 39, 237 (1893).

⁶⁶ Fragments of Science: Radiant Heat and its Relations.

and it is probably not monatomic hydrogen because that is what happens with nickel. If the first stage is a splitting off of hydroxyl, does the other hydrogen come from the adjacent carbon atom giving ethylene direct or does it come from the same carbon atom, forming a substituted methylene, CH₂CH, which then rearranges to ethylene? The decomposition of ether by alumina apparently must lead to 2 CH₂CH + H₂O as one of the intermediate stages. How does nickel decompose ether?

180l. In at least two instances it should be relatively simple to determine the reacting radicals. If we pass a mixture of ethyl acetate and hydrogen over pulverulent nickel, it is probable that some or all of the initial products will be reduced before they have time to react in the normal way. A study of the reaction products will therefore throw light on the probable mechanism of the reaction which occurs in the absence of hydrogen. If we obtained CH₄ and HCO₂C₂H₅, for instance, we should conclude that the original break had been into CH₈ and CO₂C₂H₅. If we found CH₂CO₂H and C₂H₆, we should conclude that these were reduction products of CH₆CO₂ and C₂H₅. If the reaction products were CH₄, C₂H₆, and CO₂ or some reduction product of this last, we should undoubtedly assume that ethyl acetate splits simultaneously into CH₂, CO₂ and C₂H₅.

180m. If ether is passed over pulverulent nickel, the dissociation will probably be to $C_2H_5O + C_2H_5$ or to $C_2H_5O + C_2H_4 + H$. In the first case the final products will be 2 C_2H_4 and H_2O just as with alumina. In the second case they are likely to be $CH_3CHO + C_2H_4 + H_2$, though the ethylene and hydrogen may combine more or less completely to form ethane.

180n. These two illustrations are sufficient to indicate the kind of work that ought to be done and the organic chemists will undoubtedly be able to develop this suggestion in most unexpected ways.

The following cases are worth considering, though it must not be assumed that the reactions run as written for one hundred per cent yield.

With nickel we get the following decompositions of the esters:

$$\begin{array}{c} \mathrm{CH_{2}CO_{2}CH_{2}CH_{2}} = \mathrm{CH_{2}CH_{2}CH_{2}} + \mathrm{CO_{2}} \\ \mathrm{CH_{2}CO_{2}CH_{2}} = \mathrm{CH_{2}CH_{2}} + \mathrm{CO_{2}} \\ \mathrm{HCO_{2}CH_{2}} = \mathrm{CH_{4}(?)} + \mathrm{CO_{2}} \end{array}$$

With thoria the decomposition is quite different:

2
$$CH_{2}CO_{2}CH_{2}CH_{2} - CH_{2}COCH_{3} + CO_{2} + (C_{2}H_{3})_{2}O$$

 $- CH_{2}COCH_{3} + CO_{2} + C_{2}H_{3}OH + C_{2}H_{4}$
2 $CH_{2}CO_{2}CH_{3} - CH_{3}COCH_{3} + CO_{2} + (CH_{3})_{2}O$
2 $HCO_{3}CH_{3} - HCHO + CO_{2} + (CH_{2})_{3}O$

With titania there is a third set of products:

$$CH_sCO_sCH_sCH_s - CH_sCO_sH + C_sH_s$$

 $2CH_sCO_sCH_s - 2CH_sCO_sH + C_sH_s$
 $+CO_sCH_s - HCO_sH + CH_s - CO + CH_sOH.$

The decompositions are regular and characteristic with each catalytic agent and the molecules must break or slip at different points in the different cases. It would help a great deal towards formulating a theory of the behavior of these oxides if we knew exactly what happened in each case. Of course, a study of this sort should include the chlorinated esters. There is some evidence to show that the decomposition may shift from one type to another with increasing substitution of hydrogen by chlorine.

180o. While we have no satisfactorily developed theories of contact catalysis at present, our theoretical knowledge in regard to the poisoning of catalytic agents is in good shape, though it is not supported as yet by adequate experimental evidence. Since the reaction takes place in or at the surface, it follows that any substance, which cuts down the rate at which the reacting substances reach the catalytic surface 67 or which prevents them from reaching it, will decrease the reaction velocity and may destroy the catalytic action entirely. Berliner 68 has shown that traces of fatty vapors from the air or from the grease on the stop-cocks will decrease the adsorption of hydrogen by palladium from nearly nine hundred volumes practically to nothing. Faraday 69 has shown that traces of grease destroy the catalytic action of platinum on oxyhydrogen gas. De Hemptinne 70 has apparently shown that carbon monoxide cuts down the adsorption of hydrogen by palladium, though his method of presenting his results is very obscure. Harbeck and Lunge 71 found that carbon monoxide inhibits practically completely the catalytic action of platinum on a mixture of ethylene and hydrogen. Schönbein 72 pointed out that the hydrides of sulphur, tellurium, selenium, phosphorus, arsenic, and antimony act very energetically in cutting down the catalytic action of platinum on mixtures of air with hydrogen or ether. He considered that the hydride must decompose, giving rise to a solid film. This is not necessary in order to account for the phenomenon; but he seems to have been right in at least one case, for Maxted 78 has found that

```
67 TAYLOR, Trans. Am. Electrochem. Soc., 36 (1919).
```

⁶⁸ Wied. Ann., 35, 903 (1888).

⁶⁹ Experimental Researches on Electricity, 1, 185 (1839).

⁷⁰ Zeit. phys. Chem., 27, 249 (1898).

⁷¹ Zeit. anorg. Chem., 16, 50 (1898).

⁷² Jour. prakt. Chem., 29, 238 (1843).

⁷³ Jour. Chem. Soc., 115, 1050 (1919).

hydrogen sulphide is decomposed by platinum black with evolution of hydrogen, and that the platinum then does not adsorb hydrogen. Paal and Hartmann ⁷⁴ state that the catalytic action of palladium hydrosol and its adsorption of hydrogen are destroyed by metallic mercury or by the oxide of mercury.

180p. Langmuir 75 believes that oxygen prevents dissociation of hydrogen by a heated tungsten filament because it cuts down the adsorption of the hydrogen.

180q. Harned ⁷⁶ has shown that the rate of adsorption ⁷⁷ of chlorpicrin by a charcoal which has been cleaned by washing with chlorpicrin is much greater at first than by a charcoal which has not been so cleaned, although the final equilibrium is apparently about the same in the two cases. This is analogous to the evaporation of water when covered by an oil film. The oil cuts down the rate of evaporation very much but has practically no effect on the partial pressure of water at equilibrium. Taylor points out that normally the time of contact between a gas and the solid catalytic agent is extremely small and consequently anything which decreases the rate of adsorption will cut down the reaction velocity very much.

180r. It is easy to see that the piling up of the reaction products will cut down the reaction velocity, if they prevent the reacting substances from coming in contact with the catalytic agent. Bunsen apparently recognized this as early as 1857 for he is quoted ⁷⁸ as saying that it is only when the products of decomposition are removed and new matter is brought into contact that the reaction continues. This has been observed experimentally in the contact sulphuric acid process. ⁷⁰ The explanation that the decrease in the reaction velocity is due to a decreased adsorption of the reacting substances was first given by Fink, ⁸⁰ who is the real pioneer in this line. Although the reaction between carbon monoxide and oxygen is practically irreversible at ordinary temperature, Henry ⁸¹ recognized that the presence of the reaction product might slow up the rate of reaction and he proved his point by increasing the reaction velocity when he removed the carbon dioxide with caustic potash. Water vapor checks

```
74 Berichte, 51, 711 (1918).
```

¹⁵ Jour. Am. Chem. Soc., 38, 2272 (1916).

⁷⁶ Jour. Am. Chem. Soc., 42, 372 (1920).

⁷⁷ TAYLOR, Trans. Am. Electrochem. Soc., 36 (1919).

⁷⁸ DEACON, Jour. Chem. Soc., 25, 736 (1872).

⁷⁰ BODLÄNDER and KOPPEN, Zeit. Elektrochemie, 9, 566 (1903); BERL, Zeit. anorg. Chem., 44, 267 (1905).

⁸⁰ Bodenstein and Fink, Zeit. phys. Chem., 60, 61 (1907).

⁸¹ Phil. Mag. (3), 9, 324 (1836).

the catalytic dehydration of ether ** and of alcohol ** and hydrogen cuts down the catalytic dehydrogenation of alcohol.

180s. When catalytic poisons are present or are formed during the reaction, the apparent equilibrium may vary with the amount of the catalytic agent.⁸⁴ With only a small amount present, the catalytic agent will be poisoned before the reaction has run very far. In the hydrolysis of ethyl butyrate by enzymes, the reaction apparently runs to different end-points depending on the relative amounts of enzyme.⁸⁵

While our theoretical knowledge in regard to the poisoning of catalytic agents is fairly adequate, we know literally nothing except empirically in regard to the action of the so-called promoters. It has recently been found that the addition of small amounts of a substance which does not in itself have any very marked catalytic action may make the catalyst considerably more active. Such substances were called promoters in the patents of the Badische Anilin and Soda Fabrik, and the term is now in common use. Rideal and Taylor say: "Thus far no theory put forward to account for the acceleration of reaction by minute quantities of promoters added to the main catalyst material is completely satisfactory. A possible mechanism, which. however, has received no experimental test, may be advanced by considering the case of ammonia synthesis from mixtures of nitrogen and hydrogen. Reduced iron is an available contact substance, the activity of which may be regarded as due to the simultaneous formation of the compounds, hydride and nitride, with subsequent rearrangement to give ammonia and unchanged iron. Or, maybe, the activity of the iron is due to simultaneous adsorption of the two gases. The particular mechanism of the catalysis is unimportant for the present considerations. Now such bodies as molybdenum, tungsten, and uranium have been proposed, among others, as promoters of the activity of iron. It is conceivable that these act by adjusting the ratio in which the elementary gases are adsorbed by or temporarily combined with the catalytic material to give a ratio of reactive nitrogen and hydrogen more nearly that required for the synthesis, namely, one of nitrogen to three of hydrogen. From the nature of the materials suggested as promoters, it would seem that they are in the main nitride-forming materials, which on the above assumption of mechanism would lead to the conclusion that the original iron tended

⁸² IPATIEF, Berichte, 37, 2996 (1904).

⁸⁸ Lewis, Jour. Chem. Soc., 115, 182 (1919).

⁸⁴ BANCROFT, Jour. Phys. Chem., 22, 22 (1918).

²⁵ KASTLE and LOEVENHART, Am. Chem. Jour., 24, 491 (1900).

to adsorb or form an intermediate compound with a greater proportion of hydrogen to nitrogen than required by the stoichiometric ratio. The catalytic activity of reduced iron as a hydrogenation agent would tend to confirm this viewpoint.

180t. "In reference to this suggested mechanism it must be emphasized, however, that in such examples of 'promotion,' as require only minute quantities of added promoter the activity is more difficult to understand. With the case of ammonia synthesis, the promoters are added in marked concentrations. It is difficult to realize, however, that 0.5 per cent of ceria or a concentration of one molecule of ceria among 200 molecules of iron oxide, in the example cited above in reference to catalytic hydrogen production, can so far 'redress the balance' of adsorption or combination as to produce the marked increase in activity of which it is capable. It is obvious that in this phase of the problem there lies an exceedingly fascinating field for scientific investigation, with the added advantages that, being practically virgin territory, the harvest to be gained therefrom should be rich and abundant."

180u. Instead of the promoter changing the ratio of adsorption, it might be that the catalytic agent activates only one of the reacting agents or activates one chiefly, and that the promoter activates the other. Thus it might be, in the ammonia synthesis, that iron activates the hydrogen chiefly so that we have hydrogenation of the nitrogen. The molybdenum might tend to activate the nitrogen giving rise to nitridation of hydrogen, or it might increase the activation of the nitrogen. Such a state of things is not impossible theoretically. When a dye reacts with oxygen under the influence of light, the light may make the exygen active, in which case the activated oxygen oxidizes the dye, or the light may make the dye active in which case the activated dye reduces the oxygen. It is easy to decide this question by seeing whether the effective light corresponds to an adsorption band for the dye or for the oxygen.

CHAPTER IV

ISOMERIZATIONS, POLYMERIZATIONS, AND CONDENSATIONS BY ADDITION

§ 1. ISOMERIZATIONS

181. Isomerizations, that is to say, changes of structure effected within a molecule without modifying its composition, are often accomplished by the action of heat alone.

As catalysts have frequently the effect of lowering the temperature of reactions, it can be foreseen that their use will permit, in many cases, of realizing an isomerization at a lower temperature, or causing it to go more rapidly. Experiment has often verified this prediction under very varied conditions.

Strong mineral acids bring about a large number of isomerizations; the concentration of the acid has usually a great influence on the direction of the transformation. The mechanism of the change can usually be interpreted by assuming the addition of water to the original compound under the influence of the acid ions followed by a dehydration, or the reverse.

182. Change of Geometric Isomers. The transformation of fumaric acid into maleic is brought about by a large number of catalysts, for example hydrobromic or hydriodic acids in hot concentrated solution, hot hydrochloric acid, or hot dilute nitric acid.

Bromine acts, in the cold, on maleïc acid to give dibromsuccinic acid but, at the same time, a part of the maleïc acid is changed to fumaric.

Likewise, traces of *iodine* are sufficient to transform maleïc esters into fumaric.⁵

If to a solution of maleïc acid an equivalent amount of sodium thiosulphate be added and then sulphuric acid, sulphur dioxide is evolved without appreciable separation of sulphur and 25% of fumaric acid crystallizes out.⁶

- ¹ Kekulé, Annalen, Supp. Band, 1, 133 (1861).
- ² Kekulé and Strecker, Annalen, 223, 186 (1884).
- ⁸ Kekulé, Annalen, Supp. Band, 2, 93 (1862).
- 4 Petri, Annalen, 195, 49 (1879).

- ⁵ SERAUP, Monatsh., 12, 107 (1891).
- ⁶ TANATAR, J. Russian Phys. Chem. Soc., 43, 1742 (1912), C. A., 6, 1279.

When hydrogen sulphide is passed into solutions of lead, copper, or cadmium maleates, the maleic acid set free is changed to fumaric.

183. Citraconic acid warmed with dilute nitric acid, or with concentrated hydrobromic acid, or with concentrated hydriodic acid, is changed into mesaconic acid.

Warmed above 100° with a concentrated solution of caustic soda, it gives mesaconic acid with a little *itaconic*.¹¹

Itaconic acid dissolved in a mixture of ether and chloroform to which a few drops of a chloroform solution of bromine have been added, and exposed to sunlight, is transformed into mesaconic acid.¹²

Itaconic acid boiled with soda lye changes, almost entirely, into mesaconic.18

184. Small amounts of nitrous acid transform a number of cis ethylenic acids into their trans isomers, oleic into elaïdic, 4 hyprogaeic into gaïdic, 16 erucic, C₈H₁₇CH: CH(CH₂)₁₁CO₂H, into brassidic. 16

185. α -Benzaldoxime in contact with hydrogen chloride or with crystallized pyrosulphuric acid is changed into β -benzaldoxime.¹⁷ The reverse change is brought about by contact with dilute sulphuric acid.

186. Changes of Optical Isomers. Solutions of caustic soda can determine numerous stereo-isomeric changes in the sugar group and the same is true of solutions of lime and baryta and even of pure water mixed with lead and zinc hydroxides. Glucose, mannose and fructose, heated two hours under these conditions yield the same mixture of these three hexoses. In the cold and with concentrated alkalies, the same isomerization takes place in five days. In the same way, galactose gives a mixture of sorbose, tagatose, talose and galtose. Similarly baryta water transforms gulose or idose into sorbose.

- 7 SKRAUP, loc. cit.
- 8 GOTTLIEB, Annalen, 77, 268 (1857).
- ⁹ Firrig, Ibid., 188, 77 and 80 (1877).
- ¹⁰ Kekulé, *Ibid.*, Supl., 2, 94 (1862).
- ¹¹ Delisle, *Ibid.*, 269, 82 (1892). Fittig and Langworthy, *Ibid.*, 304, 152 (1899).
 - ¹² Fittig and Langworthy, *Ibid.*, 304, 152 (1899).
 - 18 Fittig and Köhl, Ibid., 305, 41 (1899).
- ¹⁴ Bouder, Ann. Chim. Phys. (2), 50, 391 (1832). LAURENT, Ibid. (2), 65, 149 (1837).
 - ¹⁵ CALDWELL and GÖSSMANN, Annalen, 99, 307 (1856).
 - 16 HAUSSENECHT, Ibid., 143, 54 (1867).
 - 17 BECKMANN, Berichte, 20, 2766 (1887).
- 18 LOBEY DE BRUYN and VAN EKENSTEIN, Rec. Trav. Chim. Pays-Bas, 14, 203 (1895) and 15, 92 (1896).
 - 19 Van Exenstein and Blanksma, Ibid., 27, 1 (1908).

187. The acids derived from the hexoses are isomerized when they are heated to 135-150° with an organic base that does not yield amides with the acids; quinoline or pyridine are usually employed. The new acid differs from the old only in the arrangement of the groups around the last asymmetric carbon atom. Furthermore, the isomerizations take place in both directions, reaching the same limit. Thus gluconic acid furnishes mannonic with quinoline and reciprocally.²⁰ Likewise with pyridine we pass from arabonic acid (with five carbon atoms) to ribonic,²¹ from lyxonic to xylonic,²² and also from the dibasic acid. talomucic. to mucic.²³

188. The sugars, glucose, laevulose, galactose, arabinose, and xylose, which are not susceptible of a molecular decomposition by the addition of water, present a special phenomenon known as multirotation; the rotatory power observed immediately after solution in water is much greater than that after some time.²⁴

Thus the rotation of glucose starts at 105° and goes down to half of this, 52.5°.25 The explanation is that there are isomeric molecular modifications of these various sugars, analogous to the three varieties that Tanret has been able to isolate for glucose.26

Of the three varieties, the one that is stable in dilute solution, called β , has exactly the rotatory power finally found, 52.5°, another form α has the value 106°. The passage to the stable isomer takes place slowly in the cold, rapidly when hot, but is greatly accelerated by the presence of *mineral acids*.²⁷

- 189. d.Menthone on long contact with sulphuric acid containing 10% of its volume of water passes to l.menthone.25
- 190. Migrations of Double and Triple Bonds. Isopropylethylene, (CH₂)₂CH.CH: CH₂, when heated under pressure at 480–500° in the presence of anhydrous alumina, is transformed into trimethylene, (CH₂)₂C: CH.CH₂.²⁹
- 191. Eugenol, when boiled with amyl alcoholic potash, changes to isoeugenol, the direct oxidation of which furnishes vanilline: ²⁰
 - 20 E. FISCHER, Berichte, 23, 801 (1890).
 - 21 FISCHER and PILOTY, Berichte, 24, 4216 (1891).
 - 22 Fischer and Bromberg, Berichte, 29, 584 (1896).
 - 22 Fischer and Morell, Berichte, 27, 387 (1894).
 - 24 DUBRUNFAUT, Ann. Chim. Phys. (3), 18, 105 (1846).
 - 25 PARCUS and TOLLENS. Annalen. 257, 160 (1890).
 - 26 TANKET, Bull. Soc. Chim. (3), 15, 195 and 349 (1896).
 - 27 ERDMANN, Jahresb., 1855, 672.
 - 28 BECKMANN, Annalen, 250, 334 (1889).
- ²⁰ IPATIEF, J. Russian Phys. Chem. Soc., 38, 63 and 92 (1906), C., 1906, (2), 86 and 87.
 - ²⁰ TIMMANN, Berichte, 24, 2871 (1891).

$$\begin{array}{cccc} CH_{2}\cdot CH: CH_{2} & CH: CH\cdot CH_{4} \\ C_{2}H_{3}-OCH_{3} & \rightarrow & C_{4}H_{4}-OCH_{4} \\ OH & OH & OH & \end{array}$$

192. The acetylene triple bond undergoes analogous transpositions under the influence of sodium or of alkalies.

Ethyl-acetylene, CH₂.CH₂.C: CH, heated with potash to 170°, changes to dimethyl-acetylene, CH₂.C:C.CH₃.²¹

Inversely, disubstituted acetylene hydrocarbons are changed into true acetylenes when they are heated with sodium, a part of the new hydrocarbon combining with the metal. e. g. methyl-ethyl-acetylene, CH₂.C:C.CH₂.CH₃, gives propylacetylene, CH₂. CH₂. CH₂. C: CH.⁸²

The same catalysts cause the transformation of allenic hydrocarbons into acetylene hydrocarbons and inversely. Thus diethylallene, CH., CH., CH.; C: CH., CH., Which under the influence of heat alone isomerizes into methyl-ethyl-butadiene, is changed by contact with metallic sodium into diethyl-allylene, CH₂. CH₃. CH,.C: C.CH,.CH,. **

Inversely isopropyl-acetylene, (CH,), CH, C: CH, heated above 150° with alcoholic potash, changes to dimethyl-allene, (CH₂)₂C:-C: CH., 84

193. Decyclizations. Cyclo-propane is not changed to propylene by heat alone below 600°, but in the presence of platinum sponge, this change takes place in the cold and very rapidly at 100°.85

The vapors of ethyl-cyclo-propane passed at 300-310° over asbestos impregnated with anhydrous alumina, are isomerized into methyl-ethyl-ethylene:

350°, gives divinyl, CH₂: CH . CH : CH₂. 87

- 21 FAWORSKII, J. Russian Phys. Chem. Soc., 19, 414 and 553 (1887); 20, 518 (1888), C., 1887, 153.
- 22 FAWORSKY, J. prakt. Chem., (2), 37, 387 (1888). BÉHAL, Bull. Soc. Chim., 50, 629 (1888).
- 28 MERBHEOVSKI, J. Russian Phys. Chem. Soc., 45, 1969 (1914), C. A., 8,
 - 84 FAWORSKY, J. prakt. Chem. (2), 37, 392 (1888).
 - 35 TANATAR, Zeit. phys. Chem., 41, 735 (1902).
 - ³⁶ ROZANOV, J. Russian Phys. Chem. Soc., 48, 168 (1916), C. A., II, 454.
 - ²⁷ MARESHKOVSKI, Ibid., 45, 2072 (1914), C. A., 9, 799.

194. Cyclizations and Transformations of Ring Compounds. Hydrobenzamide when boiled with potash changes to amarine: 22

$$\begin{array}{c} C_{e}H_{s}\cdot CH: N \\ C_{e}H_{s}\cdot CH: N \end{array} \xrightarrow{CH} \begin{array}{c} C_{e}H_{s}\cdot C\cdot NH \\ C_{e}H_{s}\cdot C\cdot NH \end{array} \xrightarrow{CH} \begin{array}{c} C_{e}H_{s}\cdot C\cdot NH \\ C_{e}H_{s}\cdot C\cdot NH \end{array}$$

195. The acetylenic pinacones when kept on the water bath with a 4% water solution of mercuric sulphate, are rapidly and completely isomerized into ketohydrofurfuranes. Doubtless there is at first addition of water to the triple bond and then dehydration of the glycol thus obtained: 30

$$\begin{array}{c} \operatorname{CH}_{\mathbf{s}} \\ \operatorname{CH}_{\mathbf{s}} \\ \operatorname{CH}_{\mathbf{s}} \end{array} \to \begin{array}{c} \operatorname{CO} - \operatorname{CH}_{\mathbf{s}} \\ \operatorname{CH}_{\mathbf{s}} \\ \operatorname{CH}_{\mathbf{s}} \end{array} \to \begin{array}{c} \operatorname{CO} - \operatorname{CH}_{\mathbf{s}} \\ \operatorname{CH}_{\mathbf{s}} \\ \operatorname{CH}_{\mathbf{s}} \end{array}$$

196. In contact with maleïc acid or with other acids, dimethyl-ketazine isomerizes into trimethyl-pyrazoline: 40

$$\begin{array}{c} \text{CH}_{\text{s}} \\ \text{CH}_{\text{s}} \end{array} \text{C: N \cdot N : C} \begin{array}{c} \text{CH}_{\text{s}} \\ \text{CH}_{\text{2}} \end{array} \rightarrow \begin{array}{c} \text{CH}_{\text{s}} \cdot \text{CH} \cdot \text{CH}_{\text{s}} \\ \text{N - NH} \end{array} \begin{array}{c} \text{CH}_{\text{s}} \\ \text{CH}_{\text{s}} \end{array}$$

197. Cyclo-heptane, heated to 210° with reduced nickel in an atmosphere of hydrogen, is transformed into methyl-cyclo-hexane, and likewise cyclo-octane gives dimethyl-cyclo-hexane.⁴¹

198. Sulphuric acid provokes many isomerizations among the terpenes. Thus pinene, warmed with sulphuric acid diluted with its own volume of water, is changed to a mixture of terpinolene, terpinene, and dipentene.⁴²

l.Pinene dissolved in glacial acetic acid and warmed to 60-70°, isomerizes into l.limonene with evolution of heat, when 5% of phosphoric acid is added. Likewise phellandrene, on contact with sulphuric acid, yields terpinene.

Thujone is isomerized to isothujone when it is warmed for nine hours with sulphuric acid diluted with two volumes of water. 45

In the presence of sulphuric acid, pseudo-ionone passes into the cyclic α - and β -ionones. Thus α -ionone (artificial extract of

- 88 Fownes, Annalen, 54, 364 (1845).
- ³⁹ DUPONT, Compt. rend., 152, 1486 (1911) and 153, 275 (1911).
- 40 CURTIUS and FORRSTERLING, J. prakt. Chem. (2), 51, 394 (1895).
- 41 WILLSTÄTTER and KAMETAKA, Berichte, 41, 1480 (1908).
- ⁴² Armstrong and Thorn, Berichte, 12, 1754 (1879).
- 48 Prins, Chem. Weekbl., 13, 1264 (1916), C. A., 11, 586.
- 44 Wallach, Annalen, 239, 35 (1887).
- 45 WALLACH, Annalen, 286, 101 (1877).

violets) is prepared by heating for 16 hours, 20 parts pseudo-ionone dissolved in 100 parts of water and 100 parts of glycerine with 2.5 parts sulphuric acid. Concentrated sulphuric acid gives mainly β -ionone. Phosphoric acid also may be employed.

199. Migration of Atoms. Migrations of halogen atoms are frequently effected by anhydrous aluminum chloride or bromide. Thus propyl bromide, CH₂. CH₂Br, boiled 5 minutes with 10% of aluminum bromide is completely transformed into isopropyl bromide, CH₃. CHBr. CH₃; while 4% of the salt will effect the change in 24 hours in the cold.⁴⁷ The mechanism is apparently a separation into propylene and hydrobromic acid and a recombination of these to form isopropyl bromide.

Propyl chloride is affected in the same way.48

In the presence of anhydrous aluminum chloride at 110°, acetylene tetrachloride, CHCl₂. CHCl₂, changes partly into the unsymmetrical tetrachlorethane, CCl₂. CH₂Cl. ⁴⁹

By warming with 15 to 20% of aluminum chloride, α -bromnaphthalene, dissolved in 3 or 4 parts of carbon disulphide, is transformed into β -bromnaphthalene.⁵⁰

200. Mercuric chloride and zinc bromide greatly accelerate the isomerization of isobutyl bromide, (CH₃)₂CH. CH₂Br, into tertiary-butyl bromide, (CH₃)₂CBr. ⁵¹

ethylene oxide gives acetaldehyde.⁵² The same transformation is accomplished by anhydrous alumina acting on the vapor of ethylene oxide at 200°.⁵⁸

- 201. Concentrated or dilute *mineral acids* frequently cause the migration of atoms in a straight chain of cyclic hydrocarbon or in a ring containing nitrogen.
 - 46 TIEMANN and KRÜGER, Berichte, 26, 2693 (1893) and 31, 808 (1898).
- 47 Kekulé and Schrotter, Berichte, 12, 2279 (1879). Gustavson, J. Russian Phys. Chem. Soc., 15, 61 (1883).
 - 48 MOUNEYRAT, Bull. Soc. Chim. (3), 21, 616 (1899).
 - 49 MOUNEYRAT, Ibid. (3), 19, 499 (1898).
 - 50 Roux, Ann. Chim. Phys. (6), 12, 344 (1887).
 - ⁵¹ Michael, Schaff, and Voiot, J. Am. Chem. Soc., 38, 653 (1916).
- ⁵² Kaschieski, J. Russian Phys. Chem. Soc., 13, 76 (1881), C., 1881, 278. Krassuski, Ibid., 34, 543 (1902), C., 1902, (2), 1095.
 - 58 IPATIEF and LEONTOWITCH, Beritche, 36, 2016 (1903).

The 1,2 dihydrotetrazines isomerize into the 1,4, when heated with alcoholic hydrochloric acid, 54 thus:

$$C_{e}H_{s} \cdot C \nearrow N \longrightarrow C \cdot C_{e}H_{s} \rightarrow C_{e}H_{s} \cdot C \nearrow N \longrightarrow C \cdot C_{e}H_{s}$$

202. Acetylchloraminobenzene, CH₂.CO.NCl.C₆H₅, is transformed into p.chloracetanilide, CH₃.CO.NH.C₆H₄Cl, under the influence of hydrochloric acid.⁵⁵ The same acid changes hydrasobenzene into benzidene.⁵⁶

203. Acids with a double bond in $\beta\gamma$ position, and hydroxyl in the α , are changed by boiling with dilute hydrochloric acid into γ -keto acids. Thus phenyl- α -hydroxycrotonic acid, C_6H_5 . CH: CH.-CH(OH). COOH, is changed into benzoyl-propionic acid, C_6H_5 .-CO.CH₂.CH₂.COOH. The mechanism of this reaction has been variously explained.⁵⁷

204. The aldoximes, R.CH: NOH, of the aliphatic series are changed to amides, R.CO.NH₂, by warming with sulphuric acid. To explain this change it is sufficient to assume that there is first a dehydration of the oxime to the nitrile which is hydrated by the mineral acid in the usual way to the amide.

205. In contact with sulphuric acid, oximes of cyclic ketones are transformed into internal amides, or iso-oximes. Thus the oxime of cyclohexanone yields the lactam of e-aminocaproic acid:

$$CH_{2} \xrightarrow{CH_{2} \cdot CH_{2}} C: NOH \rightarrow CH_{2} \xrightarrow{CH_{2} \cdot CH_{2} \cdot NH} CH_{2} \cdot CH_{2} \cdot CH_{2} \cdot CH_{3} \cdot CH_{4} \cdot CH_{4} \cdot CH_{5} \cdot CH_{5}$$

The concentrated acid, to which a little water or acetic acid has been added, is suitable for this reaction.⁵⁸

206. Alkaline solutions also can cause the migration of atoms. The potassium salt of diazobenzene heated to 130° with concentrated caustic potash is changed to the potassium salt of phenylnitrosamine. 50°

- 54 STOLLE, J. prakt. Chem. (2), 73, 299 (1906).
- 55 Acres and Johnson, Am. Chem. Jour., 37, 410 (1907).
- 56 ZININ, Annalen, 137, 376 (1865).
- ⁵⁷ Fittig, Annalen, 299, 20 (1898). Thiele and Sulzberger, Ibid., 319, 199 (1901). Erlenmeter, Jr., Ibid., 333, 205 (1904). Bougault, Ann. Chim. Phys. (8), 15, 513 and Compt. rend., 157, 403 (1913).
 - 58 WALLACH, Annalen, 312, 171 (1900).
 - 59 SCHRAUBE and SCHMIDT, Berichte, 27, 522 (1894),

$$C_0H_0 \cdot N : N \cdot OK \rightarrow C_0H_0 \cdot N \setminus_{K}^{NO}$$

207. Thiourea, CS(NH₂)₂, on contact with a solution of ethyl nitrite, isomerizes into ammonium isosulphocyanate, CSN. NH₄.60

208. In certain cases, finely divided metals, copper, nickel, etc., can bring about a migration of atoms, thus causing a change of function. Thus unsaturated alcohols are transformed into aldehydes or ketones in a way that is easy to explain.

Allyl alcohol, CH₂: CH.CH₂OH, passed in the vapor form over reduced copper at 180-300°, is changed almost entirely into propionic aldehyde, CH₃.CH₂CHO, with only slight traces of acroleine, CH₄: CH.CHO. The hydrogen produced by the decomposition of the alcohol by the copper is immediately added to the double bond of the acroleine.⁶¹

Likewise α-unsaturated secondary alcohols, R.CH: CH.-CH(OH). R', mixed with hydrogen over reduced nickel at 195–200°, are isomerized into the ketones, R.CH₂.CH₂.CO. R'. 62

§ 2. POLYMERIZATIONS

209. Frequently several molecules of the same kind, having one or more double bonds, condense to a single molecule, which is called a *polymer* of the original molecule. The presence of a catalyst frequently causes such a change or accelerates its velocity. We will examine from this point of view:

Hydrocarbons.

Aldehydes.

Nitriles and amides.

Hydrocarbons

210. Ethylene Hydrocarbons. Hydrocarbons of the ethylene series, C_nH_{2n}, frequently change into polymers of double, triple, or even quadruple, the original molecule yet retaining the same character as the original.

Sulphuric acid, either concentrated or slightly diluted, frequently causes this polymerization. In fact, its action is complex as, besides polymerization, it can cause the addition of water to form secondary or tertiary alcohols and also the formation of acid or neutral esters

- 60 CLAUS, Annalen, 179, 129 (1875).
- 61 SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 463 (1905).
- 62 Douris, Compt. rend., 157, 55 (1913).

of sulphuric acid. With hydrocarbons of moderate molecular weight, there is principally the formation of alcohols and esters. Thus sulphuric acid diluted with its own volume of water transforms trimethyl-ethylene, $(CH_3)_2C: CH: CH_3$, at 0°, chiefly into dimethylethyl-carbinol, $(CH_3)_2C(OH): CH_3: CH_$

With ethylene hydrocarbons of high molecular weight, there is chiefly production of polymers, particularly dimers. Thus duodecene is changed by sulphuric acid quantitatively into viscous tetracosene which is stable in presence of sulphuric acid. 65

The concentration of the acid determines the nature of the reaction. Thus α -hexene and γ -heptene, with 85% acid yield alkyl sulphuric acids, while they polymerize in contact with the normal acid, H_2SO_4 .

The acid, diluted with 20% of its volume of water, changes isobutene, in the cold, to tributene, boiling at 177°.66

Trimethyl-ethylene in contact with sulphuric acid diluted with half its volume of water, furnishes, at 0°, much diamylene, boiling at 154°.67

211. Zinc chloride can polymerize unsaturated hydrocarbons, e. g. trimethyl-ethylene into diamylene, triamylene, and tetra-amylene. **

Boron trifluoride transforms amylene into diamylene. 69

The use of catalysts under high pressures greatly favors the polymerization of ethylene into unsaturated hydrocarbons at high temperatures. The products obtained with anhydrous alumina, under 70 atmospheres above 250°, are the same as those produced by heat alone in the absence of the catalyst.⁷⁰

Ethylene with anhydrous zinc chloride at 275° and 70 atmospheres, gives a gas containing 36% ethylene, 3% hydrogen, and 61% saturated hydrocarbons and a complex liquid of which 85% is pentane and hexane without any methyl-cyclobutane. The remainder consists of numerous hydrocarbons including unsaturated hydrocarbons boiling above 145° and naphthenes which are particularly abundant around 250°.

Anhydrous aluminum chloride produces little effect with ethylene at 70 atmospheres and 240°, but at 280°, a gas is obtained containing

- 68 Brooks and Humphrey, J. Am. Chem. Soc., 40, 822 (1918).
- 64 WISCHNEGRADSKY, Annalen, 190, 336 (1877).
- 65 Brooks and HUMPHREY, Loc. cit.
- 66 BUTLEROW, Berichte, 6, 561 (1873).
- 67 SCHNEIDER, Annalen, 157, 207 (1871).
- 68 BAUER, Jahresb., 1861, 660.
- 69 LANDOLPH, Berichte, 12, 1578 (1879).
- ⁷⁰ IPATIEF, J. Russian Phys. Chem. Soc., 43, 1420 (1911), C. A., 6, 736.

only saturated hydrocarbons, and no liquid, but, instead, a rather abundant carbonaceous residue.⁷¹

212. Doubly Unsaturated Hydrocarbons. Acetylene is adsorbed more energetically than hydrogen by colloidal palladium and is to a great extent polymerized.⁷²

Allylene is absorbed by concentrated sulphuric acid and is polymerized into mesitylene: 78

3
$$CH_a \cdot C : CH = C_6H_a(CH_a)_a(1.3.5)$$
.

This can be explained by assuming that the acid first causes the hydration of the allylene to acetone (308) and then dehydrates 3 molecules of this according to a well-known reaction.

Similarly crotonylene, or butine (2), shaken with slightly diluted sulphuric acid (1 part water to 3 parts acid), gives hexamethyl-benzene.⁷⁴

Valerylene, C₅H₈, shaken with sulphuric acid changes into polymers, trivalerylene and polyvalerylenes.⁷⁵

213. Divinyl, or butadiene, CH₂: CH. CH: CH₂, as well as its higher homologs, piperylene, CH₃. CH: CH: CH: CH₂, isoprene, CH₂: C(CH₃). CH: CH₂, and dipropylene, CH₂: C(CH₃). -C(CH₃): CH₂, polymerize spontaneously under the influence of heat alone giving rise to various elastic solid hydrocarbons resembling natural rubber and constituting the synthetic rubbers. This polymerization is greatly accelerated by the presence of various catalysts. Thus with 5% metallic sodium or potassium, the reaction which goes on in the cold or with slight warming, is complete and is not hindered by the presence of non-polymerizable hydrocarbons.⁷⁶

214. The polymerization of isoprene by barium peroxide or benzoyl peroxide or potassium sulphide gives rise to the intermediate formation of β -myrcene,

a hydrocarbon boiling at 63° at 20 mm., which, in turn, warmed with sodium or with barium peroxide changes quantitatively into normal

- 71 IPATIEF and RUTALA, Berichte, 46, 1748 (1913).
- 72 PAAL and HOHENEGGER, Berichte, 43, 2684 (1910).
- 78 SCHROHE, Berichte, 8, 17 (1875).
- ⁷⁴ Almedingen, J. Russian Phys. Chem. Soc., 13, 392 (1881), C., 1881, 629.
- ⁷⁵ BOUCHARDAT, Bull. Soc. Chim. (2), 33, 24 (1880). REBOUL, Annalen, 143, 373 (1867).
- ⁷⁶ MATTHEWS and STRANGE, English Pat., 24,790 (1910). HARRIES, Annalen, 383, 157 (1911).

caoutchouc; the direct polymerization furnishes only an abnormal caoutchouc."

215. Glacial acetic acid and especially acetanhydride acting at 150° have been recommended for the polymerization into caout-chouc, the presence of 0.2% of sulphur or of 0.002% of sulphuric acid in the hydrocarbon being favorable to the reaction. 19

Trioxymethylene, at a high temperature in an autoclave, has also been proposed as a catalyst in this reaction.⁸⁰

216. Cyclic Hydrocarbons. Pinene heated twelve hours with formic acid changes to a hydrocarbon of double the molecular weight, $C_{20}H_{32}$. ⁸¹

Pinene is transformed into colophene, C₂₀H₈₂, by contact with concentrated sulphuric acid, boron fluoride, or phosphoric anhydride.**

Pinene heated to 50° with 20% of antimony chloride is changed into tetra-terebenthine, C₄₀H₆₄. ⁸² Aluminum, ferric, and zinc chlorides cause the formation of analogous products. ⁸⁴

217. Indene. Indene polymerizes on contact with sulphuric acid into para-indene, (C₂H₂)_x, which melts at 120°. 35

Aldehydes

- 218. The tendency to polymerize is very general among aldehydes and small traces of various materials are sufficient to cause the polymerization to take place, whether the molecules thus condensed are joined by carbon to carbon or by means of the oxygen atoms.
- 219. Aldolization. The first method of condensation is called aldolization; one of the aldehyde groups is preserved and the other is converted into a secondary alcohol group. The name comes from aldol, the first example to be studied.

Acetaldehyde kept for some time in contact with a small amount of hydrochloric acid or of zinc chloride solution condenses to give aldol, or butanalol (1.3): **

$$CH_3 \cdot CHO + CH_3 \cdot CHO = \underbrace{CH_3 \cdot CH(OH) \cdot CH_2 \cdot CHO}_{aldol}$$

- 77 OSTROMUISLENSKII and KOSHELEV, J. Russian Phys. Chem. Soc., 47, 1928 (1915), C. A., 10, 1947. OSTROMUISLENSKII, Ibid., 48, 1071 (1916), C. A., 11, 1768.
 - 78 CHEM. FABR. AUF. ACTIEN, French Patent, 433,825.
 - ⁷⁹ Badische, French Patent, 434,587.
 - 90 Gross, French Patent, 459,987.
 - 81 LAFONT, Ann. Chim. Phys. (6), 15, 179 (1888).
 - 82 SAINTE-CLAIRE-DEVILLE, Ibid. (2), 75, 66 (1839) and (3), 26, 85 (1849).
 - 88 PRINS, Chem. Weekbl., 13, 1264 (1916), C. A., 11, 586.
 - 84 RIBAN, Ann. Chim. Phys. (5), 6, 42 (1875).
 - 85 KRÄMER and SPILKER, Berichte, 23, 3278 (1890).
 - 86 WURTZ, Compt. rend., 74, 1361 (1872) and 76, 1165 (1873).

The same result is obtained more readily by leaving acetaldehyde for 18 hours in contact with a solution of neutral potassium carbonate or with a fragment of solid caustic potash.²⁷ Also in the presence of zinc turnings at 100°, acetaldehyde gives aldol and likewise crotonic aldehyde by loss of water (795).

220. Likewise benzaldehyde heated with an alcoholic solution of potassium cyanide (10% of the weight of the aldehyde), is rapidly transformed into benzoïne, C₆H₅. CH(OH). CO. C₆H₅. The original aldehyde group is in this case changed to a ketone.

Anisaldehyde, CH₂O. C₆H₄. CHO, with the same reagent, gives anisoine, CH₂. O. C₆H₄. CH(OH). CO. C₆H₄. O. CH₃. ²⁹

On heating an hour and a half, the same catalyst transforms cuminaldehyde into cuminoïne, on and in half an hour, furfural into furfuroïne. on

221. The aldolization of several molecules of aldehyde can be realized successively or simultaneously.

Under the influence of milk of lime, formaldehyde condenses to a hexose, CH₂(OH). CH(OH). CH(OH). CH(OH). CO. CH₂OH, which is racemic laevulose. Analogous condensations giving inactive arabinose and laevulose, are realized in contact with granulated tin, or with a mixture of magnesia, magnesium sulphate, and granulated lead. A similar condensation can be obtained starting with trioxymethylene, (HCOH)₂. 95

222. Second Method. The second method in which aldehydes polymerize suppresses the aldehyde function, producing bodies called paraldehydes and metaldehydes, the vaporization of which tends to reproduce the original aldehyde.

Acetaldehyde in contact with small quantities of sulphur dioxide, anhydrous zinc chloride, hydrogen chloride, or carbonyl chloride soon warms up and is converted into paraldehyde, boiling at 124°. The same result is obtained by warming it with ethyl iodide or by leaving a solution of cyanogen in acetaldehyde to stand for several days.⁹⁶

- 87 MICHAEL and Kopp, Am. Chem. Jour., 5, 190 (1883).
- 83 Wöhler and Liebig, Annalen, 3, 276 (1832). Zinin, Ibid., 34, 186 (1840).
 Breues and Zincke, Ibid., 198, 151 (1879).
 - 89 Rossmi, Ibid., 251, 33 (1869).
 - 90 Bösler, Berichte, 14, 324 (1881).
 - ⁹¹ E. Fischer, Annalen, 211, 218 (1882).
- ⁹² LOEW, Berichte, 22, 475 (1889). E. FISCHER and PASSMORE, Ibid., 22, 359 (1889).
 - 98 Loew, J, prakt. Chem. (2), 34, 51 (1886).
 - 94 Low, Loc. cit.
 - 98 SEVEWETZ and GIBELLO, Compt. rend., 138, 150 (1904).
 - ⁹⁶ Lieben, Annalen, Supl. Band, 1, 114 (1861).

A few bubbles of hydrogen chloride or sulphur dioxide passed into acetaldehyde cooled below 0°, convert it into metaldehyde, a sublimable solid.* By adding one drop of concentrated sulphuric acid to 100 cc. acetaldehyde, paraldehyde is obtained.

223. Likewise by passing a few bubbles of hydrogen chloride into propionic aldehyde cooled below 0°, crystals of metapropanal (melting at 180°) are obtained along with parapropanal, a liquid boiling at 169°. By a current of hydrogen chloride at —20°, metapropanal is formed.*

When a current of dry hydrogen chloride is passed into butanal at —20°, heat is evolved and, on stopping the gas, crystals of metabutanal (melting at 173°) separate out along with oily parabutanal.

Under the same conditions, cenanthal (heptaldehyde) gives paraheptaldehyde (melting at 20°) and metaheptaldehyde (melting at 140°).⁹⁹

224. Isobutyric aldehyde, with a concentrated solution of sodium acetate at 150°, is changed into the dialdehyde boiling at 136°. With a little chlorine, bromine, iodine, hydrochloric acid, phosphorus pentachloride or zinc chloride, meta-isobutanal, melting at 59°, 101 is produced.

With alcoholic potash it gives in succession, tri-isobutanal (b.154°), tetra-isobutanal (b.190°), penta-isobutanal (b.223°), hexaisobutanal (b.250°), and finally oily hepta-isobutanal.¹⁰²

Chloral behaves similarly in contact with various substances, forming solid insoluble metachloral with sulphur dioxide. Trimethyl amine produces the same effect rapidly; 108 fuming sulphuric acid causes the same polymerization, 104 while pyridine gives metachloral in a gelatinous form. 105

225. Third Method. Aromatic aldehydes, e. g. benzaldehyde, when warmed with alkali, undergo a special change, yielding the alcohol and acid at the same time:

$2C_6H_5 \cdot CHO + KOH = C_6H_5 \cdot CO_2K + C_6H_5 \cdot CH_2OH \cdot$

- 97 KEKULÉ and ZINCKE, Ibid., 162, 125 (1872).
- 98 ORNDORF, Amer. Chem. J., 12, 353 (1890).
- 99 Franke and Wozelka, Monatsh., 33, 349 (1912).
- 100 FOSSEK, Ibid., 2, 622 (1881).
- ¹⁰¹ BARBAGLIA, Berichte, 5, 1052 (1872) and 6, 1064 (1873). Demtschenko, Ibid., 6, 1176 (1873).
 - ¹⁰² Perkin, J. Chem. Soc., 43, 91 (1883).
 - 108 MEYER and Dulk, Annalen, 171, 76 (1874).
 - 104 Börseken, Rec. Trav. Chim. Pays-Bas, 29, 104 (1910).
 - 105 Börseken and Schimmer, Ibid., 32, 112 (1913).

Formaldehyde gives the same reaction to some extent with dilute caustic soda. On the contrary, acetaldehyde, with caustic soda or potash, polymerizes into a complex resin.

226. Isobutyric aldehyde with baryta water reacts somewhat like aromatic aldehydes, yielding isobutyl isobutyrate:

 $(CH_2)_2CH \cdot CHO + CHO \cdot CH(CH_2)_2 = (CH_2)_2CH \cdot CO \cdot OCH_2 \cdot CH(CH_2)_2 \cdot CH(C$

When the solution is warmed, the ester is saponified into isobutyl alcohol and isobuturic acid.¹⁰⁷

227. This reaction takes place with all aliphatic aldehydes in which the carbon atom next to the aldehyde group carries no hydrogen.

It is sometimes caused by the presence of ethyl magnesium iodide. With 2,2-dimethyl-propanolal the hydroxypivalic ester of 2,2-dimethyl-propandiol is obtained: 108

228. The same reaction can be brought about with the lower aliphatic aldehydes by the use of aluminum ethylate, Al(OC₂H₈)₈ (299). Thus formaldehyde is condensed into methyl formate, acetaldehyde into ethyl acetate, propionic aldehyde into propyl propionate, even chloral into trichlorethyl trichloracetate.¹⁰⁹

In the case of acetaldehyde this reaction goes quantitatively in 24 hours if 4% of ethyl aluminate be used and the mixture kept below 15°. The ethylate can be used in solution in ethyl acetate, xylene, 110 or solvent naphtha. 111

The reaction is carried out in this way: To 135 parts of acetaldehyde, 6 parts of aluminum ethylate containing 10% aluminum chloride are added little by little and the mixture let stand for ten hours. The yield is 123 parts ethyl acetate.¹¹²

- 106 H. & A. EULER, Berichte, 38, 2556 (1905).
- 107 FRANKE, Monatsh. Chem., 21, 1122 (1900).
- 108 FRANKE and Kohn, Ibid., 25, 865 (1904).
- 109 TISCHENKO, J. Russian Phys. Chem. Soc., 33, 260 (1901).
- 110 KONSORTIUM F. ELEKTBOCH. IND., English pat., 26,825 and 26,826 of 1918. J. S. C. I., 33, 666 (1914). German pat., 277,188 (1913); IMBAY, English pat., 1,288 of 1915, J. S. C. I., 35, 141 (1916).
 - ¹¹¹ German pat., 308,043 (1918), Chem. Centr., 1918 (2), 613.
- ¹¹² Konsortium F. Elektroch. Ind., French patent, 465,965. J. Soc. Chem. Ind., 33, 666 (1914).

Ketones

229. The ketones rarely polymerize but usually condense with the loss of water.

However, aldolization of acetone takes place in the cold with a concentrated solution of caustic soda. Thus:

$$CH_4 \cdot CO \cdot CH_4 + CO \cdot CH_5 = CH_5 \cdot CO \cdot CH_5 \cdot C(OH) \cdot CH_5 \cdot C(OH) \cdot CH_5 \cdot CO \cdot CH_$$

When the product is heated with the same alkali, the reaction is reversed.

Nitriles and Amides

230. Hydrocyanic acid, or formic nitrile, HCN, kept with caustic potash or an alkaline carbonate, desposits crystals of the empirical formula (CNH), which are soluble in ether and appear to be the nitrile of amino-malonic acid, CN.CH(NH₂). CN, along with brown amorphous material.¹¹⁴ The same substance is obtained when a small fragment of solid potassium cyanide is added to a water solution of hydrocyanic acid.¹¹⁵

231. Propionic nitrile, CH₂. CH₂. CN, dissolved in its own weight of anhydrous ether in contact with 20% metallic sodium is converted into dipropionic nitrile, melting at 47°. Under the same conditions, acetonitrile, CH₂. CN, is converted into diacetonitrile, CH₃. C(NH).-CH₂. CN, melting at 52°. 117

232. When the same nitriles, pure and without the ether, are heated with metallic sodium or potassium (1 of metal to 9 of nitrile), they are polymerized into their trimers, acetonitrile into cyanethine, (C₂NH₂)₂. ¹¹⁸

Benzonitrile polymerizes on contact with sulphuric acid into cyaphenine: 119

$$\begin{array}{c|c} & & & & & & & \\ & & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & \\ & & \\ &$$

- 118 KOELICHEN, Z. phys. Chem., 33, 129 (1900).
- 114 WIPPERMANN, Berichte, 7, 767 (1874).
- ¹¹⁵ LESCOEUR and RIGAUT, Compt. rend., 89, 310; Bull. Soc. Chim. (2), 34, 473 (1880).
 - 116 VON MEYER, J. prakt. Chem. (2), 38, 337 (1888).
 - 117 HOLTZWART, Ibid. (2), 39, 230 (1889).
- FRANKLAND and KOLES, Annalen, 65, 269 (1848). BAYER, Berichte, 2,
 (1869) and 4, 176 (1871). VON MEYER, J. prakt. Chem. (2), 27, 153 (1883).
 HOFMANN, Berichte, 1, 198 (1868).

85

233. Cyanamide, either in the cold in contact with concentrated caustic soda or potash, or in a hot solution to which is added a little ammonia, is transformed into dicyanamide. 120

§ 3. DEPOLYMERIZATIONS

234. Depolymerizations are far more rare than polymerizations, since the polymers usually correspond to a much more stable molecular state. In exceptional cases, polymers can be decomposed into the simple molecules by the action of heat and this return is greatly facilitated by the very catalysts that cause the polymerization.

This is the case with paraldehydes and metaldehydes. The catalysts which at low temperature polymerize the aldehydes into their trimers break these up at high temperatures to regenerate the aldehydes. A trace of concentrated sulphuric acid, hydrochloric acid, calcium or zinc chloride or the like is sufficient to change hot paraldehydes into the monomolecular aldehydes. Likewise metaldehydes are transformed into the aldehydes by heating with dilute sulphuric acid. 122

Certain aldols can be decomposed, by warming with a trace of potassium carbonate, regenerating the two molecules of the original aldehyde. But with benzoïne and analogous compounds this decomposition does not take place simply.

235. The transformation of pinene and especially of dipentene, $C_{10}H_{16}$, into isoprene, C_5H_8 , which is realized by the action of an incandescent platinum spiral, ¹²⁸ appears to be due to the eatalytic action of the metal, for this reaction can be caused by passing the vapors of the terpene over pumice impregnated with platinum black in an iron tube at a very low red. ¹²⁴

§ 4. CONDENSATIONS BETWEEN DISSIMILAR MOLECULES

Aldehydes and Ketones

- 236. Aldehydes and ketones can add molecules of other kinds, the reactions being comparable to aldolizations and aided by catalysts of the same nature.
- ¹²⁰ Haag, Annalen, 122, 22 (1862). BAUMANN, Berichte, 6, 1373 (1873). Grube and Krüger, Zeit. phys. Chem., 86, 65 (1914).
 - 121 Franke and Kohn, Monatsh. Chem., 19, 354 (1898).
 - 122 BURSTYN, Ibid., 23, 737 (1902).
- ¹²⁸ Harries and Gottlos, Annalen, 383, 228 (1911). Staudinger and Klever, Berichte, 44, 2212 (1911).
 - 124 SCHORGER and SAYES, J. Ind. Eng. Chem., 7, 924 (1915).

This reaction is general between aldehydes and nitroparaffines and gives nitroalcohols. The presence of an alkali, or better an alkali carbonate, is sufficient to cause the reaction.

By adding a small fragment of potassium bicarbonate to a mixture of equal molecules of nitromethane and acetaldehyde, with an equal volume of water, 1-nitropropanol(2) is obtained: 125

$$CH_2 \cdot CO \cdot H + CH_2NO_2 = CH_2 \cdot CH(OH) \cdot CH_2NO_2 \cdot$$

Likewise nitroethane condenses with formaldehyde in the presence of a little neutral potassium carbonate to give 2-nitropropyl alcohol, CH₂. CH(NO₂). CH₂OH.¹²⁶

Several aldehyde molecules may take part in the reaction. *Nitro-propane* and formaldehyde with a little potassium carbonate give 2-nitro-methanol(2)-butanol(1): 127

$$CH_2 \cdot CH_2 \cdot CH_2NO_2 + 2HCHO = CH_2 \cdot CH_2C(NO_2)(CH_2OH)_2 \cdot$$

A mixture of formaldehyde (commercial formaldehyde solution) and nitro-methane reacts violently on the addition of a fragment of potassium bicarbonate to give 2-nitro-methylol(2) propanediol(1,3), a nitro-triprimary alcohol melting at 158°. 128

$$3HCHO + CH_2NO_2 = C(NO_2)(CH_2OH)_2.$$

237. The mixture of glyceric aldehyde and dihydroxyacetone which is produced by the air-oxidation of glycerine in the presence of finely divided platinum (92), condenses into *i-laevulose* in contact with a water solution of caustic soda: 129

238. Acetone reacts with chloroform in the presence of solid caustic potash to give acetone-chloroform or trichlor-tertiary-butyl alcohol:

$$(CH_3)_2CO + HCCl_3 = (CH_3)_2 \cdot C(OH) \cdot CCl_3$$

To a mixture of 500 parts acetone and 100 parts chloroform, 300 parts of pulverized caustic potash are added very slowly and the mixture left for 36 hours.¹⁸⁰

239. Anhydrous aluminum chloride can sometimes cause the same

- 125 HENRY, Bull. Soc. Chim. (3), 13, 993 (1895).
- 126 HENRY, Ibid., 15, 1223 (1896).
- 127 PAUWELS, Chem. Centbl., 1898 (1), 193.
- 128 HENRY, Compt. rend., 121, 210 (1895).
- ¹²⁹ E. Fischer and Tafel, *Berichte*, 22, 106 (1882). Wohl and Neuberg, *Ibid.*, 33, 3098 (1900).
 - 180 WILLGEBOOT and GENIESER, J. prakt. Chem. (2), 37, 361 (1888).

sort of reactions Thus chloral gives an addition compound with naphthalene, C₁₀H₇. CH(OH). CCl₂. ¹⁸¹

240. Acetylation of Aldehydes. The addition of the anhydrides of monobasic organic acids to aldehydes yields esters of the ethylidene glycols corresponding to the aldehydes. This reaction is catalyzed by the presence of various metal salts, copper sulphate, zinc chloride, ferric chloride, and stannic chloride and even by sulphuric acid. Thus benzaldehyde and acetanhydride give benzylidene acetate quantitatively in the presence of copper sulphate:

$$C_6H_6 \cdot CHO + (CH_2CO)_2O = C_6H_5 \cdot CH(O \cdot COCH_2)_2 \cdot$$

In the presence of stannic chloride, vanilline gives a quantitative yield of the triacetate, the phenol group being simultaneously acetylated.¹⁸²

Hydrocarbons

241. Unsaturated hydrocarbons, ethylenic or acetylenic, may add themselves to hydrocarbons in the presence of aluminum chloride. By passing acetylene into benzene containing aluminum chloride, symmetrical diphenyl-ethane is obtained: 184

$$C_6H_6 + CH : CH + C_6H_6 = C_6H_6 \cdot CH_2 \cdot CH_2 \cdot C_6H_6$$

and also a certain amount of styrene formed by the addition of only one molecule of benzene:

$$C_6H_6 + CH : CH = C_6H_6 \cdot CH : CH_2$$

By passing ethylene into a warm mixture of diphenyl and aluminum chloride, ethyl-diphenyl is obtained:

$$C_0H_5 \cdot C_0H_5 + CH_2 \cdot CH_2 = C_0H_5 \cdot C_0H_4 \cdot CH_2 \cdot CH_3$$

along with some of the diethul derivative.185

242. In an analogous way anhydrous aluminum chloride causes the addition of carbon tetrachloride or of chloroform to ethylenic chlorine derivatives.

Thus trichlorethylene, CCl₂: CHCl, gives, with carbon tetra-

- 181 Frankforter and Daniels, J. Amer. Chem. Soc., 37, 2560 (1915).
- 182 KNOEVENAGEL, Annalen, 402, 111 (1913).
- 188 This may be considered as a case of the Friedel and Crafts reaction. A trace of water is always present and reacts with the aluminum chloride to give hydrochloric acid which adds to the hydrocarbon to form an alkyl chloride which then reacts in the usual way liberating hydrochloric acid which repeats the reaction.—E. E. R.
 - 184 VARET and VIENNE, Bull. Soc. Chim. (2), 47, 919 (1887).
- ¹⁸⁵ ADAM, Bull. Soc. Chim. (2), 47, 689 (1887) and Ann. Chim. Phys. (6), 15, 252 (1888).

chloride, heptachlorpropane, CCl_s. CHCl. CCl_s, boiling at 249°, and with chloroform, hexachlorpropane, CCl_s. CHCl. CHCl_s, boiling at 216°.

Likewise dichlorethylene, CHCl: CHCl, and chloroform give symmetrical pentachlorpropane, CHCl₂. CHCl. CHCl₂, boiling at 198°. See Chapter XX for the reverse reactions caused by aluminum chloride.

243. Stannic chloride causes an analogous addition of ethylenic or cyclohexenic chlorides to acid chlorides to form α -chlorketones. Aluminum chloride also can be used as catalyst in the reaction but is not so good.¹⁸⁷

¹⁸⁶ PRINS, J. prakt. Chem. (2), 89, 414 (1914).

¹⁸⁷ DARZENS, Compt. rend., 150, 707 (1910).

CHAPTER V

OXIDATIONS

I. Direct Oxidations by Gaseous Oxygen

- 244. The action of oxygen on various substances, or oxidations, can be divided into three groups:
- 1. Oxidations which take place spontaneously as soon as the oxidisable material and oxygen are brought together under the proper conditions of temperature and pressure.¹
- 2. Oxidations which are brought about by the simultaneous oxidation of certain substances called *auto-oxidisers*.
- 3. Oxidations effected by substances which are apparently unchanged and which are called oxidation catalysts.

At first sight only the latter seem to belong in the present treatise. But even in the first group, catalytic phenomena are of more or less importance. We have already mentioned (73) the influence of moisture on reactions. Practically, the amounts of water vapor contained in the air or in the oxygen, even when they are dried by the usual means, are sufficient to facilitate oxidations of the first class.

The case of induced oxidations, that is as a consequence of simultaneous oxidations, has been examined in Chapter III (150), and we have shown how we can sometimes pass from the mechanism of such reactions to catalytic oxidations which should be specially examined.

- 245. Platinum. The direct formation of a sort of unstable oxide on the surface of the platinum (154) permits us to explain the important rôle of this metal in many oxidations. Its activity should be proportional to its surface and it can be shown that the surface is immeasurably larger for platinum sponge and especially for the black than it is for the same amount of metal in foil or wire.
- 246. The use of platinum black enables us to effect many oxidations. Ethyl alcohol poured on platinum black is vigorously oxidised to acetaldehyde and acetic acid; the black is sometimes made incandescent and the alcohol may take fire.
- ² In cases of this kind it is practically impossible to eliminate the catalytic effect of the interior surfaces of the walls of the containing vessel and hence it is sometimes difficult to distinguish between reactions of this kind and those of Class 3.—H. D. Gress.

Formic and oxalic acids are burned to water and carbon dioxide.²
Alcohols are usually oxidised to aldehydes and even to acids.
Cinnamic aldehyde can be obtained thus from the corresponding alcohol.²

By oxidising glycerine by air in the presence of platinum black, the isomers, glyceric aldehyde and dihydroxyacetone, are obtained: 4

```
CH_2OH \cdot CHOH \cdot CH_2OH + O = H_2O + CH_2OH \cdot CHOH \cdot CHO

CH_2OH \cdot CH_2OH + O = H_2O + CH_2OH \cdot CO \cdot CH_2OH \cdot
```

However, platinum black has no effect on a mixture of carbon monoxide and oxygen.⁵

247. The results given by the black are irregular because its action is too violent, particularly at the beginning of the reaction.

By substituting for it, platinized asbestos where the active material is diluted by a large proportion of inert material, regular oxidation of vapors mixed with suitable amounts of oxygen or of air, is obtained. The manufacture of sulphur trioxide is only an application of this on the large scale.

248. Colloidal platinum (67) has intense oxidising power, greater than that of the black. It gives 50% carbon dioxide with a mixture of carbon monoxide and half its volume of oxygen.⁵

249. Platinum in very fine wire or very thin foil is employed industrially in the oxidation of ammonia gas by the oxygen of the air. The gaseous mixture, previously heated to about 300°, is passed over the metal which is thereby maintained in incandescence. Contact with the metal for one-five-hundredth of a second is sufficient to obtain a good yield of nitrous vapors which are easily transformed into nitric acid.

It furnishes also an excellent method for the regular oxidation of alcohols and of other sufficiently volatile organic substances. Trillat has described a method of operating which makes it easy to attain this end by the aid of a platinum wire which is heated by a current that can be regulated at will for any desired temperature and over which a current of air passes laden with the vapors of the substance to be oxidised.

- ² MULDER, Rec. Trav. Chim. Pays-Bas, 2, 44 (1883).
- 8 STRECKER, Annalen, 93, 370 (1855).
- 4 GRIMAUX, Bull. Soc. Chim. (2), 45, 481 (1886).
- ⁵ Paal, Berichte, 49 548 (1916).
- ⁶ The points that are used in pyrography for burning designs on wood contain leaves of platinum foil which are heated by the catalytic combustion of the mixture of air and combustible vapors forced over them.—E. E. R.
- ⁷ Better catalysts than platinum are known for the oxidation of many alcohols. See note to 254 infra.—H. D. Gibbs.
 - ³ Trillat, Bull. Soc. Chim. (3), 27, 797 (1902).

Under these conditions methyl alcohol is oxidised below 200° chiefly to formaldehyde with some methylal and water but no acid. The acid appears when the spiral reaches a dull red, at the same time that the formaldehyde and methylal increase. At a cherry red these decrease and the proportion of carbon dioxide increases with increase of incandescence.

The presence of water in the methyl alcohol favors the oxidation which goes best when 20% of water is present.

Ethyl alcohol is oxidised as low as 225° and readily at a dull red with a yield of 16.8% of acetaldehyde and 2.3% acetal. The results are less and less favorable as the molecular weight of the alcohol increases.

With propyl alcohol the yield of aldehyde is about the same as with ethyl, but is 12% for normal butyl alcohol and 5% for isobutyl. Isopropyl alcohol gives 16% acetone. Tertiary butyl alcohol breaks up, on oxidation, into formaldehyde, acetone and water.

Allyl alcohol gives 5.8% acroleïne, some acrylic acid, formaldehyde and glyoxal.

Glycol oxidises at 90°, raising the spiral to incandescence and yielding formaldehyde, glycolic aldehyde and glyoxal. Glycerine gives principally formaldehyde and acroleïne.

Aromatic alcohols likewise produce some of the corresponding aldehyde. Benzyl alcohol has furnished 4% benzaldehyde and cuminyl alcohol, 5.7% cuminic aldehyde.

Cinnamic alcohol gives some cinnamic aldehyde at a dull red and cinnamic acid and benzaldehyde at higher temperatures.

Isoeugenol oxidises at a dull red to give 2.9% vanilline mixed with the unchanged substance.¹⁰

- 250. The use of porous porcelain impregnated with platinum is advantageous for securing the complete oxidation of organic compounds in combustion analysis.¹¹
- 251. Metals of the Platinum Group. The various metals of this family may be used as sponge or better as black for the same purposes.

Palladium black gives good results.12

Osmium, a more moderate catalyst, sometimes has advantages. In the oxidation of cyclohexene, it gives some cyclohexenol accom-

⁹ TRILLAT, Bull. Soc. Chim. (3), 29, 35 (1903).

¹⁰ TRILLAT, Bull. Soc. Chim. (3), 29. 35 (1903).

¹¹ CARRABCO and BELLONI, J. Pharm. and Chim. (6), 27, 469; Chem. Centbl., 1908 (2), 95.

¹² WIELAND, Berichte, 46, 3327 (1913).

panied by adipic acid and other products. The other metals of the platinum family are not suitable for these reactions.

Tellurium may be used, but it is less active than osmium.18

Colloidal irridium can catalyze the oxidation of carbon monoxide as does colloidal platinum, but colloidal osmium is less efficient.¹⁴

252. Gold and Silver. Gold and silver can be substituted for platinum in the preparation of formaldehyde. Silvered asbestos obtained by the reduction of the nitrate by formic acid and asbestos gilded by the reduction of the chloride are more active than platinized asbestos (245).¹⁵

253. Copper. In the oxidation of methyl alcohol by the method of Trillat (248), the platinum spiral can be replaced by a roll of copper gauze heated to a dull red.

The results obtained are entirely similar. In operating thus with a current of 2.3 to 2.7 liters of air per minute, carrying 0.5 to 0.8 g. methyl alcohol, copper gauze gives a yield of 48.5% formaldehyde at 330°. There is at the same time production of carbon monoxide, carbon dioxide and water vapor.¹⁶

The direct oxidation of methane by air in contact with copper or silver is a practicable method for preparing formaldehyde. A mixture of one volume of moist air with three volumes of methane is passed over either of these metals or over a mixture of the two. The formaldehyde that is produced is taken out by contact with water and the residual gases are passed again over the catalyst.¹⁷

254. Fokin, operating under identical conditions with air saturated with methyl alcohol vapor passed over various catalysts, has obtained the following yields of formaldehyde (figured on the methyl alcohol used): 18

Gilded asbestos	71%
Silvered asbestos	64-66
Coppered asbestos	43-47
Platinized asbestos	5.2
Reduced cobalt	2.8
Manganese in powder	2
Aluminum turnings	
Reduced nickel	

- 18 WILLSTÄTTER and Sonnenfeld, Berichte, 46, 2952 (1913).
- 14 PAAL, Berichte, 49, 548 (1916).
- 18 FOKIN, J. Russian Phys. Chem. Soc., 45, 286 (1913); C. A., 7, 2227.
- ¹⁶ OBLOFF, J. Russian Phys. Chem. Soc., 39, 855 and 1023 (1907); C. A., 2, 263 and 1692.
- ¹⁷ VEREIN F. CHEM. IND., German patent, 286,731, J. Soc. Chem. Ind., 35, 73 (1916).
 - ¹⁸ Fokin, J. Russian Phys. Chem. Soc., 45, 286 (1913); C. A., 7, 2227.

A maximum yield of 84% was obtained by a mixture of silver and copper. The silvered or gilded asbestos requires an initial temperature of only 200-250° and the heat evolved is sufficient to maintain it at a suitable temperature.

Copper used alone requires continual heating, but this can be avoided by placing ahead of the copper gauze some fragments of pumice impregnated with platinum or palladium the incandescence of which heats the gas sufficiently.¹⁹

The presence of lead in the copper is unfavorable.

Ethyl, propyl, isobutyl and isoamyl alcohols may be oxidised under like conditions.²⁰ Ether is oxidised to formaldehyde and acetaldehyde.²¹ Various hydrocarbons have been submitted to regular oxidation by the same process but the products have not been fully studied.²²

255. As acetaldehyde can be prepared from acetylene (309), its direct oxidation to acetic acid is an interesting industrial problem.

It appears to be realized by the use of platinum; the aldehyde vapors carried by air or oxygen over platinized asbestos kept at 130-40° are regularly transformed into acetic acid.²⁵

256. The same metals may be used as catalysts for the direct oxidation of ammonia or amines.

Moist ammonia yields ammonium nitrite with a little nitrate and very little free nitrogen.

Moist methyl amine gives formaldehyde along with ammonium nitrite and nitrate, while ethyl amine gives some acetaldehyde.

Dimethylaniline produces formaldehyde and a complex aromatic amine.²⁴ Aniline, toluidine and pyridine are oxidised with the formation of complex oily products.²⁵

- 19 The oxidation of isopropyl alcohol has been extensively investigated by R. R. Williams and H. D. Gibbs in connection with the utilization of the waste unsaturated gases obtained in large quantities from the petroleum cracking stills. It was found that the best catalyst was brass (zinc and copper). The isopropyl alcohol is mixed with air and passed through brass gause at about 200°. With a catalytic chamber of a proper volume in relation to the radiation surface, the reaction is continuous and requires no external heat. The yield of acetone is over 90% of the theory. That the reaction is essentially a dehydrogenation is shown by passing the isopropyl alcohol over the catalyst without the oxygen of the atmosphere, acetone is formed but the necessary heat must be supplied externally. This work was done for the U. S. Government during the war but the report has not yet been published.—H. D. Gibbs.
 - ²⁰ Orloff, Ibid., 40, 203 (1908); C. A., 2, 3346.
 - ²¹ Orloff, Ibid., p. 799; C. A., 3, 1147.
 - 22 Orloff, Ibid., p. 652.
 - 28 DREYFUS, French patent, 487,412 (1918).
 - ²⁴ TRILLAT, Bull. Soc. Chim. (3), 29, 873 (1903).
 - ²⁵ ORLOFF, J. Russian Phys. Chem. Soc., 40, 659 (1908).

257. Carbon. The less combustible forms of carbon may serve as oxidation catalysts.

Coke at 200° aids in transforming toluene into benzoic acid. 26 27 Coal and lignite after being heated in the air to 300° are good oxidation catalysts between 150 and 300°; the action, being partly due to the oxide of iron which they contain, is increased by the addition of ferric oxide. They can be used in the oxidation of ethyl alcohol to acetaldehyde and acetic acid, and of toluene into benzaldehyde and benzoic acid. Anthracene gives anthraquinone and borneol forms camphor and camphoric acid. 28

258. Metallic Oxides. A large number of metallic oxides act as oxidation catalysts and for the most of them this property can be readily explained by the fact that they are readily reduced to the metals or to lower oxides by the substances to be oxidised and are readily reoxidised directly by oxygen. This is the case with the oxides of copper, nickel and cobalt. When alcohol vapors alone are passed over copper oxide moderately heated, aldehyde is formed and the oxide is reduced, but if the air is mixed with the alcohol vapors the copper is immediately reoxidised and can recommence the oxidation of the alcohol. A like explanation fits the case of ferric oxide, which can be reduced to a lower oxide which is reoxidised by the air. It is more difficult to perceive the mechanism in the case of oxides which can not be reduced to suboxides e. g. chromium sesquioxide which is, nevertheless, an excellent oxidation catalyst.²⁹

The catalytic activity of *iron sesquioxide*, such as is obtained by roasting pyrites, is utilized industrially in the manufacture of sulphuric acid by the contact process.

259. The use of metallic oxides as catalysts in the oxidation of organic compounds has until recent years been limited to copper oxide

²⁶ DENNSTEDT and HASSLER, German patent, 203,848, Chem. Centrbl., 1908, (2), 1750.

During the war various forms of carbon were extensively studied as adsorbents for gases and as catalysts for certain reactions. Very active forms of charcoal were developed by high heat treatments. These charcoals were found to be excellent clarifying agents for solutions, and some forms catalyzed certain reactions to a high degree. The reaction between chlorine and water was found to be quite rapid at low temperatures, even so low as 0° , and at 100° it is very vigorous. The reaction is $2 \text{ Cl}_2 + 2\text{H}_2\text{O} \rightarrow 4\text{ HCl} + \text{O}_2$. This would constitute a reversal of the Deacon process were it not for the fact that the oxygen does not appear as such but unites with the carbon gradually consuming the catalyst. See: The Production of Hydrochloric Acid from Chlorine and Water. Gibbs, J. Ind. and Eng. Chem., 12, 538 (1920).—H. D. Gibbs.

²⁸ Wood, Compt. rend., 145, 124 (1907); C. A., 1, 2690.

²⁹ SABATIER and MAILHE, Compt. rend., 142, 1394 (1906); C., 1906, (2), 402.

which is the real agent when copper is used, as has been said above. Sabatier and Mailhe have shown that the oxides of copper, nickel, and cobalt, as well as those of chromium, manganese, uranium, etc., have catalytic properties entirely comparable to those of finely divided platinum. When these oxides are heated to 200° in a mixture of oxygen with the vapors of aliphatic hydrocarbons (methane, pentane, hexane, and heptane), they become incandescent and maintain themselves so, giving mainly water and carbon dioxide, but also a certain amount of aldehyde and acid.²⁹

Almost simultaneously with the above work, Matignon and Trannoy have shown the possibility of realizing a *lamp without flame* by the aid of asbestos fibers impregnated with the oxides of iron, nickel, chromium, copper, manganese, cerium, and silver suspended in a mixture of air and ether vapor.⁸⁰

The use of ferric oxide between 175 and 300° permits the regular oxidation of toluene to benzaldehyde; the most favorable temperature is 280° and the yield of aldehyde may reach 20%. Above 280° the oxide becomes incandescent and there is partial charring of the products.

Employed in the same way, nickel oxide gives benzaldehyde above 150°, while at 270° incandescence begins to manifest itself.

With copper oxide (oxidised turnings), the reaction takes place between 180 and 260°. S1 S2

- MATIGNON and TRANNOY, Compt. rend., 142, 1210 (1906); C., 1906 (2), 202.
 - ⁸¹ Wood, Compt. rend., 145, 124 (1907), C. A., 1, 2690.
- 22 The catalytic oxidation of carbon monoxide at low temperatures may be brought about by certain metals such as platinum and palladium but the time of contact necessary for complete oxidation is quite great. Mixtures of certain metallic oxides are much more effective and may bring about the catalytic oxidation of carbon monoxide at room temperatures with a surprisingly short time of contact. These mixed-oxide catalysts require careful preparation in order that they may function under these conditions. Fineness of subdivision and intimacy of admixture of the ingredients are among the most essential conditions. The most important of this class of catalysts for the oxidation of carbon monoxide contains, as its essential constituent, manganese dioxide made by the method of Frémy (Compt. rend., 82, 1213 (1876). Copper oxide or silver oxide, when properly incorporated with this manganese dioxide, gives an excellent catalyst which is capable of effecting the catalytic oxidation with great rapidity even at temperatures somewhat below 0° C.

To prepare the catalyst, the Frémy oxide is washed free of sulphates and filtered on a Büchner funnel. This paste, usually containing about 60% of water, is analyzed for moisture by drying to constant weight at 130° in oxygen. A weighed amount of this paste is mixed with a large volume of cold water, care being taken to secure a uniform suspension. To this suspension is added such

260. Vanadium pentoxide is also a very active oxidation catalyst and can transform the vapors of ethyl alcohol mixed with air into acetaldehyde and acetic acid.³⁸ Acetaldehyde can also be changed to acetic acid; this oxidation is readily realized by passing a current of air through a solution of the aldehyde in glacial acetic acid containing oxides of vanadium,³⁴ uranium ³⁵ and iron.³⁶

261. Cerium oxide also can be employed for transforming acetaldehyde into acetic acid (256). The aldehyde mixed with 1% cerium oxide is submitted to the action of oxygen at two atmospheres or of air at higher pressures. The oxidation evolves heat and gives a yield of 95%.²⁷

an amount of a solution of copper or silver nitrate, as the case may be, as will give a mixture of 75% of manganese dioxide to 25% of the other oxide and, with continual vigorous stirring, a solution of sodium carbonate is run in till precipitation is just complete. The precipitate is filtered, carefully washed, and thoroughly dried at about 130°. In order to produce a harder and less friable product, it is well to compress the material in a filter press before drying. Silver oxide may be precipitated by caustic soda, but with copper, sodium carbonate must be used, the copper carbonate passing into the oxide during the drying. Both silver and copper oxides may be used in the catalyst. Certain other oxides, such as iron oxide, may be tolerated in limited amounts and appear to act only as diluents. When properly prepared, these catalysts will bring about the complete oxidation of carbon monoxide provided a sufficient amount of oxygen is present in the mixture. Moisture is rapidly absorbed by the catalyst, diminishing its activity, hence the gas mixture must be relatively dry for the oxidation to be catalytic.—J. C. W. Frazer.

- 88 NAUMANN, MOESER, and LINDENBAUM, J. prakt. Chem. (2), 75, 146 (1907).
- ³⁴ Vanadium pentoxide is an excellent catalyst for the oxidation of toluene to benzaldehyde, anthracene to anthraquinone, naphthalene to phthalic anhydride and other reactions of a similar nature.

Phthalic anhydride is produced in America almost exclusively by this process. Naphthalene is volatilized in an air stream and passed over the catalyst. The reaction begins at about 300° and attains a maximum yield at about 400 to 450°, equaling about 50% of the theoretical. [Gibbs, J. Ind. Eng. Chem., 11, 1031 (1919)].

Vanadium compounds have been extensively employed in the production of aniline black. [PINENEY, Brit. pat. 2745 of 1871, See Chem. News, 33, 116 (1876)].

Austerweil (U. S. pat. 979,247 (1910); C. A., 5, 972) used vanadium compounds in solution to catalyze the oxidization of borneol to camphor by nitric acid. — H. D. Gibbs.

- ss Recently the oxidation of benzene vapors by air in the presence of vanadium pentoxide has assumed commercial importance as a method for manufacturing maleic acid, Weiss and Downs, J. Ind. Eng. Chem., 12, 228 (1920), U. S. patents 1,318,631-2-3, Oct. 14, 1919, C. A., 14, 70; Can. pat. 192,766, Sept. 16, 1919, C. A., 13, 2683.—E. E. R.
 - ²⁶ Johnson, English patent 17,424 of 1911; J. Soc. Chem. Ind., 31, 772 (1912).
- ²⁷ FARBW. MRISTER, LUCIUS and BRÜNING, English patent 10,377 of 1914, J. Soc. Chem. Ind., 33, 981 (1914).

The use of cerium oxide permits acetic acid being made from acetylene in one operation by effecting the hydration (309) and oxidation simultaneously. It is sufficient to circulate a mixture of 130 parts acetylene and 80 to 100 parts oxygen through a mixture of 400 parts glacial acetic acid, 100 parts water, 50 parts mercuric nitrate, and 10 parts cerium oxide kept between 50 and 100°. 25

262. Anthracene can be transformed directly into anthraquinone by gaseous oxygen under pressure and in the presence of catalysts. **Osmium* peroxide in the small amount of 0.05% realizes this oxidation rapidly with oxygen under 10 atmospheres pressure. **O The same result can be obtained by keeping anthracene suspended in 30 parts water containing a little ammonia and 0.5 part copper oxide for 20 hours at 170° with compressed oxygen. **I

The mixture of oxides remaining from the manufacture of Welsbach incandescent mantles has been proposed as a catalyst for direct oxidation.⁴²

263. Metallic Chlorides. Anhydrous aluminum chloride, AlCl₂, causes the direct fixation of atmospheric oxygen by aromatic hydrocarbons. Benzene gives a certain amount of phenol and toluene yields m.cresole.⁴⁸

264. Manganous Salts. As has been stated in Chapter III (153), manganous salts are active agents of direct oxidation, particularly in water solution. This activity persists whatever be the acid constituent of the salt; it is observed in the salts of mineral acids, in the acetate, butyrate, benzoate and oxalate: it is sixteen times as great in the succinate as in the nitrate. We can assume that the manganous salt is partially hydrolyzed in water solution and that the resulting manganous hydroxide is oxidised to the dioxide by one atom of an oxygen molecule, the other oxidising the organic compound. The nascent manganese dioxide, in turn, would part with its extra oxygen to another portion of the organic compound and the manganous hydroxide thus regenerated would begin the cycle again. A trace of the manganous salt would thus be able to oxidise an unlimited amount of the oxidisable substance.

⁸⁸ Dreyfus, French patent 479,656, J. Soc. Chem. Ind., 35, 1179 (1916).

³⁹ The best catalyst yet found for oxidising anthracene to anthraquinone is vanadic oxide. The conditions are about the same as for the oxidation of naphthalene to phthalic anhydride.—H. D. Gibbs.

⁴⁰ HOFMANN, Berichte, 45, 3329 (1912).

⁴¹ German patent, 292,681.

⁴² Mason and Wilson, Proc. Chem. Soc., 21, 296 (1905); C., 1906 (1), 395.

⁴⁸ FRIEDEL and CRAFTS, Ann. Chim. Phys. (6), 14, 435 (1888).

⁴⁴ BERTRAND, Bull. Soc. Chim. (3), 17, 753 (1897).

Cerium salts may frequently act similarly (153).

265. Oxidation of Oils. The bleaching of oils can be effected by a moderate oxidation with warm air in the presence of catalytic oxides which doubtless act after being transformed into metallic soaps, the true decolorizers.

Palm oil through which a current of air is passed at 80-90° is bleached in four hours if 0.2% manganese borate is added. The same oil with 0.1% cobalt borate is bleached in 3.5 hours by the passage of less than its own volume of air. With the same proportion of nickel or iron borate, about three times as much air and 10 hours are required.⁴⁵

If the operation is carried on in an autoclave with compressed air, the addition of 0.02% of cobalt soap permits various oils to be bleached perfectly and rapidly.⁴⁶

266. The so-called drying oils, such as linseed and poppy seed, have the property of rapidly becoming thick in contact with air, which oxidises them, converting them into resinous substances which are almost insoluble in boiling alcohol. It has long been known that this drying power, depending on the oxidisability, is greatly increased by incorporating with the oils small proportions of salts of lead and particularly of manganese, the important accelerating agent appearing to be the metallic soap formed with the oil.⁴⁷

The metallic soaps that are the most active are those containing metals which are capable of several degrees of oxidation, particularly, cobalt, manganese, cerium, lead, chromium, iron, and uranium, while soaps containing bismuth, aluminum, mercury, and thallium are less active.⁴⁸

The direct oxidation of oils is retarded by moisture and accelerated by light. Elevation of temperature and increase of the pressure of the oxygen increase the velocity of the oxidation.⁴⁹

267. Metallic Silicates. Silicates can sometimes be substituted for the corresponding oxides. Kaolin (aluminum silicate) causes the union of hydrogen and oxygen at 230°.50

```
45 SASTRY, J. Chem. Soc., 107, 1828 (1916).
```

⁴⁶ RAI, J. Soc. Chem. Ind., 36, 948 (1917).

⁴⁷ LIVACHE, Compt. rend., 124, 1520 (1897); C., 1897 (2), 332.

⁴⁸ MACKEY and INGLE, J. Soc. Chem. Ind., 36, 317 (1917).

⁴⁹ FOKIN, Z. angew. Chem., 22, 1451 (1909).

⁵⁰ JOANNIS, Compt. rend., 158, 501 (1914); C. A., 9, 1866.

II. - Oxidations Carried Out with Oxidising Agents

268. Oxidations by Hydrogen Peroxide. The oxidation of organic compounds by hydrogen peroxide can be advantageously catalyzed by small quantities of ferrous or ferric salts (acetate).⁵¹

Methyl, ethyl, propyl, butyl, isobutyl, and isoamyl alcohols are oxidised to a mixture of aldehyde and acid, the acid being more abundant when ferrous oxalate is used than with the sulphate. The addition of wood charcoal favors the production of aldehyde. Manganous salts can be substituted for the iron.⁵²

Glycerine reacts vigorously to give glyceric aldehyde, along with a little dihydroxy-acetone. **Arabite yields an araboketose and dulcite, galactose. **S Malic acid passes into oxalacetic acid, HO₂C.-CO.CH₂.CO₂H.**S

Benzene is partially transformed into phenol and then to pyrocatechol; ⁵⁷ p.hydroxybenzaldehyde, HO.C₆H₄.CHO, gives protocatechuic aldehyde. ⁵⁸

Amines likewise undergo a regular oxidation to the corresponding aldehydes when they are warmed above 50° with hydrogen peroxide in presence of a ferrous salt; ethylamine giving acetaldehyde; isoamylamine, isovaleric aldehyde; benzylamine, benzaldehyde, while aminoethyl alcohol is changed, above 30°, to a mixture of glycolic aldehyde and glyoxal.⁵⁹

The use of the double cyanide of copper and potassium permits the oxidation of morphine hydrochloride by hydrogen peroxide to dehydromorphine and pseudomorphine.**

Furfural in alcoholic beverages can be destroyed slowly by the addition of 1% hydrogen peroxide and 0.01% manganese acetate.⁶¹

269. Oxidation by Nitric Acid. Vanadium pentoxide, employed

- ⁵¹ Fenton, J. Chem. Soc., 65, 899 (1894).
- ⁵² DOROSHEVSKII and BARDT, J. Russian Phys. Chem. Soc., 46, 754 (1914); C. A., 9, 1865.
 - 58 FENTON and JACKSON, J. Chem. Soc., 75, 575 (1899).
 - 54 Fenton and Jackson, Ibid., 75, 1 (1899).
 - ⁵⁵ NEUBERG, Berichte, 35, 962 (1902).
 - ⁵⁶ Fenton and Jones, J. Chem. Soc., 77, 69 (1900) and 79, 91 (1901).
 - ⁵⁷ Cross, Bevan and Herreno, Berichte, 33, 2015 (1900).
 - ⁵⁸ Sommer, German patent, 155,731, C., 1904 (2), 1631.
 - ⁵⁹ Suto, Biochem. Zeitschr., 71, 169 (1915); C. A., 9, 3059.
 - 60 DENIGES, Bull. Soc. Chim. (4), 9, 264 (1911).
 - 61 CHAUVIN, Ann. Falsif., 6, 463 (1913); C. A., 8, 981.

in the ratio of 0.1 g. to 50 g. cane sugar and 500 cc. nitric acid (density 1.4) causes the complete oxidation of the sugar in 20 to 30 hours in the cold to oxalic acid without the formation of saccharic, mucic, tartaric acids, etc., as by-products. Above 70°, carbon dioxide and water are obtained instead of oxalic acid.⁶²

In the presence of *mercuric nitrate*, nitric acid oxidises anthracene to *anthraquinone*. The reaction is finished in three hours if 117 parts anthracene suspended in 300 parts nitrobenzene are warmed to 30° with 460 parts 31% nitric acid in which three parts of mercury have been dissolved.⁶³

In the nitration of aromatic compounds by mixtures of nitric and sulphuric acids, the presence of a mercuric salt has no influence, but with nitric acid of density 1.3, it causes oxidation along with nitration or the substitution of a nucleus hydrogen by the phenolic hydroxyl group. Thus benzene, toluene, and ethyl-benzene give nitrophenols. It is possible to prepare 2,4-dinitrophenol and picric acid by heating benzene on the steam bath under reflux with 8 times its weight of nitric acid, density 1.3, and 15% mercuric nitrate. The oxidation must precede the nitration, since nitrobenzene is not oxidised by this treatment.⁶⁴

- 62 NAUMANN, MOESER, and LINDENBAUM, J. prakt. Chem. (2), 75, 148 (1907).
- 68 U.S. patent, 119,546.
- 64 Wollfenstein and Böters, Berichte, 46, 586 (1913).
- 65 In addition to vanadium and mercury compounds, a number of other substances have been found to accelerate oxidation by nitric acid. Disregarding the mechanism of reaction, oxides of nitrogen and nitrous acid may be considered as catalysts for oxidation by nitric acid. For instance, VELEY (Proc. Roy. Soc., 48, 458-9 (1891)) found that the presence of nitrous acid initiated the oxidation of copper, mercury and bismuth by 30% nitric acid. Oxides of nitrogen are mentioned a number of times in the patent literature as being necessary or desirable for the purpose of starting oxidation of organic compounds by nitric acid, especially in the manufacture of camphor. Molybdenum compounds, salts of manganese, iron, cerium and palladium, and even salts of calcium and magnesium have, under various conditions, been found to accelerate oxidations by nitric acid. Probably, in many cases, the acceleration produced by foreign substances is due to the reducing action of the substance on the nitric acid, with consequent formation of oxides of nitrogen. Thus the Commercial Research Company proposes to start the oxidizing action of nitric acid on aromatic hydrocarbons with side chains by means of formaldehyde, copper, zinc, starch or other reducing substance (Brit. Pat., 141,333 (1920)).

Nitration by means of nitric acid is likewise accelerated by dissolved oxides of nitrogen. KLEMENC and EKL (Monatsh. 39, 641-98 (1918)) studied the nitration of a number of phenol derivatives and concluded that pure nitric acid, free from dissolved nitrogen peroxide or nitrous acid, does not cause nitration.

HOLDERMANN (Berichte, 39, 1250 (1906)) obtained negative results in efforts to influence the position of the entering nitro-groups by nitrating in the pres-

270. Oxidations by Hypochlorites. The addition of a very small amount of a cobalt or nickel salt to a solution of an alkaline hypochlorite, or chloride of lime, causes the evolution of oxygen in the cold. ••

This oxidising mixture may be used for oxidising organic substances. It transforms o.nitrotoluene into o.nitrobenzaldehyde and acid.⁶⁷

By the same means, phenanthridene is oxidised to phenanthridene: 68

$$\begin{array}{cccc}
C_0H_4 - CH & C_0H_4 - CO \\
 & & & & & & \\
C_0H_4 - N & C_0H_4 - NH
\end{array}$$

and acridine into acridone:

$$C_{\theta}H_{\bullet}$$
 $C_{\theta}H_{\bullet}$
 $C_{\theta}H_{\bullet}$
 $C_{\theta}H_{\bullet}$
 $C_{\theta}H_{\bullet}$
 $C_{\theta}H_{\bullet}$
 $C_{\theta}H_{\bullet}$

271. Oxidations by Chlorates. The oxidation of aniline hydrochloride, in the preparation of aniline black, is carried out in the cold by a solution of potassium or sodium chlorate with the aid of metal catalysts, the most active of which is vanadium pentoxide, V₂O₅, of which one part is sufficient for 270,000 parts of aniline and the corresponding amount of chlorate. Salts of cerium and, to a less extent, those of copper and iron are useful catalysts but less powerful.

Osmium peroxide, OsO4, is at least as powerful as vanadium pent-

ence of catalysts, but an appreciably greater yield of dinitrobensene, from nitrobensene, was obtained by nitrating with, rather than without, a small amount of mercuric nitrate, under conditions otherwise similar (28.0% and 23.5% of theory respectively). Also, Holdermann obtained evidence that mercuric nitrate acts as catalyst in the nitration of beta-methylanthraquinone. For the control of the position of the entering nitro-group, the use of considerable quantities of different acids mixed with the nitric acid is more promising than the use of small amounts of metal salts. See Tingle and Blanck (J. Amer. Chem. Soc., 30, 1395 and 1587 (1908)).

Additional data on simultaneous nitration and oxidation in the presence of mercury compounds are given by Wolffenstein and Paar (Berichte, 46, 589 (1913)) and Vignon (Bull. Soc. Chim., 27, 547-50 (1920)). There are also a number of patents on this subject. Silver, copper and aluminum salts are said to act as catalysts as well as mercury.—A. S. RICHARDSON.

⁶⁶ FLEITMANN, Annalen, 134, 64 (1865).

⁸⁷ BADISCHE, German patent, 127,388, C., 1902 (1), 150.

⁶⁸ PICTET and PATET, Berichte, 26, 1962 (1893).

oxide and its use makes it possible to oxidise anthracene to anthraquinone by means of chlorates. 69

272. Oxidations by Sulphur Trioxide. Fuming sulphuric acid is frequently used as an oxidiser for organic compounds, the trioxide being reduced to the dioxide, but its action is not rapid enough in the absence of metallic catalysts, the most active being mercuric sulphate between 290 and 390°. The sulphates of potassium, magnesium, manganese, and cobalt are without effect, while those of nickel and iron act feebly. Only the sulphate of copper can replace that of mercury in practice but it is disadvantageous. It should be mentioned that a mixture of the sulphates of copper and mercury is more active than the two taken separately.

It has been proposed to add to the sulphuric acid the mixture of the rare earths (oxides of cerium, lanthanium, etc.) which is a byproduct in the manufacture of thorium nitrate, but this has not proved to be of any advantage.⁷²

In the Kjeldahl method for estimating nitrogen in organic compounds, the substances are boiled for a long time with fuming sulphuric acid. During the oxidation of the carbon and hydrogen, all the nitrogen passes into ammonia which is retained by the sulphuric acid without being burned. The addition of 0.5% mercuric sulphate triples the speed of the oxidation. In practice, 1 to 2 g. of mercury to 20 cc. acid is used for 5 to 7 g. of sample to be analyzed.

273. The chief application of oxidation by fuming sulphuric acid is the preparation of *phthalic acid* from *naphthalene*, a reaction which is the basis of one of the methods for making artificial indigo. When naphthaline is moderately heated with the acid, sulphonation takes place, but above 200° oxidation sets in. At 275° the oxidation rate is quintupled by 1% of mercuric sulphate.

274. In the presence of mercuric sulphate, fuming sulphuric acid can oxidise anthraquinone and further oxidise the hydroxyanthraquinones first formed. Thus anthraquinone and 1-hydroxyanthraquinone give quinizarine, 1,4-C₁₄H₂O₂(OH)₂.76

At 200-250°, alizarine gives quinalizarine, 1,2,5,8-C₁₄H₄O₂(OH)₄,

- 69 HOFMANN and SCHUMPELT, Berichte, 48, 816 (1915).
- ⁷⁰ Graebe, Berichte, 29, 2806 (1896).
- 71 Bradig and Brown, Z. physik. Chem., 46, 502 (1903).
- ⁷² Drz, Chem. Zeit., 29, 581 (1905); C., 1905 (2), 485.
- 78 WILFARTH, Chem. Centr., 1885, 17 and 113.
- 74 BADISCHE, German patent, 91,202.
- ⁷⁵ This process is being replaced by the high temperature air oxidation process. See note to 260 supra.—H. D. Gibbs.
 - ⁷⁶ WACKER, J. prakt. Chem. (2), 54, 88 (1896).

and 1,3,5,7-tetrahydroxyanthraquinone heated with 20 parts of sulphuric acid of 66° Bé. to the same temperature in the presence of 0.05 part mercuric sulphate, yields 1,3,4,5,7,8-hexahydroxyanthraquinone or anthracene blue. The addition of boric acid greatly favors these reactions.

275. Oxidations by Permanganates. The oxidation of aliphatic alcohols by potassium permanganate in presence of ferrous sulphate readily gives aldehydes but, on the contrary, in the presence of ferrous oxalate, the acids are formed quantitatively.⁷⁷

276. Oxidations by Persulphates. The persulphates of the alkalies mixed with nitric acid and a small quantity of silver nitrate are useful for oxidising organic compounds. The active agent is a silver peroxide or pernitrate which is constantly regenerated by the persulphate.⁷⁸

Benzene is transformed into quinone by this means, and oxalic acid is burned to carbon dioxide. Quinone is broken up into a number of products among which is found maleic acid. 79

277. Oxidations by Nitrobenzene. In the dye industry nitrobenzene is frequently used as an oxidising agent, being reduced to aniline; the presence of ferrous salts aids in these oxidations.

⁷⁷ DOROSHEVSKII and BARDT, J. Russian Phys. Chem. Soc., 46, 754 (1914); C. A., 9, 1865.

⁷⁸ KEMPF, Berichte, 38, 3963 (1905). BABOROVSKY and KUZMA, Z. Elektroch., 14, 196 (1908).

⁷⁹ KEMPF, Berichte, 39, 3715 (1906).

CHAPTER VI

VARIOUS SUBSTITUTIONS IN MOLECULES

§ 1. — INTRODUCTION OF CHLORINE, BROMINE AND IODINE

Chlorinations

278. The presence of anhydrous chlorides is a great aid in the direct chlorination of organic compounds, whether the chlorides are added as such or as the elements which are immediately transformed into the chlorides by the chlorine. There is no need to distinguish between these two.

Iodine or Iodine Chloride. Iodine, or iodine monochloride, in presence of an organic substance and of chlorine is changed to the trichloride which gives up chlorine to the organic substance, being itself reduced to the monochloride which starts all over again. With 2 to 12% of iodine it is easy to chlorinate benzene, toluene, the xylenes, etc., and also to transform carbon disulphide into carbon tetrachloride.

The chlorine compounds thus obtained are always mixed with a small amount of iodine derivatives formed by catalytic induction.

- 279. Bromine. This can catalyze chlorinations in the same manner as iodine, particularly in the preparation of carbon tetrachloride from the disulphide, but its use is less advantageous.
- 280. Sulphur. The immediate chlorination of sulphur by chlorine to several degrees of chlorination makes of it a good chlorinating agent of moderate activity which gives excellent results in some cases. Thus to transform acetic acid into chloracetic, chlorine is
- ¹ Müller, J. Chem. Soc., 15, 41 (1862); Jahresb., 1862, 414 and 1864, 524. Jungfleisch, Ann. Chim. Phys. (4), 15, 186 (1868).
- Bellstein and Geitner, Annalen, 139, 334 (1866). Limpricht, Ibid., 139,
 326 (1866). Hübner and Majert, Berichte, 6, 790 (1873).
- ⁸ Wollrath, Zeit. f. Chem., 1866, 488. Krüger, Berichte, 18, 1755 (1885). Kluge, Ibid., 18, 2099 (1885). Koch, Ibid., 23, 2319 (1890).
 - 4 English patent, 18,890 of 1899.
- ⁵ With iodine as a catalyst, the reaction may be stopped at the intermediate stage, Cl₂CSCl, though with iron, carbon tetrachloride is formed at once. (Helfrich and Reid, J. Amer. Chem. Soc., 43, 593 (1921)).—E. E. R.

passed into the boiling acid containing a small amount of sulphur. In two hours 8 parts of acetic acid are changed to 10 parts chloracetic containing but little acetyl chloride. In the cold, with a little sulphur or sulphur chloride, only acetyl chloride is obtained.

281. Phosphorus. Red phosphorus can be substituted for sulphur in the preparation of chloracetic acid (280).

The presence of phosphorus trichloride greatly facilitates the formation of benzyl chloride from toluene. By passing a current of chlorine into 100 parts of boiling toluene containing 1 part phosphorus trichloride (as far as possible in the sunlight), 80 parts of the desired product are obtained in eight hours.

282. Charcoal. Wood charcoal readily causes the chlorination of hydrogen to hydrochloric acid without explosion. By passing a mixture of equal volumes of carbon monoxide and chlorine through a long tube filled with fragments of charcoal, carbonyl chloride is obtained.⁸ Animal black gives even better results, a 30 cm. tube being sufficient.⁹ 10

A charcoal made by calcining blood with potassium carbonate can serve as a catalyst for the chlorination of organic substances between 250° and 400°. The progressive and complete chlorination of ethyl chloride can thus be readily obtained.¹¹

Carbon can likewise serve as a catalyst in the preparation of carbon tetrachloride from carbonyl chloride by a kind of auto-chlorination:

$$2\text{COCl}_2 = \text{CO}_2 + \text{CCl}_4$$

The carbonyl chloride vapors are passed through a succession of towers filled with coke or animal charcoal.12

- AUGER and BÉHAL, Bull. Soc. Chim. (3), 2, 145 (1889). RUSSANOF, J. Russian Phys. Chem. Soc., 1891, 1, 222; Berichte, 25, Ref. 334 (1892).
- 7 If sunlight is used no other catalyst is required. The chlorine reacts as fast as it can be passed in, even at 0°.—E. E. R.
 - 8 SCHIEL, Jahresb., 1864, 359.
 - 9 PATERNO, Gas. Chim. Ital., 8, 233 (1878).
- ¹⁰ Using 10 g. charcoal prepared from ox bones, ATKINSON, HEYCOCK and POPE (J. Chem. Soc., 117, 1410 (1920)) caused carbon monoxide and chlorine to combine at 40 to 50° as rapidly as the mixture could be passed into the U-tube containing the catalyst. After the preparation of 10 k. of phosgene this catalyst had lost none of its activity.

They found the activated charcoal from Army box respirator to be more active still, it being extremely efficient even at 14°. Even at 50° this catalyst does not cause the formation of hydrogen chloride in mixtures of chlorine and carbon monoxide containing hydrogen.— E. E. R.

- 11 DAMOISEAU, Compt. nend., 83, 60 (1876).
- 12 U. S. patent, 808,100.

283. Metallic Chlorides. Activity is possessed by the chlorides of polyvalent metals which have several degrees of chlorination, such as iron, thallium, molybdenum, antimony, tin, gold, vanadium, uranium, etc., and also by aluminum chloride and to a certain extent by zinc chloride but not by the chlorides of the alkaline or alkaline earth metals or of nickel, cobalt, manganese or lead.¹²

Moisture is usually unfavorable to their action.

284. Aluminum Chloride. Anhydrous aluminum chloride, or aluminum turnings, is an excellent chlorination catalyst.¹⁴ It readily realizes the transformation of carbon disulphide into carbon tetrachloride.¹⁵ The addition of 3% of it to benzene permits the progressive introduction of chlorine, going from the monochlor- to hexachlor-benzene.¹⁷

A mixture of equal volumes of chlorine and carbon monoxide passed over fragments of anhydrous aluminum chloride at 30-35°, is partially transformed into phosgene. The yield is better when the mixture of the gases is passed into chloroform saturated with aluminum chloride.¹⁸

285. Ferric Chloride. A little ferric chloride, for which may be substituted iron scale, iron sesquioxide or sulphide, ferrous carbonate, or even iron sulphate, gives good results with the substitution of chlorine in aromatic compounds.

By using one part ferric chloride and one of iron powder to 300 parts of benzene, one obtains a yield of 335 parts of monochlor-benzene with 37 parts of poly-chlor-. 19 20

- 18 WILLIERODT, J. prakt. Chem. (2), 34, 264 (1886) and 35, 391 (1887).
- 14 SHELIG, Annalen, 237, 178 (1887).
- ¹⁵ GOLDSCHMIDT and LARSEN, Z. physik. Chem., 48, 424 (1904). BORNWATER and HOLLEMAN, Rec. Trav. Chim. Pays-Bas, 31, 221 (1912).
 - 16 MOUNEYRAT, Bull. Soc. Chim. (3), 19, 262 (1898).
 - 17 MOUNEYRAT and Pouret, Compt. rend., 127, 1026 (1898); C., 1899 (1), 199.
 - 18 PLOTNIKOV, J. Russian Phys. Chem. Soc., 48, 457 (1916).
 - 19 FAHLBERG, LIST & Co., German patent, 219,242.
- ²⁰ It is usually assumed that the action of ferric chloride depends on the polyvalency of iron, supposing that a part of its chlorine is abstracted by the benzene leaving ferrous chloride which then combines with free chlorine to regenerate the ferric chloride.

In order to find whether benzene actually takes chlorine away from ferric chloride the following experiments were tried in my laboratory by H. K. Parker. Ferric chloride was sublimed, as it was formed, into a dry flask which was repeatedly evacuated to remove free chlorine. To this ferric chloride, 100 g. of benzene was added and kept at 40° for 30 hours, after which water was added. No chlorine was found in the benzene layer. The water layer contained 2.90 g. ferric chloride and 0.36 g. ferrous. Into a similar mixture of ferric chloride and benzene, dry chlorine was passed at 40° for 2 hours and extensive chlorination

It is equally satisfactory for the chlorination of toluene 21 or the xylenes.22

The use of ferric chloride facilitates the commercial preparation of carbon tetrachloride from carbon disulphide:

$$CS_2 + 3 Cl_2 = S_2Cl_2 + CCl_4$$

because it catalyzes the chlorination of the carbon disulphide by the sulphur chloride according to the equation:

$$CS_2 + 2 S_2Cl_2 = 6 S + CCl_4$$
.

The reaction commences at 60° and is continued at the boiling temperature of the mixture.^{28 24}

286. Molybdenum Chloride. Molybdenum chloride, MoCl_s, is an excellent catalyst in the aromatic series and, when used to the amount of 0.5%, permits successive stages of chlorination. Its use is of no advantage in the aliphatic series.²⁵

287. Antimony Chlorides. The chlorides of antimony (which can be replaced by the powdered metal or by the oxide) are frequently employed as carriers in chlorinations. They are more active than iodine and permit the complete chlorination of benzene.²⁶

They are useful in transforming carbon disulphide into the tetrachloride.²⁷

The successive use of iodine and of antimony pentachloride enables us to pass directly from benzyl chloride to hexachlor- and heptachlortoluene.²⁸

288. Tin Chloride. Stannic chloride (which can be replaced by the metal or the oxide) can also give good effects.²⁹ Its action, as

took place. At the end there was 30 g. bensene still unchlorinated and treatment with water showed only 0.04 g. ferrous iron.

These experiments show that the reduction of ferric chloride by a large excess of benzene is very slight. It seems to me best to regard the action of ferric chloride as analogous to that of aluminum chloride in this reaction, see note to 157.—E. E. R.

- 21 SEELIG, Annalen, 237, 152 (1887).
- ²² Claus and Burstmet, J. prakt. Chem. (2), 41, 552 (1890).
- ²⁸ MÜLLER and DUBOIS, German patent, 72,999. English patent, 19,628 of 1893.
- ²⁴ With iron as catalyst, it is impossible to stop at the intermediate, Cl_{*}CSCl.—E. E. R.
 - ²⁵ Aronheim, Berichte, 8, 1400 (1875). Similio, Annalen, 237, 152 (1887).
 - 26 Müller, Zeit. Chem. Pharm., 1864, 40.
 - 27 HOFMANN, Annalen, 115, 264 (1860).
 - 28 Beilstein and Kuhlberg, Annalen, 150, 306 (1869).
 - 29 PÉTRICOU, Bull. Soc. Chim. (3), 3, 189 (1890).

without doubt is the action of all chlorides used to aid direct chlorinations, is proportional to its concentration.²⁰

289. Aluminum Bromide. Its use permits the direct preparation of *perchlorethane*, CCl₂. CCl₃, starting with acetylene tetrabromide, CHBr₂. CHBr₂, or with ethylene bromide.³¹

Brominations

- 290. Anhydrous chlorides and bromides are more or less active agents in bromination just as in chlorination. The hydrobromic acid produced in the reaction is the product most readily followed.²²
- 291. Iodine. Iodine, or rather iodine bromide, which is the immediate product, is frequently used and leads especially to the bromination of the aromatic nucleus.²²
- 292. Manganese. Powdered metallic manganese is an excellent catalyst for the bromination of benzene, toluene, and xylene. With 3 g. of the powdered metal and bromine, 18 g. benzene is completely converted into monobrombenzene in 90 hours in the cold, without the metal suffering any appreciable attack.³⁴ The slight traces of bromide formed on the surface are doubtless sufficient to catalyze the reaction.
- 293. Aluminum Chloride. A small proportion is sufficient to effect the regular bromination of most organic compounds. Thus 1 g. can cause the bromination of 120 g. benzene.²⁵

We may put alongside of the brominations catalyzed by aluminum chloride the migration, which it causes, of the bromine of tribromphenol to benzene, so or toluene, which are thereby transformed to brombenzene or m.bromtoluene with the production of phenol.

Aluminum bromide causes a regular bromination of toluene.²⁸
Zinc Chloride and Bromide. Zinc chloride or metallic zinc which is changed to the bromide may be effective.³⁹

- ⁸⁰ Goldschmidt and Larsen, Z. physik. Chem., 48, 424 (1904).
- 81 MOUNEYRAT, Bull. Soc. Chim. (3), 19, 262 (1898).
- ⁸² Gustavson, J. prakt. Chem. (2), 62, 281 (1900).
- ³⁸ Rillier and Ador, Berichte, 8, 1287 (1875). Jacobsen, Ibid., 17, 2372 (1884) and 18, 359 (1885). Bruner, Chem. Cent., 1900 (2), 257.
 - 34 DUCELLIEZ, GAY, and RAYNAUD, Bull. Soc. Chim. (4), 15, 737 (1914).
- ³⁵ FITTIG, Annalen, 121, 361 (1862). Lamot, Bull. Soc. Chim. (2), 48, 210 (1887). Roux, Ann. Chem. Phys. (6), 12, 347 (1887).
 - ³⁶ Kohn and Milliam, Monatsh. Chem., 30, 407 (1909).
 - ²⁷ Kohn and Bum, Ibid., 33, 923 (1912).
 - 38 GUSTAVSON, J. Russian. Phys. Chem. Soc., 9, 286 (1877).
 - 89 SCHIAPARRILI, Gaz. Chim. Ital., 11, 70 (1882).

Ferric Chloride or Bromide. Ferric chloride or finely divided iron (which changes to the bromide) is a good bromination catalyst.40

Cyclobutene bromide, · · · , brominates in the presence of CH₂ - CHBr

iron powder to tetrabrombutane, the ring being opened.41

Mercuric Chloride or Bromide. These may be used as brominating agents.42 Without doubt the simultaneous formation of aluminum and mercuric bromides is the cause of the remarkable activity of aluminum amalgam as a bromination catalyst.48

Introduction of Iodine

294. The direct introduction of iodine into organic molecules is very difficult but may sometimes be accomplished by the aid of ferric chloride, as is the case with benzene. The yield of iodide thus formed is low.44

§ 2. — ADDITION OF SULPHUR

295. Anhydrous aluminum chloride can cause the addition of sulphur to benzene at 75-80°. Thiophenol, C.H. SH, and products derived from it, phenyl sulphide and phenylene sulphide, are thus obtained.45

296. The direct sulphuration of diphenylamine, by heating the amine with sulphur, requires a temperature of 200 to 265° for 6 to 8 hours: 46

$$NH \frac{C_6H_6}{C_6H_5} + 2S = H_2S + S \frac{C_6H_6}{C_6M_6}NH.$$

In the presence of iodine the reaction is complete in 10 minutes at 185°, giving a quantitative yield of thiodiphenyl-amine instead of 50 to 60%. Thiodinaphthyl amines, etc., are also prepared in good yields.47

- 40 SCHENFELEN, Annalen, 231, 164 (1885).
- 41 WILLSTÄTTER and BRUCE, Berichte, 40, 3979 (1907).
- 42 LAZAREW.
- 48 Cohen and Dakin, J. Chem. Soc., 75, 893 (1899).
- 44 LOTHAR MEYER, Annalen, 231, 195 (1885).
- 45 Friedel and Crafts, Bull. Soc. Chim. (2), 39, 306 (1883).
- 46 BERNTHSEN, Annalen, 230, 77 (1885).
- 47 Knozvenagel, J. prakt. Chem. (2), 89, 11 (1914).

§ 3. — ADDITION OF SULPHUR DIOXIDE

297. Benzene warmed with aluminum chloride absorbs sulphur dioxide readily giving benzene sulphinic acid, C₆H₅. SO₂H. ⁴⁸ The reaction is accelerated by the presence of hydrochloric acid and is doubtless due to the formation of an unstable addition product which reacts with the benzene in the presence of the aluminum chloride and hydrochloric acid.⁴⁹

§4.—ADDITION OF CARBON MONOXIDE

298. The direct addition of carbon monoxide to hydrocarbons is an exceptional reaction which can be realized in only a small number of cases. However, the use of aluminum chloride or bromide makes it possible with aromatic hydrocarbons.

A mixture of carbon monoxide and hydrogen chloride is passed for several hours into benzene containing aluminum chloride and 10% cuprous chloride at 40 to 50°.

It can be assumed that the carbon monoxide dissolves on account of the cuprous chloride and forms formyl chloride, H.CO.Cl, which then reacts as an acid chloride on the benzene in the presence of aluminum chloride (891). We have in the end:

$$C_6H_6 + CO = C_6H_5$$
. CHO.

The yield is 90%.⁵⁰ Likewise from toluene and aluminum chloride, p.toluic aldehyde, CH₃. C₆H₄. CHO, with a yield of 73%; ⁵¹ o.Xylene gives, by the same method, 1,2 dimethyl-benzaldehyde (4). p.Xylene and mesitylene give analogous results.⁵²

The presence of the cuprous chloride and the hydrogen chloride seem to be superfluous and it is sufficient to cause the carbon monoxide under pressures of from 40 to 90 atmospheres to act on the benzene in the presence of aluminum chloride and a little hydrogen chloride.⁵⁵

- 48 FRIEDEL and CRAFTS, Ann. Chim. Phys. (6), 14, 443 (1888).
- ⁴⁹ Knoevenagel and Kenner, Berichte, 41, 3315 (1908). Andrianowski, Bull. Soc. Chim. (2), 31, 199 and 495 (1879).
- ⁵⁰ REFORMATSKI, J. Russian Phys. Chem. Soc. 33, 154 (1901); C., 1901 (1), 1226.
- ⁵¹ GATTERMANN and Koch, Berichte, 30, 1623 (1897) and GATTERMANN, Ibid., 31, 1149 (1898). English patent, 13,709 of 1897.
- ⁵² BAYER AND Co., Chem. Cent., 1898, 932. HARDING and COHEN, J. Amer. Chem. Soc., 23, 594 (1901).
 - 58 English patent, 3,152 of 1915; J. Soc. Chem. Ind., 35, 384 (1916).

§ 5. — INTRODUCTION OF METALLIC ATOMS

Formation of Alcoholates

299. Aluminum alcoholates are formed by the direct action of aluminum amalgam on alcohols thoroughly freed from water.⁵⁴ But the presence of a catalyst enables them to be prepared directly from aluminum. It is sufficient to add a little mercuric chloride, iodine or even ethyl iodide. Thus ordinary absolute alcohol readily gives aluminum ethylate, Al(OC₂H₅)₈, a solid melting at 134° which can be isolated by distilling at 15 mm. pressure.⁵⁶

Production of Mixed Organo-Magnesium Compounds

300. The production of mixed organo-magnesium compounds from organic halides is equivalent to the addition of the magnesium atom to the organic molecule:

$$Mg + RBr = Mg$$
 R
 Br

This reaction is usually carried out in anhydrous ether which plays the rôle of catalyst in their formation. It is possible to carry out the reaction in benzene in the presence of a small amount of ether. Without doubt, we have in succession:

$$RBr + (C_2H_5)_2O = \frac{C_2H_5}{C_2H_5}O \frac{R}{Br}$$

and then:

$$C_2H_5 O R + Mg = Mg R + O C_2H_5$$

$$C_2H_5 O Br + Mg = Mg R + O C_2H_5$$
Therefore the state of the state of

The regenerated ether can repeat the first reaction.

301. The ethyl ether as catalyst can be replaced by other ethers, amyl ether, etc., or even by a small quantity of a tertiary amine such as dimethyl aniline, the reaction taking place in benzene, toluene, hexane, or ligroïne. In this case the temporary addition product would be: 56

$$\begin{array}{c|c} C_0H_0 & R \\ CH_2 & N \\ CH_2 & Br. \end{array}$$

⁵⁴ TISTCHENKO, J. Russian Phys. Chem. Soc., 31, 483 (1899).

MHISTER, LUCIUS and BRÜNING, German patent, 286,596; J. Soc. Chem.
 Ind., 34, 1168 (1915).
 TSCHELINZEFF, Berichte, 37, 4534 (1904).

302. The formation of organo-magnesium halides is easy with organic bromides or iodides but it is greatly facilitated by the presence of a suitable catalyst, iodine, hydriodic acid or an alkyl iodide such as ethyl iodide.

The addition of such catalysts is indispensable for the formation of these derivatives from aliphatic or cycloaliphatic chlorides, but even with this assistance they can not be prepared from aromatic chlorides.

According to Zelinski, iodine and magnesium produce in anhydrous ether some of the compound, MgI₂.2(C₂H₅)₂O, which he was able to isolate and which would start the reaction.⁵⁷

303. A certain number of substances hinder the formation of the organo-magnesium compounds. We may mention anisol, ethyl acetate, chloroform, and carbon disulphide which act as negative catalysts (11).

304. For the preparation of mixed organo-zinc compounds, Blaise uses pure *ethyl acetate* as a catalyst instead of ether and operates in a toluene or petroleum ether solution. Actually one-third of a molecule of ethyl acetate is used for one molecule of the alkyl iodide.⁵²

⁵⁷ Zelinski, J. Russian Phys. Chem. Soc., 35, 399 (1903).

⁵⁸ BLAISE, Bull. Soc. Chim., 1911, Conference, 7.

CHAPTER VII

HYDRATIONS

305. Hydration reactions can be separated into two distinct groups according to whether the water is added without splitting the molecule or whether the addition of the water causes the original molecule to break up into two or more new ones.

As examples of the first group we have the addition of water to unsaturated hydrocarbons giving alcohols or ketones, or to nitriles and imides.

Reactions of the second class are more frequent, such as the saponification of esters, the decomposition of acetals and glucosides, the hydrolysis of amides, oximes, hydrazones, semicarbazones, etc.

More or less concentrated *mineral acids* are very powerful agents for realizing the various hydration reactions, whether in concentrated form, they give rise to unstable temporary addition products which decompose to form the hydration products and to regenerate the acids, or whether they act in dilute solution in consequence of their electrolytic dissociation, the chief factors being the hydrogen ions so liberated.

Water solutions of the strong bases, either the alkalies or alkaline earths, can often realize hydrations which water alone can usually accomplish but at a much slower rate or a much higher temperature.

1. - Fixation of Water by Addition

306. Ethylene Compounds. Moderately concentrated sulphuric acid enables us to add, in the cold, a molecule of water to isobutylene, (CH₂)₂C: CH₂, to give trimethyl-carbinol, (CH₂)₃. COH.¹² By adding amylene, little by little, to a mixture of concentrated sulphuric acid and ice, diluting with ice water, washing with soda, and distilling the product, dimethyl-ethyl carbinol is obtained with a yield of 85 to 92% of the amylene.²

Likewise 3-methylpentene, CH₂: CH . CH(CH₂) . CH₃ . CH₃, adds water to give the corresponding tertiary alcohol.

- ¹ Butlerow, Annalen, 144, 22 (1867).
- ² Isopropyl alcohol is now manufactured by absorbing in sulphuric acid the propylene from the gases resulting from the cracking of heavy hydrocarbons.—E. E. R.
 - ⁸ Adams, Kamm, and Marvel, J. Amer. Chem. Soc., 40, 1950 (1918).

With 85% sulphuric acid hexene-1 and heptene-3 give the secondary alcohols, accompanied by a certain amount of the sulphuric acid esters, while the 100% acid only polymerizes the hydrocarbons.

At 45° sulphuric acid effects the addition of water to iso-oleic acid which is changed to hydroxystearic acid.

307. Dilute *nitric acid* provokes the rapid hydration of *pinene*, $C_{10}H_{16}$, in alcohol solution, at the ordinary temperature to form terpine, $C_{10}H_{20}O_2$.

Hydrochloric acid also can cause the addition of water. By digesting for three hours in the light a mixture of croton aldehyde, $CH_3 \cdot CH \cdot CHO$, and hydrochloric acid, the aldol, β -hydroxy-butyric aldehyde, $CH_3 \cdot CH(OH) \cdot CH_3 \cdot CHO$, is formed.

308. Doubly Unsaturated Compounds. Acetylene hydrocarbons and their allenic isomers can add water in the presence of sulphuric acid and other catalysts to give ketones:

and
$$R.C:C.R' + H_2O = R.CO.CH_2.R'$$

$$R'' + H_2O = R'CH.CO.CH'R''$$

With sulphuric acid, the reaction is carried out by dissolving the hydrocarbon in the cold concentrated acid and pouring this solution immediately on to ice.*

The mechanism appears to be the formation of an unstable sulphuric acid derivative which decomposes on contact with water to form an unsaturated alcohol which immediately isomerizes into the ketone. Thus with ethyl-acetylene, we should have successively:

$$CH_{2}.CH_{2}.C:CH + H_{2}SO_{4} = CH_{2}.CH_{2}C CH_{2}$$

$$CH_{2}.CH_{2}.CH_{2}.CH_{2}$$

$$CH_{3}.CH_{2}.CH_{2}.CO(OH):CH_{2}$$

$$CH_{3}.CH_{2}C(OH):CH_{3} = CH_{3}.CH_{2}.CO.CH_{3}.$$

In the case of true acetylene hydrocarbons, the product is a methyl ketone. With disubstituted acetylenes, two isomeric ketones are obtained. This is the case with methylamylacetylene.

- 4 Brooks and Humphrey, Ibid., 40, 822 (1918).
- ⁵ SAYTZEFF, J. prakt. Chem. (2), 37, 284 (1888).
- ⁶ Wiggers, Annalen, 57, 247 (1846).
- 7 WURTZ, Bull. Soc. Chim. (2), 42, 286 (1884).
- 8 BÉHAL, Bull. Soc. Chim. (2), 47, 33 (1887).
- ⁹ Béhal, *Ibid*. (2), 50, 359 (1888).

Acetylene should give acetaldehyde, but this condenses with loss of water (795) and crotonic aldehyde is the chief product.¹⁰

309. Water solutions of mercuric salts, the chloride, bromide, and sulphate, cause the same formation of ketones in consequence of the temporary production of combinations of the hydrocarbon and the salt, which are then decomposed by water. Thus allylene, CH₂.-C:CH, gives acetone, CH₂.CO.CH₃.

Acetylene behaves normally, yielding acetaldehyde: 11

$$CH:CH + H_2O = CH_3.CHO.$$

Acetylene is absorbed at 25 to 45° in a solution of mercuric oxide in water containing 45% or less sulphuric acid, or 25% phosphoric acid. The solution saturated with the gas is warmed to 80 to 100° when acetaldehyde is given off. The solution is then cooled and made to take up more gas and so on. By a number of repetitions the mercuric salt produces 20 times its own weight of acetaldehyde.¹² A stronger solution of sulphuric acid is unfavorable as it would cause the formation of crotonic aldehyde and other condensation products. (See the direct formation of acetic acid (261).)

- 310. This hydration of acetylene to acetaldehyde can likewise be accomplished by passing the moist gas over zinc, nickel, or ferrous oxides at 300°. There is the formation of a certain amount of acetal-dehyde and also of crotonic aldehyde. If the moist acetylene contains ammonia, the formation of acetaldehyde is shown by the deposit of crystals of aldehyde ammonia.¹⁸
- 311. Nitriles. Nitriles dissolved by gentle warming in sulphuric acid diluted with 20% of water, are transformed into amides. The same transformation can be effected also by caustic soda and potash; but, especially if the operation is carried on in alcohol solution in the neighborhood of 100°, the hydration may go so far as to break the amide down into ammonia and the acid, which is at least partially neutralized by the alkali.
- 312. Imides. It is the same way with imides, succinimide, CH₂-CO
 NH, warmed with a small amount of baryta water, gives CH₂-CO at first amido-succinic acid, H₂N.CO.CH₂.CH₂.CO₂H,¹⁴ the further hydration of which yields succinic acid.
 - 10 BERTHELOT, Compt. rend., 128, 336 (1899).
 - ¹¹ Kutscheroff, Berichte, 17, 13 (1884).
 - ¹² Dreyfus, French patent, 487,411 (1918).
- ¹⁸ CHICHIBABINE, J. Russian Phys. Chem. Soc., 47, 703 (1915); C. A., 9, 2512 (1915).

¹⁴ TEUCHERT, Annalen, 134, 136 (1865).

aqueous ammonia, adds a molecule of water to give asparagine, HOOC.CH,.CH(NH,).CONH,.15

Acetaldehyde in water solution causes cyanogen to add two molecules of water to form oxamide, H.N. CO. CO. NH₂. 16

II. - Hydrations with Decomposition

313. A hydration which results in the decomposition of the molecule is usually called hydrolysis.

§ 1. HYDROLYSIS IN WATER SOLUTION

Hydrolysis of Esters. The hydrolysis of esters is known as saponification.

When a water solution of methyl acetate or ethyl acetate is kept in the cold, there is a slow decomposition by water to give the alcohol and free acid:

$$CH_{2}$$
, CO_{2} , $C_{2}H_{5} + H_{6}O = CH_{2}$, $CO_{2}H + C_{2}H_{5}$, OH

The reverse reaction of esterification tends to reform the ester so the decomposition is never complete; the reaction tends to an equilibrium, the more water there is present, the more ester will be decomposed, but this limit is not reached at ordinary temperatures till after some years. In several days, the amount of ester decomposed is only about 1%. On the contrary, if a small amount of hydrochloric acid, or other strong acid, be added to the mixture, the reaction becomes very rapid, the limit being reached in 24 hours.

Furthermore, the acid added is in no way changed. It is entirely precipitated by silver nitrate which shows that it has not formed an appreciable amount of ethyl chloride. Up to a certain limit, the saponifying power of the acid is proportional to its concentration; and for different acids, at the same molecular concentrations, the saponifying power is proportional to the strength of the acid which is measured by its electrolytic dissociation, and consequently this activity is defined by the number of hydrogen ions in a unit volume of the solution. Hydriodic, hydrobromic, nitric and chloric acids, which are strongly ionized, are consequently powerful catalysts, and so is sulphuric acid which is most commonly used for this purpose.

¹⁵ KOERNER and MENOZZI, Gas. Chim. Ital., 17, 173 (1887).

¹⁶ Liebig, Annalen, 113, 246 (1860).

- 314. This action is general and applies equally well to the saponification of fats which are esters of glycerine with the fatty acids. A fat heated with water and 4% sulphuric acid to 120° is completely hydrolyzed in 12 hours, 42% of the fatty acids being liberated in the first hour. To produce a like decomposition with water alone requires heating to 220° in an autoclave.¹⁷
- 315. In very dilute solutions the velocity of saponification is the same with hydrochloric, hydrobromic, hydriodic, nitric, chloric and methyl sulphuric acids of the same acidity and is proportional to the concentration of the acid.

It is the same for all the esters of a given organic acid with different primary alcohols.¹⁸

- 316. On the other hand, the velocity changes greatly when the organic acid from which the ester is derived is changed. Thus for ethyl esters the saponification velocity of the *formate* is 25 times that of the *acetate*, 50 times that of the *isobutyrate*, 100 times that of the *valerate*, and 4,000 times that of the *benzoate*.
- 317. The presence of neutral metallic salts modifies the velocity, chlorides accelerating the saponification by hydrochloric acid, while sulphates retard the action of sulphuric acid.

Pressure may also have an effect; in the case of the saponification of methyl acetate by hydrochloric acid, pressure increases the velocity.

318. The soluble bases, potassium, sodium, barium, and calcium hydroxides have an analogous effect which, at first sight, would be attributed to the affinity of the base for the acid liberated, were the amount of ester saponified not disproportionate to the amount of base reacting. The real reason for the saponification is again found in the dissociation of the bases in dilute solution into ions.

The saponification of fats has given clear evidence on this point. The amount of lime required to saturate all of the fatty acids of a fat is about 9.7%, and in practice might reach 12 to 14%, but experience shows that 1% is sufficient for complete saponification in water at 190° under 12 atmospheres pressure, while 3% produces this result in 8 to 10 hours, at 170° and 8 atmospheres.²⁰

It appears from the above figures that, in spite of the additional energy liberated by the combination of the bases with the acids, the catalytic activity of the bases is less than that of the strong acids.

¹⁷ Lewkowitsch, Confér. à la Soc. Chim., 1909, 12.

¹⁸ Löwenherz, Z. phys. Chem., 15, 395 (1894).

¹⁹ TEST, J. prakt. Chem. (2), 34, 353 (1886). Eules, Z. phys. Chem., 32, 348 (1900).

²⁰ LEWKOWITSCH, loc. cit., p. 8.

319. For different strong bases, in very dilute solution and at the same concentration, the saponification velocity is independent of the nature of the base, whether it be potassium, sodium, barium or calcium hydroxide, but is proportional to the concentration of the base.

For esters derived from the same acid, the velocity changes greatly with the alcohol, thus methyl acetate is saponified twice as rapidly by caustic soda in the cold as is isobutyl acetate.²¹ The influence of the nature of the acid is less than it is in saponification by strong acids. Thus when methyl esters are saponified by caustic soda at 14°, the velocity for the acetate is about double that for the isobutyrate, six times that for the isovalerate, and quadruple that for the benzoate.

In the saponification of ethyl acetate by bases, the presence of neutral salts cuts down the velocity.²¹

320. The saponification of chlorine derivatives is not usually possible, but benzal chloride and benzotrichloride are hydrolyzed to benzaldehyde and benzoic acid by water alone when heated under pressure.

The saponification of benzal chloride, C₆H₅CHCl₂, by water alone requires a temperature of 140–160°.²² In the factory it is usually effected by means of milk of lime. In the presence of *iron powder*, the reaction can be carried out at 90–95°.²⁸

It is the same way with benzotrichloride which is readily transformed into benzoic acid in the presence of iron, ferric chloride, oxide or benzoate.²³

321. Ethers. Water alone breaks up ethers into two molecules of the alcohols very slowly.²⁴ The addition of small quantities of sulphuric acid greatly accelerates the reaction.²⁵

322. Acetals. Acetals can be regarded as mixed ethers derived from alcohols and the unstable gylcols of which aldehydes and ketones are the anhydrides. Their hydrolysis cannot be accomplished by water alone, nor by alkalies, even when hot. On the contrary, it is easily effected by boiling with either dilute hydrochloric acid or with sulphuric acid diluted with four volumes of water, the aldehyde and the alcohol being set free.

323. Polysaccharides. Polyoses and polysaccharides such as sucrose, lactose, maltose, trehalose and even starch, dextrine and

²¹ ARRHENIUS, Z. phys. Chem., 1, 110 (1887).

²² Limpricht, Annalen, 139, 319 (1866).

²⁸ SCHULTZE, German patent, 82,927.

²⁴ LIEBEN, Annalen, 165, 136 (1873).

²⁵ ERLENMEYER, Zeit. f. Chemie, 4, 343 (1868).

cellulose can be regarded as ethers or acetals involving the many alcohol groups and the aldehyde or ketone groups of the simple hexoses. Their hydrolysis into the simple sugars can be realized more or less readily by the aid of small quantities of acids as catalysts.

324. The inversion of cane sugar, that is, its complete hydrolysis into glucose and fructose, can be brought about by traces of mineral acids and is, perhaps, the earliest catalytic reaction to be observed. Hydrochloric, sulphuric, or even oxalic acid can be used and the velocity of the hydrolysis is proportional to the concentration of the hydrogen ions resulting from the electrolytic dissociation of the acid. Concentrated sugar solutions are rapidly inverted by traces of acid. A solution containing 80 g. sugar to 20 g. water, with the addition of 0.004 g. hydrochloric acid, is completely inverted by boiling for an hour. Even carbonic acid can cause this reaction, slowly in the cold, rapidly when heated. A sugar solution saturated with carbon dioxide and heated for an hour in a sealed tube is completely inverted.

The velocity of the inversion of sugar by strong acids is increased by the addition of neutral salts.²⁹

When the reaction is carried on in alcohol solution, the velocity varies considerably with the proportion and nature of the solvent.²⁰ Increase in the concentration of the sugar increases the velocity.²¹

High pressure diminishes the velocity of inversion by hydrochloric acid, the diminution being about 1% for 100 atmospheres.²²

325. The hydrolysis of maltose is slower than that of cane sugar, requiring at least three hours of boiling with 3% sulphuric acid.³³

Trehalose is slowly hydrolyzed into glucose by warming with dilute sulphuric acid.34

326. Sulphuric acid diluted with 33 parts of water is used in the commercial preparation of glucose by the hydrolysis of starch at 100° for three hours. The addition of a little nitric acid to the sul-

- 26 CLEMENT and DESORMES, Ann. Chim. Phys., 59, 329 (1806).
- 27 WOHL, Berichte, 23, 2086 (1890).
- 28 LIPPMANN, Berichte, 13, 1823 (1880).
- ²⁹ SPOHE, Z. physik. Chem., 2, 194 (1888). ARRHENIUS, Ibid., 4, 226 (1889). Euler, Ibid., 32, 348 (1900).
 - 80 Burrows, J. Chem. Soc., 105, 1260 (1914).
 - ²¹ Rosanoff and Potter, J. Amer. Chem. Soc., 35, 248 (1913).
- ⁸² Röntomn, Wiedermann's Annalen (3), 45, 98 (1892). Rothmund, Z. physik. Chem., 20, 170 (1896).
 - 38 MEISSL, J. prakt. Chem. (2), 25, 120 (1882).
- ³⁴ MUNTZ, Jahresber., 1873, 829. BERTHELOT, Ann. Chim. Phys. (3), 55, 272 (1859).

phuric acid seems to shorten the time required. There is an intermediate formation of *dextrine* which is, in turn, hydrolyzed by the dilute acid.

327. Glucosides. The various substances designated by this name are numerous among vegetable products and have a constitution analogous to that of the polysaccharides. Their hydrolysis by pure water can usually be accomplished only by heating to high temperatures in sealed tubes, but by boiling with dilute mineral acids, they are decomposed into a sugar (usually glucose) and one or more substances of various kinds.

The action of acids is comparable to that of soluble ferments, such as *emulsine*, but is more rapid and more drastic, the product of hydrolysis being sometimes altered by the peculiar action of the acid.

328. Thus arbutine when boiled with dilute sulphuric acid is hydrolyzed into 1 molecule of glucose and 1 molecule of hydroquinone, 35 which is identical with the results obtained by long contact with emulsine in the cold.

Helicine is hydrolyzed by dilute acids into glucose and salicylic aldehyde, so and quercitrine into isodulcite and quercitine (tetrahydroxyflavanol. so

The ruberythric acid of madder root gives alizarine and 2 molecules of glucose when boiled with dilute acids.38

329. Salicine heated to 80° with 10 parts of fuming hydrochloric acid (d. 1.25), gives 2 molecules of glucose ³⁰ and saliretine, O(C₆H₄.CH₂.OH)₂, while the action of emulsine in the cold or boiling with dilute acid leads to saligenine, o.HO.C₆H₄.CH₂OH.⁴⁰

Amygdaline is decomposed by boiling with dilute hydrochloric or sulphuric acid, just as it is in the cold by emulsine, into benzaldehyde, hydrocyanic acid, and 2 molecules of glucose:

$$C_{20}H_{27}NO_{11} + 2H_2O = C_6H_5 \cdot CHO + HCN + 2C_6H_{12}O_6.$$

But when acids are used the hydrocyanic acid formed is rapidly hydrolyzed into formic acid and ammonia.⁴¹

Coniferine is split by emulsine into glucose and coniferyl alcohol, but when the hydrolysis is carried out by boiling with dilute acids, the alcohol is resinified.⁴²

- 85 KAWALIER, Annalen, 84, 356 (1852).
- ⁸⁶ Piria, Annalen, 56, 64 (1845).
- 87 RIGAUD, Annalen, 90, 289 (1856).
- 88 GRAERE and LIEBERMANN, Annalen, Supl., 7, 296 (1870).
- 89 KRAUT, Annalen, 156, 124 (1870).
- 40 Piria, Annalen, 56, 37 (1845).
- 41 Ludwig, Jahresber., 1855, 699 and 1856, 679, Arch. Pharm. (2), 87, 273.
- 42 TIEMANN and HARMANN, Berichte, 7, 611 (1874).

330. The dilute acids can be replaced by dilute soluble bases such as sodium, potassium and barium hydroxides or even by a solution of zinc chloride (for example with helleborine).48

331. Acidamides and Analogous Compounds. The derivatives formed by the loss of a molecule of water between an organic acid and ammonia, an amine, hydroxylamine, hydrazine, phenylhydrazine or semicarbazid can be more or less readily hydrolyzed into the molecules from which they were derived. This hydrolysis can be accomplished by mineral acids which combine with the ammonia or other base or by aqueous alkalies which unite with the organic acid.

Amides which can be hydrolyzed by boiling with pure water are much more rapidly hydrolyzed by heating with dilute mineral acids or with dilute alkalies.

332. The hydrolysis of oximes takes place on contact with hot concentrated hydrochloric acid, which combines with the hydroxylamine that is liberated along with the aldehyde or ketone.

Phenylhydrazones are hydrolyzed in the cold with concentrated hydrochloric acid which combines with the phenylhydrazine.

by sulphuric acid diluted with 4 molecules of water to give 2 molecules of hydrazine and 2 molecules of oxalic acid.44

333. Heating in a sealed tube with a concentrated solution of hydrochloric acid causes the hydrolysis of sulphocyanic esters:

$$CN.SR + 2H_2O = R.SH + CO_2 + NH_3$$

as well as of mustard oils, or isosulphocyanic esters:

$$CS : NR + 2H_2O = H_2S + CO_2 + H_2NR.$$

334. On the contrary, the hydrolysis of isocyanic esters, or alkyl carbylamines, is carried out by boiling with aqueous 45 or alcoholic 40 potash:

$$CO: NR + H_2O = CO_2 + H_2NR$$

and the activity of the base can be attributed to its affinity for carbonic acid.

- 48 HUSEMANN and MARMÉ, Annalen, 135, 55 (1865).
- 44 CURTIUS and LANG, J. prakt. Chem. (2), 38, 532 (1888).
- 45 WURTZ, Ann. Chim. Phys. (3), 42, 43 (1854).
- 46 HALLER, Bull. Soc. Chim. (2), 45, 706 (1886).

335. The hydrolysis of amides and of alkyl amides can be carried out by acids or alkalies indifferently. In the case of aliphatic amides, the reaction is usually effected by heating with alcoholic potash or soda and takes place slowly, sometimes requiring heating for several days. We have:

$$R \cdot R \cdot CO \cdot NH_2 + H_2O - R \cdot CO_2H + NH_2$$

and we may believe that the affinity of the potash for the acid determines the reaction.

336. In most of the above reactions the catalytic rôle of the acids and bases does not appear, at first sight, to be well established. We can, however, assume that it is really catalytic, since in the reactions that have been most closely studied, such as the hydrolysis of amides, amounts of acid far less than equivalent to the amides greatly accelerate the reaction. In the case of the hydrolysis of acetamide by dilute mineral acids, it has been found that the activity of the acids is proportional to their ionization and to the concentration of the ions in the system.⁴⁷

§ 2. — HYDROLYSIS IN GASEOUS SYSTEMS

337. The Saponification of Esters. Titania, TiO₂, which readily causes the esterification of alcohols by aliphatic acids (767), between 280 and 300°, just as readily reverses the reaction and brings about the saponification of esters by water. The rapid passage of a mixture of water and ester vapors in equivalent amounts over the oxide at 280 to 300° is sufficient to reach about 30% hydrolysis and this percentage is increased as the relative amount of water is increased till practically complete hydrolysis is effected by very large amounts of water.

Thoria can produce the same effect but with less activity.48

338. Ethers. Thoria, ThO₂, which effectively catalyzes the formation of phenyl oxide, (C₆H₅)₂O, from phenol at 400 to 500°, can equally well decompose it when phenyl ether and water vapor are passed over the heated oxide, the decomposition reaching 50%.⁴⁹

339. Hydrolysis of Carbon Disulphide. The reaction of water vapor on carbon disulphide in the presence of appropriate catalysts, such as ferric oxide, can be considered a case of hydrolysis. The reaction is incomplete but goes in this direction:

$$CS_2 + 2H_2O = CO_2 + 2H_2S$$
.

- ⁴⁷ OSTWALD, J. prakt. Chem. (2), 27, 1 (1883).
- 48 SABATTER and MAILHE, Compt. rend., 152, 494 (1911).
- 49 SABATTER and ESPIL, Bull. Soc. Chim. (4), 15, 228 (1914).

This reaction applied to illuminating gas suppresses 67% of the carbon disulphide which it contains and, if the hydrogen sulphide is absorbed as rapidly as it is formed, all of the carbon disulphide is eliminated.⁵⁰

III. — ALCOHOLYSIS

340. The action of alcohols on esters can be compared to the saponification of esters by water and is likewise catalyzed by small quantities of strong mineral acids, hydrochloric and sulphuric.⁵¹

If a primary aliphatic alcohol, R'OH, is mixed with the ester derived from an acid, RCOOH and a complex alcohol, MOH, we shall have: 52

$$R.CO.OM + R'.OH = MOH + R.CO.OR'.$$

The alcohol, MOH, is set free. This is what takes place when methyl, ethyl, propyl alcohols and the like attack the esters of borneol, glycerine, etc., in the presence of a minute amount of hydrochloric acid.

Thus bornyl acetate in a methyl alcohol solution, containing 1% hydrochloric acid, is rapidly decomposed into borneol and methyl acetate.

Glycerides dissolved in absolute alcohol containing a few per cent of hydrogen chloride yield glycerine and the fatty acid ethyl esters. 52

341. Haller has designated these saponifications which take place readily with all the fats, by the name of alcoholysis. They can be carried out by mixing 100 g. of a fat with 200 g. dry methyl alcohol containing 1 or 2 g. hydrogen chloride and heating on a steam bath under reflux till the mixture becomes homogeneous. If necessary

- 50 GRILLET, Soc. Tech. de l'Ind. du gaz en France, 1918, 245.
- ⁵¹ Sodium alcoholate is an even better catalyst than hydrochloric acid. In the transformation of methyl benzoate into the ethyl ester, sodium ethylate was found to be about 4,000 times as efficient as an equivalent amount of hydrochloric acid. (Red, Amer. Chem. J., 45, 506 (1911)).—E. E. R.
- ⁵² This reaction is a perfectly general one and simple alcohols may be replaced as well as "complex," thus methyl alcohol replaces ethyl and vice versa as shown by Red (Amer. Chem. J., 45, 479 (1911)), and recently by Redmer and Downes (J. Amer. Chem. Soc., 43, 945 (1921)).

For a number of references on alcoholysis see article by Parden and Rum (J. Ind. and Eng. Chem., 12, 129 (1920)). It is a reversible reaction, the equilibrium point depending on the concentrations and activities of the two alcohols competing for the acid and hence can never be complete, no matter how much one alcohol predominates.—E. E. R.

58 ROCHLEDER, Annalen, 59, 260 (1846). BERTHELOT, Ann. Chim. Phys. (3), 41, 311 (1854).

more hydrogen chloride may be added during the reaction. The mixture is finally poured into brine which dissolves the glycerine and causes the methyl esters of the fatty acids to separate as a top layer.⁵⁴

This reaction is rapid with cocoa butter ⁵⁵ and castor oil, with which heating for several hours is sufficient, ⁵⁶ and is slower with drying oils such as linseed. ⁵⁷ It goes just as well with ethyl, propyl, and isobutyl alcohols. ⁵⁶

- 54 HALLER, Compt. rend., 143, 657 (1906).
- 55 HALLER and Youssouffian, Compt. rend., 143, 803 (1906).
- 56 HALLER, Compt. rend., 144, 462 (1907).
- ⁵⁷ HALLER, Compt. rend., 146, 259 (1908).

CHAPTER VIII

HYDROGENATIONS

HYDROGENATIONS IN GASEOUS SYSTEM, GENERALITIES. USE OF NICKEL

342. Historical. The catalytic properties of finely divided platinum discovered by Davy and Doebereiner at the beginning of the nineteenth century, have shown its power to cause oxidations. Several chemists attempted to apply the special powers of platinum sponge to other reactions and particularly to the direct addition of hydrogen to various substances. In 1838, Kuhlmann showed that nitric oxide, or the vapors of nitric acid, warmed with hydrogen in the presence of platinum sponge gave ammonia. In 1852, Corenwinder observed that the same agent caused hydrogen to combine rapidly, though incompletely, with iodine between 300 and 400°.2° In 1863, Debus, with the aid of platinum black, accomplished the addition of hydrogen to hydrocvanic acid to form methyl amine.3 and found that ethyl nitrite is transformed into alcohol and ammonia under the same circumstances. In 1874, von Wilde succeeded in transforming acetylene into ethylene and then into ethane, by platinum black at room temperature.4

343. In a series of investigations continued since 1897, Sabatier and Senderens (1897–1905), then Sabatier and Mailhe (1904–1908), and Sabatier and Murat (1912–1914) have established and extended to a large number of cases a general method of direct hydrogenation of volatile organic compounds, based on the use of finely divided catalytic metals and particularly on the use of nickel recently reduced from the oxide.⁵

- ¹ KUHLMANN, Compt. rend., 7, 1107 (1838).
- ² CORENWINDER, Ann. Chim. Phys. (3), 34, 77 (1852).
- ³ Debus, Annalen, 128, 200 (1863).
- 4 VON WILDE, Berichte, 7, 352, (1874).
- ⁵ These investigations have been published in a large number of original articles of which more than 50 are in the Comptes Rendus de l'Academie des Sciences as well as in various collective memoirs of which the chief are: Sabatier, Vth Congress of Pure and Applied Chemistry, Berlin, 1904, IV, 663. Sabatier and Senderens, Confer. Soc. Chim., Paris, 1905. Sabatier, Rev. Gen. Sc., 16, 842 (1905). Sabatier, Rev. Gle. Chim., 8, 381 (1905). Sabatier and Senderens,

As early as 1902 this new method was taken up in many French and foreign laboratories and numerous chemists have contributed, along with the above authors, to widen its application.

344. Essentially the process consists in passing the vapors of the substance mixed with hydrogen over a layer of the catalytic metal, platinum black, or even nickel, cobalt, iron, or copper reduced from the oxides in the same tube in which the hydrogenation is to be carried on, maintained at a suitable temperature, sometimes room temperature but more commonly somewhere between 150 and 200°. A temperature around 180° is very frequently found to be the most suitable.

Of the five metals mentioned above, nickel is the most active and it and cobalt are the only ones capable of effecting certain hydrogenations such as that of the benzene nucleus. Copper is less powerful and platinum and iron are between cobalt and copper.

345. The apparatus employed by Sabatier and his co-workers comprises:

- 1. A hydrogen generator.
- 2. A working tube to contain the catalytic metal.
- 3. An arrangement for introducing the vapors to be hydrogenated along with the hydrogen.
 - 4. A receiver to collect the product of the reaction.
- 346. The Hydrogen Generator. The hydrogen can be prepared by the action of commercial hydrochloric acid diluted with half its volume of water on ordinary granulated zinc. The continuous generator of Sainte-Clair Deville consists of two large flasks of 10 to 15 l. of which the lower tubulures are connected by large rubber tubing. One flask is filled with granulated zinc and the other with hydrochloric acid. The gas is washed with strong caustic soda and then with concentrated sulphuric acid. A graduated safety tube in the acid wash bottle serves to indicate the gas pressure. Between the two wash bottles is a stop cock to regulate the gas and beyond the acid wash bottle is a pinch cock for further adjustment of the pressure. To secure a regular delivery of the gas it is sufficient to maintain the acid in the safety tube at a constant height. On account of

Ann. Chim. Phys. (8), 4, 319 (1905). SABATIER, VIth. Congress Pure and Appl. Chem., Rome, 1906, Xth. Sect. 174. SABATIER and MAILHE, Ann. Chim. Phys. (8), 16, 70 (1909). SABATIER, Berichte, 44, 1984 (1911). SABATIER, Address at Stockholm on the reception of the Nobel Prize, Rev. Scient., 1, 289 (1913). SABATIER, Confer. à Toulouse au Congrès du gaz, Le Gaz, 57, 1914. SABATIER, Confér. à University of London, Rev. Gle. Chim., 17, 185 and 221 (1914). SABATIER and MURAT, Ann. de Chim. (9), 4, 253 (1915).

the large dimensions of the apparatus, a constant evolution of gas can be maintained for at least six hours.

The hydrogen must be carefully freed from impurities derived from the zinc or from the acid (hydrogen sulphide, arsine, phosphine and hydrochloric acid vapors). For this purpose it passes through a tube of Jean glass, filled with copper turnings kept at a dull red, which stops the major part of the impurities. The purification is completed by passing the gas through a long tube filled with slightly moist fragments of caustic potash which retains acid vapors as well as any remaining hydrogen sulphide. The purified gas passes to the reaction tube.

The complete drying of the gas appears superfluous as it has been shown that moist hydrogen hydrogenates benzene or phenol, over nickel, at least as well as dry.

Electrolytic hydrogen, which is on the market in steel cylinders at high pressures, can be used to advantage. These cylinders fitted with suitable reducing valves, furnish a nearly pure gas which can be freed from the small amount of oxygen which it contains by passing over red hot copper in a tube followed by a drying tube containing caustic potash.

347. The Reaction Tube. In a glass tube 65 to 100 cm. long and 14 to 18 mm. inside diameter, a longer or shorter (35 to 80 cm.) thin layer of platinum black or of the oxide, from which the catalytic metal is to be prepared, is spread. The tube is heated in a gas furnace such as is used for organic combustions but in which the burners have wing tips with little holes so that there are a large number of little flames equal in size and close together distributing the heat evenly.

The tube is laid in a semicircular trough and rests on a rather thick layer of calcined magnesia or fine sand. The temperature is taken simply with a glass thermometer graduated to 450° which is embedded in the trough by the side of the tube and which may be moved from place to place to test the evenness of the heating.

The temperature read on the thermometer is always a little lower than that in the tube, the difference being greater at higher temperatures. For temperatures around 180 to 200° the difference is hardly more than 10 to 15°, while at 350° it may be as great as 35°. The limits between which the reactions go on are usually wide enough so that this approximate determination of the temperature is sufficient.

⁶ SABATIER and ESPIL, Bull. Soc. Chim. (4), 15, 228 (1914).

⁷ SABATTER and MAILHE, Ann. Chim. Phys. (8), 20, 296 (1910).

348. If more exact determinations are desired a rectangular copper oven 12 x 15 x 65 cm. down the centre of which runs a copper tube is used. The thermometer and the tube containing the previously prepared catalysts are placed in this tube side by side. A metallic regulator contained in a copper tube parallel to the first controls the gas and maintains the temperature at which it is set. The copper box is filled with a liquid which up to 270° may be boiled linseed oil, or for higher temperatures a mixture of equal weights of sodium and potassium nitrates which is liquid above 225°. For delicate hydrogenations with such substances as benzoic esters, Sabatier and Murat have employed a massive bronze block, 65 cm. long, 10 cm. wide and 7 cm. high, of rectangular cross section, with rounded corners. Two symmetrically placed holes 25 mm. in diameter run from one end to the other of the block: the one contains the tube carrying the nickel and the other the metallic regulator which controls the gas supply of the furnace. Any desired temperature is thus obtained very uniformly on account of the large mass of the good conducting metal. On account of this conductivity, the temperature may be raised quickly. Small holes parallel to the large ones receive the thermometers.

The temperature may be first carried to 350° for the preparation of the nickel and then lowered to any desired temperature, such as 180°, for carrying out the hydrogenation.

In case nickel-coated pumice is used as catalyst (126) a very useful arrangement is to fill the two limbs of a vertical U-tube with the catalyst. This tube may be heated in an air bath to 350° for reducing the nickel and then lowered into an oil bath kept at 180° or into the vapors of boiling aniline, 185°, for the hydrogenation.

Heating on the furnace is less regular and requires close attention but has the advantage that the interior of the tube may be watched.

349. Heating by electric resistance may be conveniently employed. The reaction tube is surrounded by asbestos paper on which is wound a 1 mm. ferro-nickel spiral which, in turn, is surrounded by a second layer of asbestos paper. By the aid of suitable resistances the current is regulated to show the proper readings on an ammeter. The temperatures in the centre of the tube corresponding to various ammeter readings are previously determined by experiment.⁸

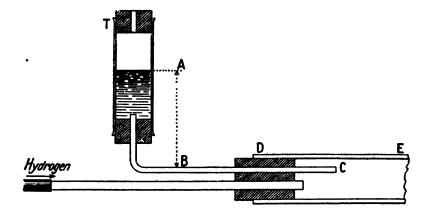
This method of heating has the advantage, as compared to the open furnace, that the tube is heated uniformly around its whole

⁸ The conditions of the experiment must be exactly duplicated during the calibration since otherwise incorrect estimates of temperatures are possible as varying amounts of heat are removed by varying currents of gas through the tube.— E. E. R.

circumference, and, with it it is best to employ nickeled pumice filling the whole tube rather than a layer of nickel resting in the bottom.

350. Introduction of the Substance. The method of introducing the substance to be hydrogenated varies, of course, according to its physical state.

If it is a gas the forward end of the tube containing the catalyst carries a two-hole stopper with two tubes, one for the gas and one for the hydrogen. The gas is furnished by a continuous generator (as with acetylene or carbon dioxide) or by a metal or glass gasometer into which it is measured in advance (carbon monoxide, propylene, nitrous oxide), or even by a discontinuous apparatus which can be operated sufficiently regularly (as for ethylene, or nitric oxide). A wash cylinder with pressure indicator interposed between a stop cock and a screw pinch cock as has been described above for hydrogen (346), serves to admit the gas at any desired constant rate. In the case of discontinuous generators, a safety valve is arranged by having a side outlet tube dipping under mercury so that the excess of gas may escape.



351. For most *liquids*, Sabatier and Senderens have devised an extremely simple apparatus. The liquid is conducted by a capillary tube to the interior of the reaction tube. The liquid is placed in a large vertical tube T, the lower end of which carries a stopper through which passes the vertical portion of a bent capillary tube, the horizontal portion of which passes through the stopper in the end of the reaction tube.

For a given liquid, the flow is more rapid the larger the bore of the

⁹ BRUNEL, Ann. Chim. Phys. (8), 6, 205 (1905).

capillary tube and the greater the head of liquid, AB. By maintaining this head constant, a regular flow of liquid is obtained.

It is well to arrange it so that the liquid does not fall from the end of the capillary tube in drops, but flows steadily from its end either on to the wall of the reaction tube or over the surface of the cork in its end.

The selection of the capillary tube depends on the viscosity of the liquid, a smaller tube being used for mobile liquids.

It is evident that there are two independent ways of regulating the flow of the liquid, by changing the diameter of the capillary tube or altering its height. Besides, the capillary tube can be fed by a reservoir with as large a surface as may be desired and, for experiments of long duration, the tube A can be placed in communication with a flask of large size in which the variations of level are very slow.

It is convenient for the stopper D to be at some distance from the heated portion of the tube; 3 to 4 cm. is sufficient. The layer of metal should not commence for a little distance, about 10 cm. from the stopper. The liquid introduced by the capillary volatilizes regularly in this open space. It is important to watch that the liquid does not wet the catalyst which is frequently altered by contact with the liquid.

352. We may also operate by bubbling the hydrogen through the liquid to be hydrogenated, thus carrying along the vapors. If the liquid is very volatile (acetaldehyde, propionic aldehyde, nitrogen peroxide, etc.) cooling is necessary so that the amount of the vapors carried along will not be too great.

If the liquid is only slightly volatile, heating may be required, always selecting a temperature so that the hydrogen will be in excess of that required for complete hydrogenation.⁹⁶

353. For solid substances which melt below 100°, the same appa-

he liquid through which the hydrogen is bubbled must be kept at such a temperature that its vapor pressure is 380 mm. For some liquids this temperature may be found from tables in the literature. The vapor pressure curves for various classes of liquids are not quite similar, owing to different degrees of association, but for most organic liquids, except the lower alcohols, the vapor pressure is 380 mm. at from 20 to 24° below their boiling points. To have a little more than 1 molecule of hydrogen to 1 of the vapor the liquid should be kept at from 25 to 30° below its boiling point. These same liquids have vapor pressures approximately one third of an atmosphere at 32 to 36° below their boiling points and should be kept at such temperatures to obtain 2 molecules of hydrogen to 1 of the vapor or at somewhat lower temperatures if an excess of hydrogen is desired, as is usually the case. Similar calculations may be made when a larger number of molecules of hydrogen to one of the compound are desired.—E. E. R.

ratus may be used by surrounding the capillary tube and the vertical tube T with a sort of cylindrical air bath, the lower end of which is heated by a Bunsen burner. The current of warm air is sufficient to maintain the substance in the liquid condition. This method may be used with phenol, the cresoles, the nitronaphthalines and naphthaline

A thick copper capillary tube brazed on to a copper vertical tube T may be used, and this may be heated directly by a small flame.

When the substance melts above 100°, it is placed in long porcelain boats in the forward part of the tube, a long tube being selected. The volatilization of the substance is effected by careful heating, a portion at a time, starting from the end next to the heated metal. The reaction is of course limited to the amount of material in the boats and is consequently intermittent.

Solids melting below 180° may be kept fused by a suitable air bath and the vapors carried on by the hydrogen which is bubbled through.

354. When the product of the hydrogenation is a liquid, it is frequently sufficient to mix some of it with the solid to be hydrogenated, thus lowering the melting point so that the usual apparatus for liquids may be employed. This is the case with phenol and with ortho and meta cresoles.

The use of solvents which can not be hydrogenated, such as water, paraffine hydrocarbons (hexane, heptane, etc.) usually gives poor results, particularly when water is used.

355. Apparatus for Collecting the Reaction Products. If the products of the hydrogenation are all gases, they are collected at the end of the catalyst tube in a gas holder over water, care being taken to saturate the water with common salt to diminish the solubility of the gases. It is well to time the collection of issuing gas in a graduated tube. A comparison of the rate at which the gases come out with the rate at which they are passed in, frequently gives valuable information as to the exact course of the reaction.

356. If the products are partly or entirely liquid, the reaction tube is connected with a condenser. When the substances are only slightly volatile this may be simply a double-necked flask. When the volatility, at room temperature, is considerable, a U-tube is employed from the bottom of which a tube leads down into a flask in which the liquid collects. The U-tube is placed in an inverted tubulated bell-jar which is filled with cold water, ice, or a freezing mixture. The gas issuing from the other limb of the U-tube is collected over water and measured.

357. Solid reaction products are collected by prolonging the re-

action tube and cooling the further end. The tube should be long enough to project a considerable distance from the furnace and the end should be inclined downward so that condensed liquids will not run back towards the catalyst.

HYDROGENATIONS BY MEANS OF NICKEL

358. In Chapter II the conditions have been described under which nickel may be used to advantage as a catalyst for hydrogenations (53), and methods have been given for obtaining a metal of excellent catalytic properties. Nickel reduced at a red heat below 700° is capable of effecting all sorts of hydrogenations and in particular can hydrogenate benzene to cyclohexane; 10° but that reduced above 750°, or which has been heated to that temperature after having been reduced at a lower, is incapable of hydrogenating benzene, is no longer pyrophoric and does not gain in weight when exposed to cold air. It is then capable of only certain hydrogenations, such as the reduction of nitro derivatives.

359. As has been stated above (112), the presence of chlorine, bromine, or iodine, even in traces, in the metal paralyzes its catalytic activity. An oxide prepared by precipitation from the chloride can not be used, but good results can be obtained with an oxide produced by calcining the sulphate at a red heat.

Whatever care one may take, it is never possible to avoid all the causes of poisoning the metal catalyst and particularly in consequence of the progressive fouling of the metal which is more or less rapid according to the work done with it, a gradual diminution of the catalytic power, its *senilescence*, so to speak, is noticed.

360. Darzens believes that nickel exists in three forms, α , β , and γ . The very active γ form is said to be obtained by reduction below 260° and is considered unstable, remaining in metastable state below 260°. Above that temperature it passes into the less active β nickel, then at a bright red into the α form which is entirely inactive for hydrogenations. According to this author the power to hydrogenate benzene belongs exclusively to γ nickel, which is contrary to the observations of Sabatier and Espil quoted above. These transformations of γ nickel, rapid at high temperatures, would take place slowly even at low temperatures and would explain the senilescence of the metal apart from many poisoning effects. 12

¹⁰ SABATIER and ESPIL, Bull. Soc. Chim. (4), 15, 779 (1914).

¹¹ DARZENS, Bull. Soc. Chim. (4), 15, 771 (1914).

¹² Darzens, Compt. rend., 139, 869 (1904).

361. Choice of Reaction Temperature. A given hydrogenation can be realized only within a well-defined temperature interval.

In practice, a lower temperature limit is set by the necessity of maintaining in the vapor state in the reaction tube, not only the compounds to be transformed but also the products of the reaction.

To a certain extent elevation of temperature accelerates the reaction and consequently raises the proportion of the substance hydrogenated during its passage through the tube. But beyond a certain limit, sometimes not far above the temperature at which the reaction begins, there is a profound modification of the phenomenon, it being possible to completely reverse the reaction in some cases. Thus the hydrogenation of benzene may be accomplished as low as 70°, and it increases in velocity as the temperature is raised till a maximum is reached at 180–200°. Then it decreases till 300° is reached, at which benzene is no longer hydrogenated, but, on the contrary, cyclohexane is decomposed into benzene and hydrogen.

362. By hydrogenating around 300°, the aromatic nucleus remains almost unaffected while any unsaturated side-chains are hydrogenated. Thus styrene, C₆H₅CH: CH₂, hydrogenates almost completely at 300° to ethyl-benzene, C₆H₅.CH₂.CH₃, while if the temperature be reduced to 180°, this is further changed into ethyl-cyclohexane, C₆H₁₁.CH₂.CH₃.

If the temperature is raised above 300°, the aromatic nucleus is, little by little, broken up, and particularly in the case of benzene the reaction:

$$C_6H_6 + 9H_2 = 6CH_4$$
methane

tends to become more and more important.14

363. When a compound can add several molecules of hydrogen in succession, we can sometimes contrive, by suitably choosing the temperatures, to produce one after the other of the various combinations. In the hydrogenation of anthracene over nickel, at 180°, perhydro-anthracene, C₁₄H₂₄, is obtained along with the dodecahydro-, at 200°, the octohydro-, and at 260°, the tetahydro-anthracene. 16

364. The easy hydrogenations are those which take place over a wide range of temperatures, as the saturation of ethylene bonds or the reduction of nitro compounds. The more difficult cases are those

¹⁸ SABATHER and MURAT, Ann. de Chim. (9), 4, 255 (1915).

¹⁴ SABATTER and SENDERENS, Ann. Chim. Phys. (8), 4, 334 (1905).

¹⁸ SABATTER and MAILHE, Compt. rend., 137, 240 (1903).

¹⁶ GODCHOT, Ann. Chim. Phys. (8), 12, 468 (1907).

where the possible temperature interval is narrow, as is the case in the hydrogenation of the aromatic nucleus, especially with diphenols, pyrogallol,¹⁷ benzoic esters, and quinoline.¹⁸

365. As has been stated above (167), the hydrogenating activity of nickel is attributed to the rapid formation of a hydride formed directly by the hydrogen gas on the surface of the metal. This hydride is readily dissociated, and if it is brought into contact with substances capable of taking up hydrogen, it gives it to them very rapidly, regenerating the metal which can again form the hydride, repeating these reactions indefinitely.

The well-attested impossibility of carrying on all sorts of hydrogenations with any sort of nickel leads to the idea that there are several stages of combination with hydrogen. The nickel produced above 700° can doubtless form only the first hydride, comparable to that formed by copper, and capable of reacting with nitro groups or with an ethylene hydrocarbon. Only powerful nickel, such as is furnished by the reduction at a low temperature of the oxide prepared from the nitrate, can form a perhydride capable of hydrogenerating the aromatic nucleus (167).

RESULTS OBTAINED BY HYDROGENATION OVER NICKEL IN GASEOUS SYSTEM

- 366. The results obtained by hydrogenation over reduced nickel can be divided into four groups:
 - 1. Simple reductions without fixation of hydrogen,
 - 2. Reductions with simultaneous fixation of hydrogen,
- 3. Addition of hydrogen to molecules which contain multiple bonds between various atoms,
- 4. Hydrogenations accompanied by decomposition of the molecule.

REDUCTIONS EFFECTED WITHOUT FIXATION OF HYDROGEN

- 367. The reduction effected by the aid of nickel corresponds most frequently to the elimination of oxygen in the form of water; it can also remove sulphur as hydrogen sulphide.
- 368. Nitrous Oxide. The first case is furnished by nitrous oxide which is reduced to nitrogen, even at the ordinary temperature, with-
 - 17 SABATIER, Berichte, 44, 1997 (1911).
 - 18 SABATIER and MURAT, Compt. rend., 158, 309 (1914).

out any production of ammonia or hydrazine. By increasing the proportion of nitrous oxide in the hydrogen, the heat evolved raises the first portions of the nickel to incandescence, and there results a partial decomposition of the nitrous oxide with the appearance of red nitrogen peroxide, the hydrogenation of which carried on by the neighboring hot nickel gives a little ammonia.¹⁹

369. Aromatic Alcohols. The hydrogenation of aromatic alcohols over nickel at 350-400° replaces the hydroxyl group by hydrogen and leads to the corresponding aromatic hydrocarbon.²⁰

Benzyl alcohol is changed to toluene, phenylethyl alcohol to ethylbenzene, benzhydrol, C₆H₈.CH(OH).C₆H₈, is changed quantitatively into diphenyl-methane, C₆H₈.CH₂.C₆H₈, and phenyl-p.cresyl carbinol, into phenyl-p.cresyl-methane.

Likewise, vapors of triphenyl carbinol, carried along by benzene vapors and hydrogen over nickel at 400°, readily yield triphenylmethane.

This reaction is particularly easy when the alcoholic hydroxyl is attached to a carbon atom adjoining a carbon atom united to hydrogen in the same paraffine side-chain. The mechanism of the reaction may then correspond to a dehydration into the phenyl-ethylene hydrocarbon, which is at once hydrogenated into the saturated hydrocarbon. Thus tolyl-dimethyl carbinol, CH₃.C₆H₄.C(OH).(CH₃)₂, which is very readily dehydrated, gives with a nickel only slightly active cymene, which may be transformed into menthane if an active nickel is used below 180°.²¹

370. Phenols and Polyphenols above 250°. Phenol hydrogenated at 250 to 300° over nickel, gives only benzene with the elimination of water:

$$C_6H_6.OH + H_2 - H_2O + C_6H_6.$$

But the reaction is slow and much of the phenol passes by unchanged. If the attempt is made to hasten the reaction by raising the temperature, the benzene is attacked with the formation of methane. The three *cresoles* behave the same way and yield toluene.

At 250° the *diphenols* (pyrocatechin, resorcine, and hydroquinone) undergo a similar reaction, the hydroxyl groups being successively replaced by hydrogen, phenol being first formed and then benzene.²²

¹⁹ SABATTER and SENDERENS, Compt. rend., 135, 278 (1902).

²⁰ SABATIER and MURAT, Ann. de Chim. (9), 4, 258 (1915).

²¹ Smirnof, J. Russian Phys. Chem. Soc., 41, 1374 (1909).

²² SABATUER and Sundmenns, Ann. Chim. Phys. (8), 4, 429 (1905).

371. Furfuryl Alcohol. This alcohol submitted to a careful hydrogenation over nickel at 190°, yields methylfurfurane.²⁸

372. Carbon Disulphide. Carbon disulphide submitted to hydrogenation over nickel below 200°, gives an addition product having a very disagreeable odor (492), but if the operation is carried on at 450-500°, in excess of hydrogen, the reaction takes place thus:

$$CS_2 + 2H_2 - 2H_2S + C$$
.

This reaction is utilized in freeing coal gas from carbon disulphide which it contains up to 0.02%.

The gas is freed from hydrogen sulphide by chemical purification in the Laming absorbers and is then heated to 400° and passed through steel tubes 7 cm. in diameter containing porous earth impregnated with nickel and heated to between 400 and 500°. The gas is cooled when it passes out of the tubes and is freed by a second passage through the chemical absorbers from the hydrogen sulphide which has been formed. On account of the deposition of carbon and also on account of a certain sulphurization of the surface, the nickel loses its activity rather rapidly. It is regenerated by passing air which burns up the carbon and converts the nickel to the oxide which is again reduced by the first portions of gas that enter. The installation of this process at the Greenwich gas works is capable of handling 500,000 cu. m. per day.²⁴

REDUCTIONS WITH SIMULTANEOUS FIXATION OF HYDROGEN

- 373. These reductions can be considered as true substitutions of hydrogen either for oxygen or, in a few cases, for the halogens, chlorine or bromine.
- 374. Oxides of Nitrogen. Although the oxides of nitrogen are outside of the scope of this treatise, yet their close connection with organic nitro and nitroso compounds justifies us in mentioning the conditions of their catalytic hydrogenation.
 - ²⁸ Padoa and Ponti, Lincei, 15 (2), 610 (1909); C., 1907 (1), 570.
- ²⁴ CARPENTER, J. Gas Lighting, 126, 928 (1914). Evans, J. Soc. Chem. Ind., 34, 9 (1915).

Nitric oxide, NO, is readily reduced above 180° with the formation of ammonia and water according to the equation:

$$NO + 5H = NH_s + H_sO$$
.

But the nitric oxide reacts with the ammonia more and more rapidly the higher the temperature, giving nitrogen and water according to the equation:

$$2NH_{3} + 3NO = 5N + 3H_{2}O.$$

By progressively increasing the proportion of nitric oxide, the metal becomes incandescent and this greatly increases the production of nitrogen.²⁵

375. If hydrogen which has passed through a thin layer of liquid nitrogen peroxide, cooled a little below 0°, is passed over cold reduced nickel, a slight evolution of heat is noticed which is due to the formation of nickel nitride.²⁶

If it is heated to 180°, white fumes of ammonium nitrate and nitrite appear which, when hydrogenated further, give ammonia and water. We have finally:

$$NO_2 + 7H = NH_3 + 2H_2O.$$

If the proportion of nitrogen peroxide in the hydrogen is increased by warming the vessel containing the nitrogen peroxide, the white fumes are produced in abundance and incandescence of the nearest portion of the metal layer is noticed and a violent explosion soon takes place.²⁵

376. The vapors of *nitric acid* mixed with hydrogen and passed over nickel at 290° give much ammonium nitrate. At 350° only water, ammonia, and free nitrogen are produced.²⁵

377. Aliphatic Nitro Compounds. Nitromethane is completely hydrogenated between 150 and 180° to methyl-amine without any side reactions. But above 200° and particularly towards 300°, there is partial hydrogenation of the methyl-amine into methane and ammonia:²⁷

$$CH_3 \cdot NO_2 + 4H_2 = CH_4 + NH_3 + 2H_2O$$

and at the same time, the formation of certain amounts of dimethyland trimethyl-amines along with the ammonia by a reaction identical with that which has been described in the hydrogenation of nitriles.

²⁵ SABATIER and SENDERENS, Compt. rend., 135, 278 (1902).

²⁶ SABATIER and SENDERENS, Ann. Chim. Phys. (7), 7, 413 (1895).

²⁷ SABATTER and SENDERENS, Compt. rend., 135, 226 (1902).

Likewise nitroethane is readily transformed at 200° into ethylamine accompanied by diethylamine, triethylamine and ammonia. At 350° the matter is complicated by the formation of ethane and also of methane which is due to the dissociation of the ethane by the nickel. But this secondary formation of the hydrocarbon is less than with nitromethane.

378. Aromatic Nitro Compounds. Above 200° nitrobenzene is rapidly transformed into aniline, but the aniline is immediately hydrogenated to form cyclohexylamine, etc. (466). If only slightly active nickel is used, the nucleus is not hydrogenated and aniline is the only product.²⁸

Above 250°, a part of the nitrobenzene is reduced to benzene and ammonia:

$$C_0H_5$$
. $NO_2 + 4H_2 = C_0H_0 + NH_3 + 2H_2O$.

This reaction is more in evidence above 300° and even the benzene is broken up to form methane:

$$C_4H_5$$
. NO, + 13H₂ = 6CH₄ + NH₅ + 2H₂O.

Ortho and meta nitrotoluenes behave similarly with a nickel catalyst at 200 to 250°, and as the further hydrogenation of the resulting toluidines does not take place readily, the toluidines are obtained nearly pure.

These reactions can be used in the factory and it has been proposed to prepare aniline by passing a current of hydrogen and steam through nitrobenzene maintained at 120° and then into a long tube containing reduced nickel also kept at 120°. A theoretical yield is claimed.²⁹

379. α -Nitronaphthalene gives at 300° beautiful white needles of α -naphthyl amine, but if the temperature is raised to 330°, or better, to 380°, ammonia is evolved and there condense, along with the diminishing naphthyl amine, naphthalene and tetrahydronaphthalene.²⁸ We have:

$$C_{10}H_7 \cdot NO_2 + 4H_2 = C_{10}H_8 + NH_2 + 2H_2O.$$

380. Dinitro derivatives are transformed with the same facility. The dinitrobenzenes give the corresponding diamines at 190-210°. At 250°, there is the splitting off of ammonia to form aniline.³⁰ Like-

- 28 SABATIER and SENDERENS, Compt. rend., 135, 226 (1902).
- 29 FARBW. MEISTER, LUCIUS and BRÜNING. German patent, 282,492 (1913).
- ⁸⁰ Mignonac, Bull. Soc. Chim. (4), 7, 154 (1910).

wise the dinitrotoluenes yield the cresyl-diamines at 175-180°, but above 190° ammonia is split off and the toluidines are the chief products.^{\$1}

381. The nitrophenols hydrogenated over nickel at 160–190°, yield the amino-phenols regularly; but there are simultaneously produced certain amounts of ammonia and phenol and also a little aniline.³²

382. Esters of Nitrous Acid. It is stated in all the textbooks that a fundamental distinction between the nitrohydrocarbons and their isomers, the nitrites, is that the nitro compounds yield amines on hydrogenation, while the nitrous esters are either not affected or give the alcohols and ammonia without any amine.

Gaudion has found that nitrous esters are regularly hydrogenated by nickel to give amines exactly like their isomers. This author has worked at 180° with methyl and ethyl nitrites, at 200° with propyl and isopropyl, and at 220° with isobutyl and isoamyl.

As a consequence of the secondary reaction already mentioned, all three amines, primary, secondary, and tertiary, are obtained, the secondary always in the largest quantity. Thus from isoamyl nitrite, 31% mono-, 62% di-, and 7% tri-isoamyl-amines are obtained.

The discussion of these facts has led Gaudion to assume that there is an isomerization of the nitrous esters into the nitro bodies at the temperature of the reaction.³⁸

The reality of this transformation by heat alone has since been established. It begins at 100° and is rapid at 125-130°.34

By carrying on the hydrogenation at low temperatures, around 125–130°, over nickeled asbestos, the unchanged nitrous esters are hydrogenated along with the nitro bodies into which they are partly isomerized so that there is simultaneous production of ammonia and the corresponding alcohol from the nitrite and of the amine from the nitro compound; while when a nitro compound is hydrogenated, the primary amine alone is formed without any secondary or tertiary. This is the case with methyl, ethyl, propyl, isobutyl and isoamyl nitrites.³⁵

383. Oximes. In the aliphatic series, aldoximes and ketoximes are readily reduced by hydrogen in the presence of nickel at 180-220° to give primary and secondary amines with a small amount of tertiary.

With acetoxime, the chief product is diethyl amine, while with

- 81 MIGNONAC, Bull. Soc. Chim. (4), 7, 823 (1910).
- 82 MIGNONAC, Bull. Soc. Chim. (4), 7, 270 (1910).
- ⁸⁸ GAUDION, Ann. Chim. Phys. (8), 25, 129 (1912).
- ⁸⁴ NEOGI and CHOWDURG, J. Chem. Soc., 109, 701 (1916).
- 35 NEOGI and CHOWDURG, J. Chem. Soc., 111, 899 (1917).

heptaldoxime, C_eH₁₈. CH: N.OH, the primary amine is the most abundant.

The oxime of acetone gives isopropyl-amine, with twice as much of the di- and a little of the tri-isopropyl-amine. Analogous results are obtained with butanoxime(2), C₂H₅(CH₅)C: N.OH, pentanoxime(2), pentanoxime(3), and 2,4-dimethyl pentanoxime(3).

By this means the secondary amines from secondary alcohols can be prepared with good yields, a class of substances otherwise difficult to obtain.

384. This method can also be applied to aromatic aldoximes in spite of the difficulty of vaporizing them without decomposition. It is best to operate with a rapid current of hydrogen and at as low a temperature as possible. Acetophenone-oxime, C₆H₅.C(:NOH).-CH₃, carried thus over nickel at 250–270°, gives a small amount of the primary amine, C₆H₅.CH(NH₂).CH₃, a larger amount of the secondary amine and some acetophenone regenerated by the action of the resulting water on the oxime.

The results are not so good with *propiophenone-oxime*, from which small amounts of the primary and secondary amines are obtained along with much *phenylpropylene* and *phenylpropane*, and still poorer results are obtained with *butyrophenone oxime*.

On the contrary, the method serves well with benzophenone-oxime from which up to 70% of the primary amine, (C₆H₅)₂CH.NH₂, is obtained with a certain amount of the secondary.²⁷

385. The ketoximes of the cycloparaffines react in an analogous manner.

The hydrogenation of cyclohexanone-oxime over nickel at 190-200° gives cyclohexyl-amine regularly with a little dicyclohexyl amine and aniline. The results are not so good with the three methyl-cyclohexanone-oximes, as the yields of the amines are poor.

The hydrogenation of cyclopentanone-oxime,

CH₂.CH₂

C: NOH

over nickel at 180°, proceeds smoothly to give a mixture of the three cyclopentyl-amines, the secondary forming half of the product and the primary and tertiary, each about one-fourth. Analogous results are obtained with methyl-cyclopentanone-oxime.³⁹

³⁶ Mailh, Compt. rend., 140, 1691 (1905), Ibid., 141, 115 (1905) and Bull. Soc. Chim. (4), 15, 327 (1914).

³⁷ MAILHE and MURAT, Bull. Soc. Chim. (4), 9, 464 (1911).

²⁴ AMOROUX, Bull. Soc. Chim. (4), 9, 214 (1911).

⁸⁹ SABATTER and MAILHE, Compt. rend., 158, 985 (1914).

Menthone-oxime yields the primary and secondary amines and a little regenerated menthone.40

Camphoroxime, when hydrogenated over nickel gives the corresponding amine in good yield.41

386. Aliphatic Amides. Acetamide is readily hydrogenated at 230° by nickel with the production of water and ethylamine and also some dimethylamine, due to the decomposition of the primary amine by the metal, and a small amount of ammonia.

Propionamide, CH₂.CH₂.CO.NH₂, gives results entirely similar.⁴²

- 387. Ethyl Acetoacetate. Ethyl acetoacetate, the ester of an unstable β -keto-acid, gives, when hydrogenated over nickel, a triple reaction: ⁴⁸
 - 1. A hydrogenation by substitution:

$$CH_8 \cdot CO \cdot CH_2 \cdot CO_2 \cdot C_2H_5 \rightarrow CH_8 \cdot CH_2 \cdot CH_2 \cdot CO_2 \cdot C_2H_5$$
ethyl butyrate

- 2. A breaking up of the molecule into the fragments CH₃. CO.-CH₂- and -CO₂. C₂H₅ which are hydrogenated separately, the first into acetone and then isopropyl alcohol (435), the second into ethyl formate which is decomposed, under the reaction conditions, into ethyl alcohol and carbon monoxide which may go into methane (867).
- 3. A condensation of the molecule with the formation of solid dehydroacetic acid, (CH₂.CO)₄, which is produced by the action of heat alone on ethyl acetoacetate,⁴⁴ and which the presence of the nickel, without the hydrogen, causes to be formed at 250°:

2 CH₂ . CO . CH₂ . CO₂ . C₂H₅
$$-$$
 (CH₂CO)₄ $+$ 2C₂H₅ . OH.

388. Aromatic Aldehydes. Contrary to what takes place with aliphatic aldehydes, the hydrogenation of aromatic aldehydes over nickel does not reduce them to alcohols, but tends to replace the oxygen by hydrogen, H₂, to give the aromatic hydrocarbons, which below 250° may be more or less hydrogenated to the cyclohexane hydrocarbons. There is, at the same time, some decomposition of the aldehyde into the hydrocarbon and carbon monoxide (618). Thus between 210 and 235°, benzaldehyde gives toluene and benzene according to the two reactions:

$$C_eH_a$$
. CHO + 2H₂ - H₂O + C_eH_a . CH₃
 C_eH_a . CHO - CO + C_eH_a ,

- 40 MAILHE and MURAT, Bull. Soc. Chim. (4), q, 464 (1911).
- 41 ALOY and BRUSTIER, Bull. Soc. Chim. (4), 9, 734 (1911).
- 42 MAILHE, Bull. Soc. Chim. (3), 35, 614 (1906).
- 44 SABATIER and MAILHE, Bull. Soc. Chim. (4), 3, 232 (1908).
- 44 GEUTHER, Zeit. f. Chem., 2, 8 (1866).

and these are accompanied by certain proportions of methylcyclohexane and cyclohexane, the carbon monoxide being partly reduced to methane (393).⁴⁵

389. Aromatic Ketones. The hydrogenation of aryl-aliphatic ketones, effected rapidly over a nickel of only moderate activity or at a temperature above 250°, is limited to replacing the ketone oxygen by H₂ with the production of the corresponding aromatic hydrocarbon. Thus acetophenone, C₆H₅. CO. CH₃, gives ethylbenzene, C₆H₅. C₂H₅; methyl-p.cresyl ketone, CH₃. C₆H₄. CO. CH₃, yields p.methyl-ethyl benzene; p.tert-butyl-acetophenone, (CH₃)₂C. C₆H₄. CO. CH₃, gives p.ethyl-tert-butyl-benzene; and benzyl-acetone, C₆H₅. CH₂. CH₂. CO. CH₃, yields butyl-benzene.

But when the hydrogenation is carried on at 180°, with an active nickel which is capable of hydrogenating the nucleus, the aromatic hydrocarbon is reduced to the cyclohexane derivative. One can be sure of avoiding this complication if nickel is used that has been so altered that it can not hydrogenate benzene or if the operation is carried on about 300°, the temperature at which cyclohexane derivatives are dehydrogenated even in excess of hydrogen.⁴⁷

It is the same way with diaryl ketones which are quantitatively reduced to the corresponding hydrocarbons by nickel at 300°.

Thus benzophenone at 300° is entirely reduced to diphenylmethane, while with an active nickel at 160°, dicyclohexylmethane is formed.

Desoxybenzoïne, C₆H₅. CH₂. CO. C₆H₅, yields dibenzyl, C₆H₅. CH₂. CH₂. C₆H₅, at 350°. Likewise dibenzyl ketone, C₆H₅. CH₂. CO. CH₂. C₆H₅, is 70% transformed at 400° into symmetrical diphenylpropane, which is accompanied by toluene formed by the breaking up of the molecule with the separation of carbon monoxide which is reduced to methane. The same hydrocarbon is formed by the hydrogenation over nickel at 350° of phenyl-phenylethyl ketone, C₆H₅. CO. CH₂. CH₂. C₆H₅. 48

390. Likewise methyl- α -naphthyl-ketone yields α -ethylnaphthalene, and methyl- β -naphthyl and the propyl-naphthyl ketones behave in a similar manner.⁴⁹

Hexahydroanthrone is hydrogenated at 200° into octohydroanthracene: 50

⁴⁵ Sabatier and Senderens, Compt. rend., 137, 301 (1903).

⁴⁶ Darzens, Compt. rend., 139, 868 (1904).

⁴⁷ Sabatier and Murat, Ann. Chim. (9), 4, 263 (1915).

⁴⁸ SABATIME and Murat, Ann. Chim. (9), 4, 264 (1915).

⁴⁹ DARZENS and Rost, Compt. rend., 146, 933 (1908).

⁵⁰ Godchot, Bull. Soc. Chim. (4), 1, 712 (1907).

$$C_0H_{10} \xrightarrow{CH_3} C_0H_4 \rightarrow C_0H_{10} \xrightarrow{CH_2} C_0H_4$$

Likewise methyl(1) cyclopentanone (3) is advantageously transformed at 250° into methylcyclopentane. 51

drogenated at 180° to form methyl-isopropyl-cyclopentane, boiling at 132°.52

391. Aromatic Diketones. Similarly to the monoketones, the aromatic diketones, when hydrogenated over nickel, give the hydrocarbons.⁵⁸

Dibenzoyl, C_0H_5 . CO. CO. C_0H_5 , which is an α diketone, is hydrogenated over nickel at 220° to symmetrical diphenylethane, or dibenzyl, C_0H_5 . CH_2 . CH_2 . C_0H_5 , beautiful crystal flakes, without appreciable secondary reactions.

Benzoïne, C₆H₅.CH(OH).CO.C₆H₅, gives the same hydrocarbon as the sole product at 210-220°.

Benzoyl-acetone, C_8H_8 . CO. CH_2 . CO. CH_8 , which is a β diketone, when hydrogenated over nickel at 200°, reacts in two ways:

- 1. Butylbenzene is formed to an extent of about 80%.
- 2. Following a general tendency of β diketones, there is a breaking up into two fragments, C_6H_8 . CO- and -CH₂. CO. CH₂, which are hydrogenated separately, the one into toluene and the other into acetone, and then into isopropyl alcohol.
- 392. Anhydrides of Dibasic Acids. The anhydrides of dibasic acids which have been submitted to hydrogenation at low temperatures, have given only the corresponding lactones.

Succinic anhydride gave butyrolactone: 54

$$\begin{array}{cccc} \text{CH}_{2}.\text{CO} & \rightarrow & \text{CH}_{2}.\text{CH}_{3} \\ \text{CH}_{2}.\text{CO} & \rightarrow & \text{CH}_{2}.\text{CO} \end{array} \rangle 0.$$

Over nickel at 200°, phthalic anhydride yields phthalid quantitatively:

$$C_0H$$
 CO
 $O \rightarrow C_0H$
 CH_2
 O
 O

- ⁵¹ Zelinbey, Berichte, 44, 2781 (1911).
- 52 Godchor and Taboury, Compt. rend., 156, 470 (1913).
- 58 SABATIER and MAILHE, Compt. rend., 145, 1126 (1907).
- 54 EIJKMANN, Chem. Weekblad, 4, 191 (1907).

Even by operating at 130° with very active nickel it is impossible to replace the second carbonyl.⁵⁵

In the same manner camphoric anhydride is changed into campholid exclusively: 54

393. Carbon Monoxide. The direct hydrogenation of carbon monoxide over nickel gives a simple method for the synthesis of methane:

$$CO + 3H_2 - H_2O + CH_4$$

The reaction commences around 180-200° and goes on rapidly without complications at 230-250°. With the theoretical mixture of hydrogen and carbon monoxide, 3:1, the reaction is practically complete, the resulting gas being nearly pure methane.

The nickel is not sensibly altered by the reaction when it is carried on below 250° and can be used indefinitely. On cooling it is found to be slightly carbonized but still pyrophoric and completely soluble in dilute hydrochloric acid without carbonaceous residue.

The reaction is less complete when the carbon monoxide is in excess; in an experiment carried out with 85 volumes of carbon monoxide to 51 volumes of hydrogen, almost one third of the hydrogen passed through the tube without combining, although the velocity of the gas was no greater than in the experiment quoted above.

394. If the operation is carried on above 250°, complications arise due to the special effect that finely divided nickel has on carbon monoxide which it breaks up into carbon and carbon dioxide (614):

$$2CO = C + CO_2$$

The carbon dioxide which is thus formed is partially hydrogenated. Its proportion is greater, the higher the temperature, since the secondary reaction which produces it is greatly accelerated by rise of temperature.

Thus when operating at 380° with the theoretical mixture which gives methane completely at 250°, a gas is obtained which contains:

 Carbon dioxide
 10.5% by vol.

 Methane
 67.9

 Hydrogen
 21.6

⁵⁴ EIJKMANN, Chem. Weekblad, 4, 191 (1907).

⁵⁵ Godchor, Bull. Soc. Chim. (4), 1, 243 (1907).

At the same temperature, water gas, equal volumes of hydrogen and carbon monoxide, gives 52.6% carbon dioxide, 39.8% methane, and 7% hydrogen.

When the percentage of carbon monoxide is still further increased, the hydrogenation is greatly weakened; much hydrogen passes through and the proportion of carbon dioxide becomes very large.⁵⁶

395. Carbon Dioxide. Like the monoxide, carbon dioxide is readily hydrogenated over nickel to form methane:

$$CO_{\bullet} + 4H_{\bullet} - CH_{\bullet} + 2H_{\bullet}O.$$

The reaction begins at a higher temperature than that with carbon monoxide, namely, around 230°, and is rapid above 300° and does not offer any considerable complications up to 400°. The theory calls for four volumes of hydrogen to one of carbon dioxide. With gas mixtures containing a larger proportion of hydrogen, the carbon dioxide disappears almost completely.

Thus in an experiment with 82% of hydrogen and 18% carbon dioxide, passed through the tube containing the nickel at the rate of 55 cc. per minute, the issuing gas contained:

- 396. Carried on at 300° with an excess of hydrogen, this reaction gives a very advantageous method for preparing *pure methane* if liquid air is available for condensing the methane. The gas is washed with caustic potash to free it from traces of carbon dioxide, dried and the methane condensed, leaving the hydrogen as gas.⁵⁷
- 397. Application to the Manufacture of Illuminating Gas. The production of methane by the direct hydrogenation of carbon dioxide over nickel can be used for the commercial preparation of a gas rich in methane having a high calorific power and capable of being used either for heating or for lighting by using incandescent mantles.⁵⁶

If hydrogen is available (produced electrolytically or by the action of iron on steam at a red heat), the hydrogenation of carbon dioxide over nickel at 300 to 400° is an excellent way to prepare methane.⁵⁰

But the preparation of the hydrogen costs too much for it to be used for the manufacture of illuminating gas. One must start with

⁵⁴ SABATIER and SENDERENS, Compt. rend., 134, 514 (1902).

⁵⁷ SABATIER and SENDERENS, Compt. rend., 134, 689 (1902).

⁵⁸ SABATIER, VI Internat. Cong. Pure and App. Chem., Rome, 1906, IV sect. p. 188.

⁵⁹ SABATIER, French patent, 356,471, June 17, 1905.

a cheap commercial gas such as water gas, Riché gas, Siemens gas, etc. Various methods may be followed.

398. First method. Water gas obtained by the action of steam on red-hot carbon varies in composition according to the temperature at which it is prepared.

At a bright red, there are equal volumes of hydrogen and carbon monoxide:

$$C + H_{\bullet}O = CO + H_{\bullet}$$

At a lower temperature (a very dull red) there are only carbon dioxide and hydrogen:

$$C + 2H_2O = CO_2 + 2H_2$$

If the temperature is intermediate (cherry red), the reaction is intermediate:

$$2C + 2H_2O = CO + CO_2 + 3H_2$$
.

If in this case the carbon dioxide be removed by any method there remains the mixture $CO + 3H_2$. The carbon dioxide may be absorbed by a solution of potassium carbonate which is changed to the bicarbonate, but is regenerated with evolution of carbon dioxide by boiling. The carbon dioxide may be solidified by refrigeration or absorbed in cold water under pressure. The residual mixture, $CO + 3H_2$, is converted into pure methane by passing over nickel at 230–250°, 5 volumes of water gas thus furnishing 1 volume of methane. A practical difficulty arises from the fact that the catalyst must be kept between 230 and 250°, since above 250° there is charring with loss of carbon and fouling of the nickel resulting in a rapid diminution of its catalytic power.

399. Second Method. The operation is carried on in two phases: Water gas prepared at a high temperature and very nearly CO + H₂ is passed over nickel at 400 to 500°, by which all the carbon monoxide disappears forming either methane with the available hydrogen, or splitting up into carbon dioxide (614) and finely divided carbon which is deposited on the nickel. If from the gas so produced, the carbon dioxide is absorbed, the remainder is very rich in methane. For the conditions cited above (394), the composition would be 83.8% methane and 15% hydrogen with a calorific power of 7,800 calories per cu. m., while the original gas had only 2,880. This is the gas of the first phase.

If steam be passed over the intimate mixture of carbon and nickel obtained above, kept at 400 to 500°, the carbon reacts rapidly tending to give hydrogen and carbon dioxide which being in the nascent

state react to give a certain proportion of methane. The final product is a mixture of hydrogen, methane and carbon dioxide and if the carbon dioxide is eliminated, there remains a mixture of hydrogen and methane of high calorific power which can be used. This is the gas of the second phase, less rich in methane than the first. Its formation has eliminated the carbon from the nickel which is then ready to repeat the first phase of the reaction.⁶⁰

400. Third Method. The gas of the second phase can be obtained alone by preparing at first the intimate mixture of nickel and carbon by the action of finely divided nickel on various gases rich in carbon monoxide such as Siemens gas or producer gas. The carbon monoxide disappears leaving carbon dioxide and carbon. It is sufficient to maintain this carbonaceous mass at 400 to 500° and pass superheated steam over it to have a mixture of methane, hydrogen and carbon dioxide which can be used after the latter is eliminated.⁶¹

401. Fourth Method. The two phases of the reaction that have just been described can be combined in practice. All that is required is to maintain finely divided nickel at 400 to 500° and pass over it a mixture of suitable proportions of water gas (or Riché gas 62) and superheated steam. Under these conditions the carbon monoxide disappears and is replaced by hydrogen, methane and carbon dioxide, and if the latter is eliminated, we have in one operation a usable mixture of hydrogen and methane.

This method of operating appears economical. The amount of nickel required for the reaction is less than 1 k. for making 1 cu. m. of gas per hour. Besides, if the carbonated gases introduced are suitably purified and if this purification is completed by passing over copper turnings heated to 600°, the nickel may be said to retain its catalytic power indefinitely. By starting with water gas a gas is obtained having an average composition of 48% methane and 52% hydrogen and having a calorific power of 5,800 calories per cu. m. This gas does not contain an appreciable amount of carbon monoxide which is present in coal gas in considerable amount (from 8 to 15%) and which renders it decidedly toxic.

In fact the reactions that take place with these conditions under the influence of nickel between water gas and steam can be summed up in this equation:

$$\frac{5(CO + H_2)}{\text{water can}} + H_2O = 2CH_4 + 2H_2 + 3CO_2.$$

⁶⁰ SABATHER, French patent, 355,900, July 5, 1905.

⁶¹ SABATTER, French patent, 355,900, 1905.

⁶² The Riché gas is a mixture of carbon monoxide, hydrogen, methane and carbon dioxide prepared by heating woody or cellulose materials.

Theoretically 5 volumes of perfect water gas should give 2 volumes of the mixture containing 50% methane. In practice, as the water gas contains some carbon dioxide, about 3 volumes are required on the average for 1 volume of the finished gas.

402. The use of industrial refrigeration permits a very advantageous modification of the first process (398). The water gas should be prepared at the highest possible temperature so as to contain $CO + H_2$ and a little carbon dioxide and nitrogen. By suitable refrigeration 75% of the carbon monoxide can be liquefied and a mixture of $CO + 4H_2$ obtained which passing over nickel at 200 to 250° would furnish exactly the gas $CH_4 + H_2$ equivalent to coal gas. The refrigeration condenses all of the substances that may be toxic to the nickel (sulphurous gases, etc.) and hence guarantees the long life of the catalyst.

The carbon monoxide separated by the liquefaction may be used for heating the catalyzers or for driving motors. 68

403. Aromatic Halogen Derivatives. The direct reduction of aromatic halogen derivatives by hydrogen in the presence of nickel may take place more or less readily: it is easy with chlorine derivatives, less easy for bromine derivatives, and difficult for iodine compounds; the reason being easy to find in the decreasing affinity of the halogens for hydrogen as we pass from chlorine to iodine, since the simultaneous formation of the hydro-acid determines the substitution of hydrogen.

When the vapors of chlorbenzene are carried by hydrogen over reduced nickel at 160°, a strong absorption of hydrogen is noted at once and a little cyclohexane is condensed without any chlorcyclohexane. The chlorine remains, fixed by the nickel, the surface of which loses all activity by being changed to the chloride. After a short time the chlorbenzene passes through unchanged.

But if the temperature is raised above 270° a vigorous evolution of hydrochloric acid is observed and a readily separated mixture of benzene and chlorbenzene is condensed. At the same time there is the formation of crystals of diphenyl.

In contact with nickel at 270° or above, chlorbenzene gives nickel chloride, and the liberated residue, C₆H₅-, combines with hydrogen to give benzene and unites with itself to form a small amount of diphenyl. But at this temperature the nickel chloride is reduced by hydrogen forming hydrochloric acid and regenerating the nickel which repeats the reaction indefinitely.

404. An analogous reduction is observed when the polychlor-

⁶⁸ Saratier, Second Congress on Refrigeration, 1, 115 (1912).

derivatives of benzene are acted on by hydrogen in the presence of nickel above 270°; the chlorine atoms are progressively replaced by hydrogen.

Thus m.dichlorbenzene gives a mixture containing:

Benzene	. 30%
Monochlorbenzene	. 60%
Unchanged dichlorbenzene	. 10%

p.Dichlorbenzene gives 35% benzene and 65% monochlorbenzene. Perchlorbenzene, C_eCl_e, acts in the same way at 270° and gives a mixture of the trichlorbenzenes (particularly the 1,2,4), dichlorbenzenes, monochlorbenzene, and benzene.

The presence of aliphatic side-chains and hydroxyl groups facilitates the reduction, the *chlortoluenes* being more readily reduced than chlorbenzene.

2,4,6-Trichlorphenol is readily reduced at 270° and gives 70% of phenol accompanied by monochlorphenols, particularly the ortho.

The reduction goes even better with amino derivatives, such as the chloranilines which give aniline hydrochloride at 270°.

The chlornitrobenzenes suffer simultaneous reduction of the nitro group and elimination of the chlorine, furnishing aniline hydrochloride at 270°.64

- 405. It can be foreseen that the reduction of bromine derivatives will be more difficult, since the temporary nickel bromide is less easily reduced by hydrogen. However, the reaction can be carried out well with monobrombenzene at 270° and also with p.bromtoluene, the bromanilines and the bromanitrobenzenes.
- 2,4,6-Tribromphenol readily yields phenol accompanied by p.bromphenol and 2,4-dibromphenol.
- 406. The difficulties are greater for the iodine derivatives. Iodobenzene passed over nickel with hydrogen at 270° gives no lasting evolution of hydriodic acid; some benzene and diphenyl are formed, but the reaction stops, since the nickel is not restored by the hydrogen and does not continue the reaction.

If pure hydrogen is passed into the tube, fumes of hydriodic acid appear, hence nickel iodide is reduced by hydrogen at 270° but not in the presence of iodobenzene, doubtless because this compound gives iodine to the nickel faster than the hydrogen can remove it. Practically, the reduction of iodobenzene can be forced by alternately passing pure hydrogen and hydrogen mixed with iodobenzene vapors

⁶⁴ SABATTER and MAILHE, Compt. rend., 138, 245 (1904).

over the nickel at 270°. But under these conditions the metal is not a true catalyst. 65

- 407. Esters of Halogenated Aliphatic Acids. Vapors of ethyl mono-, di-, and tri-chloracetates, when passed over nickel at 300°, with excess of hydrogen, are reduced to ethyl acetate, the chlorine atoms being successively replaced by hydrogen. Ethyl bromacetate is as readily reduced to ethyl acetate. 66
 - 65 SABATIER and MAILHE, Compt. rend., 138, 245 (1904).
 - 66 SABATIER and MAILHE, Compt. rend., 169, 758 (1919).

CHAPTER IX

HYDROGENATIONS (Continued)

HYDROGENATIONS IN THE GAS PHASE — USE OF NICKEL (Continued)

ADDITION OF HYDROGEN

- 408. Many hydrogenations correspond to the fixation of hydrogen by addition. This addition takes place either to free carbon, which is rare, or to complex molecules containing double or triple bonds between the atoms. We will examine these in the following order:
 - 1. Direct fixation by carbon,
- 2. On double bond between two carbon atoms, so-called ethylene bond, C:C,
- 3. On triple bond between two carbon atoms, called the acetylene bond, C : C.
 - 4. Triple bond between carbon and nitrogen, C: N,
 - 5. Quadruple bond between carbon and nitrogen, C: N,
 - 6. Double bond between carbon and an oxygen atom, C:O,
 - 7. Aromatic nucleus.
 - 8. Various rings,
 - 9. Carbon disulphide.

1. Direct Fixation of Hydrogen by Carbon

409. Berthelot noted the direct union of hydrogen and carbon at the temperature of the electric arc¹ to form acetylene which was necessarily accompanied by some methane and ethane resulting from the pyrogenetic decomposition of the acetylene.

Bone and Jerdan state that carbon unites directly with hydrogen at 1200° forming 1 to 2% methane.²

But Berthelot, carrying out the reaction with pure carbon in a quartz tube, could not confirm the formation of methane and concluded that it must have come from impurities in the carbon used by the English chemists.³

- ¹ Berthelot, Ann. Chim. Phys. (4), 13, 143 (1868).
- ² Bonn and Jerdan, J. Chem. Soc., 71, 42 (1897).
- ⁸ BERTHELOT, Ann. Chim. Phys (8), 6, 183 (1905).

410. According to Henseling, the formation of methane by carbon and hydrogen begins at 300° in the presence of finely divided nickel.

Sabatier and Senderens, by passing hydrogen at 250° over the intimate mixture of carbon and nickel which is formed by the action of reduced nickel on carbon monoxide between 250 and 300°, have definitely proved the production of methane, but also detected water vapor. After some time the formation of methane ceased though there was still much carbon with the metal. They attributed the formation of methane and water to the presence of a nickel carbonyl combination formed by the action of the carbon monoxide. The same chemists found no methane when the carbonaceous mixture had been prepared above 400°, a temperature at which carbonyl compounds can not exist.⁴

411. Mayer and Altmayer have confirmed the very slow formation of methane from carbon in contact with nickel or cobalt. At all temperatures above 250° methane is decomposed by nickel into carbon and hydrogen, the amount remaining being fixed for each temperature, and the same whether the limit be approached from above or from below as is true with all reversible reactions (19), and not altered when cobalt is substituted for nickel. The amounts of methane at equilibrium are:

\mathbf{At}	250°	• • • • • • • • • • • • • • • • • • • •	98.8%	by	volume
	536°	• • • • • • • • • • • • • • • • • • • •	51.2%		
	625°	• • • • • • • • • • • • • • • • • • • •	24.7%		
	850°		1.6%		

But this formation is very slow and could never be used for the preparation of methane. The velocities of the mixtures of gases passed over the mixture of carbon and nickel to obtain the equilibrium were not over 0.2 to 0.3 cc. per minute.⁵

2. Ethylene Double Bond

- 412. The ethylene double bond is very easily attacked by direct hydrogenation over nickel and adds two atoms of hydrogen. This is readily accomplished by nickel reduced above 500° and even by nickel which has been weakened by the action of poisons.
- 413. Hydrocarbons. Ethylene is hydrogenated by nickel from 30° up, the reaction which continues indefinitely, with evolution of heat, gives ethane exclusively. The hydrogenation is more rapid toward 130-150°.
 - 4 SABATTER and SENDERENS, Bull. Soc. Chim. (4), 1, 107 (1907).
 - ⁵ MAYER and ALTMAYER, Berichte, 40, 2134 (1907).
 - 6 SABATTER and SENDERENS, Compt. rend., 124, 1359 (1897).

In the presence of excess of hydrogen, all the ethylene disappears, while with excess of ethylene all the hydrogen is used up and a mixture of ethane and ethylene is obtained from which it is easy to remove the latter by bromine water leaving the ethane pure.

This reaction has been used for the manufacture of ethane for refrigerating machines. The mixture of equal volumes of ethylene and hydrogen is passed through tubes 1 m. long and 7.5 cm. in diameter containing reduced nickel and heated to 200°. With a velocity of 2 cu. m. per hour a gas containing 80% of ethane is obtained. In order to complete the union of hydrogen and ethylene the mixture is compressed to 30 or 40 atmospheres in a vessel filled with nickeled pumice.

Above 300° nickel decomposes ethylene (912) with the liberation of carbon, and the production of methane and certain amounts of higher paraffines which can be liquefied.

414. Other ethylene hydrocarbons can be transformed into the corresponding saturated hydrocarbons below 160° without any complications. But above 200° and particularly above 300° there can be partial breaking of the carbon chain with the formation of saturated hydrocarbons with smaller numbers of carbon atoms and also more complicated.

With propylene, CH₂.CH:CH₂, the reaction commences in the cold and up to 200° nothing but propane, CH₃.CH₂.CH₃, is produced so long as the hydrogen is in slight excess. When the propylene is in excess, particularly above 290°, small amounts of higher liquid hydrocarbons with petroleum odors are formed, and at higher temperatures there is more and more deposition of carbon with splitting up of the propane.

Trimethyl-ethylene, or 2-methyl-butylene, (CH₂)₂C: CH.CH₂, is totally hydrogenated by excess of hydrogen into pure 2-methyl-butane or isopentane, at 150°.

Likewise hexene (2) gives hexane; and caprylene, or octene (1), octane without complications below 160°.

By the hydrogenation of 2,2-dimethyl-methylene (3)-pentane, over nickel at 160°, 2,2,3-trimethyl-pentane, boiling at 110.5° is obtained, and likewise 2,5-dimethyl-heptane, boiling at 135° from 2-ethyl-5-methyl-hexene.

- 6 SABATTER and SENDERENS, Compt. Rend., 124, 1359 (1897).
- ⁷ SPRENT, J. Soc. Chem. Ind., 32, 171 (1913).
- * SABATTER and SENDERENS, Compt. rend., 134, 1127 (1902).
- ⁹ CLARK and JONES, J. Amer. Chem. Soc., 34, 170 (1912). CLARK and Beggs, Ibid., 34, 54 (1912).

Likewise nonene (2) is transformed entirely into nonane. 10

Methyl-propyl-octene gives the corresponding methyl-propyl-octane, and 4-cyclohexyl-heptene, the 4-cyclohexyl-heptane.¹¹

415. In the case of phenyl- or polyphenyl-ethylene hydrocarbons, when the hydrogenation is carried out with a weakened nickel such as is not capable of hydrogenating benzene (56), or with active nickel at 300°, the aliphatic double bonds are saturated without hydrogenating the aromatic nuclei.

Thus styrene, C_eH₅. CH: CH₂, gives only ethyl-benzene, C_eH₅. CH₂CH₅.

The ortho, meta, and para, cresyl-propenes (2) are regularly changed into the ortho, meta, and para cymenes.¹²

1-Phenul-2-propul-pentene vields 1-phenul-2-propul-pentane.11

Stilbene, or symmetrical diphenyl ethylene, C_6H_5 .CH:CH.C $_6H_5$, is readily transformed by a slightly active nickel at 240° into dibenzyl, C_6H_5 .CH₂.CH₂.C $_6H_5$. Likewise $\alpha\alpha$ -Diphenyl-ethylene is readily changed to $\alpha\alpha$ -diphenyl-ethane, 1,2-diphenyl-propene(1) and 1,1-diphenyl-propene(2) furnish the corresponding diphenyl-propanes and similar statements hold for the diphenyl-butenes and diphenyl-pentenes.¹⁸

Ocimene, (CH₃)₂C:CH.CH₂.CH:C.CH:CH₂, or 2,6-dimethyl-octa-CH₃

triene (2,5,7), of oil of basil is readily hydrogenated over nickel at 130-140° to the corresponding 2,6-dimethyl-octane boiling at 158°.14

416. Unsaturated Alcohols. The fixation of hydrogen frequently takes place without alteration of the alcohol group.

Propenol, or allyl alcohol, CH₂: CH.CH₂OH, is readily hydrogenated at 130-170° over nickel, to give nearly pure propyl alcohol containing only a slight amount of propionic aldehyde.¹⁵

Geraniol, (CH₃)₂C:CH.CH₂.CH₂.C:CH.CH₂OH, or 2,6-dimethyl-

ĊH,

octadiene (2,6) ol (8), is readily hydrogenated at 130-140° to give the corresponding dimethyl-octanol. At the same time a little of it is reduced to the saturated hydrocarbon.

- 10 CLARK and JONES, J. Amer. Chem. Soc., 37, 2536 (1915).
- 11 MURAT and AMOUROUX, J. Pharm. Chim. (7), 5, 473 (1912), C. A., 7, 1494.
- 12 SABATIER and MURAT, Compt. rend., 156, 184 (1913).
- 18 SABATTER and MURAT, Ann. Chim. (9), 4, 284-297 (1915).
- 14 ENKLAAR, Berichte, 41, 2085 (1908).
- 15 SABATIER, Compt. rend., 144, 879 (1907).

The hydrogenation of linalool, or 2,6-dimethyl-octadiene (2,7)-ol(6), (CH₃)₂C: CH. CH₂. CH₂. C(OH). CH: CH₂, furnishes the

CH,

same products.16

Citronellol, (CH₃)₂C: CH. CH₂. CH₂. CH₂. CH₂OH, like-CH₃.

wise gives dihydrocitronellol.17

We have seen (208) that the hydrogenation, over nickel at 200°, of secondary α -unsaturated alcohols gives the isomeric saturated ketone instead of the saturated secondary alcohol, by a simple migration of the hydrogen of the alcohol group.

417. Esters. The esters of unsaturated acids are readily hydrogenated over nickel whatever be the position of the double bond.

Esters of acrylic acid give esters of propionic at 180°.

Ethyl dimethyl-acrylate likewise gives ethyl isovalerate, and ethyl undecylenate, the undecylate.

It is the same way with ethyl enanthylidene-acetate, C_eH₁₈.-CH: CH. CO₂. C_eH₂₈.

The same fixation of hydrogen takes place with the esters of unsaturated aromatic acids without the hydrogenation of the nucleus. Methyl cinnamate, C_6H_5 . CH: CH. CO₂. CH₃, gives methyl phenyl-propionate.

Ethyl phenyl-isocrotonate, $C_0H_5 \cdot CH : CH \cdot CH_2 \cdot CO_2 \cdot C_2H_5$, acts in a similar manner.¹⁸

418. Ethers of Unsaturated Alcohols. The vapors of allyl ether, carried by an excess of hydrogen over nickel at 138-140° are totally changed to propyl ether.¹⁹

the side chain to dihydrosafrol without affecting the ether group.20

419. Unsaturated Aldehydes. Acroleine, CH₂: CH . CHO, is hydrogenated over nickel at 160° to propionic aldehyde,²¹ which can be further hydrogenated, by a slower reaction, to propyl alcohol.

Likewise crotonic aldehyde over nickel at 125° is changed to buty-

- ¹⁶ ENKLAR, Rec. Trav. Chim. Pays-Bas, 27, 411 (1908), and Berichte, 41, 2085 (1908).
 - ¹⁷ Haller and Martine, Compt. rend., 140, 1303 (1905).
 - ¹⁸ Darzens, Compt. rend., 144, 328 (1907).
 - ¹⁹ Sabatier, Compt. rend., 144, 879 (1907).
 - ²⁰ HENRARD, Ch. Wkbld., 4, 630-2; Chem. Cent., 1907 (2), 1512.
 - ²¹ Sabatier and Senderens, Ann. Chim. Phys. (8), 4, 399 (1905).

ric aldehyde with a yield of 50%, with about 20% of butyl alcohol resulting from the subsequent hydrogenation of the aldehyde.²²

420. Unsaturated Ketones. The fixation of hydrogen on ethylene double bonds is so rapid that it can be effected before the ketone group, -CO-, is changed to the secondary alcohol group, -CH(OH)-.

Mesityl oxide, (CH₂)₂C: CH.CO.CH₂, is transformed at 160-170° into 2-methyl-pentanone (4), ²² accompanied by some of the corresponding alcohol and even of isopentane. ²⁴ Likewise methyl-hexenone, (CH₂)₂C: CH.CH₂.CO.CH₂, gives the corresponding methyl-hexanone.

3-Methyl-heptene (3) one (5) is transformed at 180° into 3-methyl-heptanone (5), and likewise 2,4,8-trimethyl-nonene (4) one (6) gives the corresponding saturated ketone.²⁶

Phorone, (CH_s)₂C: CH. CO. CH: C(CH_s)₂, when hydrogenated over nickel at 160-170°, is totally changed to di-isobutyl-ketone, or isovalerone.²⁶ By operating at 225° the ketone is accompanied by the alcohol and the saturated hydrocarbon.²⁷

421. By hydrogenating pulegone rapidly over nickel at 140-160°, the unsaturated side chain can be hydrogenated without affecting the ketone group to give pulegomenthone: ²⁸

$$(CH_3)_2C:C \xrightarrow{CO.CH_2} CH.CH_3 \rightarrow (CH_3)_2CH.CH \xrightarrow{CO.CH_2} CH.CH_4.$$

$$CH_2.CH_2 \rightarrow CH.CH_3 \rightarrow (CH_3)_2CH.CH \xrightarrow{CO.CH_2} CH.CH_4.$$

$$CH_3.CH_2 \rightarrow CH.CH_4 \rightarrow (CH_3)_2CH.CH_3 \rightarrow (CH_3)_2CH.CH_4.$$

$$CH_4.CH_2 \rightarrow CH.CH_3 \rightarrow (CH_3)_2CH.CH_4 \rightarrow (CH_3)_2CH.CH_3 \rightarrow$$

over nickel at 130° to give dihydrocamphorone, boiling at 182°.20

422. Unsaturated Acids. Their hydrogenation is readily carried out over nickel without any damage to the catalytic metal. The vapors of crotonic acid, CH₂. CH: CH. COOH, at 190° give butyric acid quantitatively. The vapors of oleic acid, carried along by a violent current of hydrogen over nickel at 280–300°, are readily transformed into solid stearic acid, and the same is true of its isomer elaidic acid.²⁰

- ²² Douris, Bull. Soc. Chim. (4), 9, 922 (1911).
- 28 DARZENS, Compt. rend., 140, 152 (1905).
- 24 SKITA, Berichte, 41, 2938 (1908).
- 25 Bodroux and Taboury, Compt. rend., 149, 422 (1909).
- 26 SABATTER and MAILHE, Ann. Chim. Phys. (8), 16, 79 (1909).
- 27 SKITA, Loc. cit.
- 28 HALLER and MARTINE, Compt. rend., 140, 1298 (1905).
- 29 GODCHOT and TABOURY, Compt. rend., 156, 470 (1913).
- ⁸⁰ SABATTER and MAILHE, Ann. Chim. Phys. (8), 16, 73 (1909).

3. The Acetylene Triple Bond

423. If hydrogen mixed with a small proportion of acetylene is passed over cold reduced nickel, the metal becomes warm, the more so when the proportion of acetylene is increased. With 2 volumes of hydrogen to 1 of acetylene the spontaneous evolution of heat may heat the end of the nickel train to 150°. The contraction of the gas is enormous, greater than corresponds to the formation of ethane:

$$C_2H_2 + 2H_2 - C_2H_6$$
.

The volume is reduced to one fourth, although a little unchanged acetylene and some *ethylene* remain, showing incomplete hydrogenation, because there is produced at the same time a considerable proportion of higher hydrocarbons, part of which are liquefied. The nickel is coated with a little carbon which is readily separated by dilute acids.

On the contrary, the formation of ethane is complete in the presence of an excess of hydrogen.

424. Inversely if the proportion of acetylene in the mixture is increased, the metal heats up more, the liquids formed become more abundant and the presence of hydroaromatic and aromatic hydrocarbons can be shown. Finally, an incandescence is noticed similar to that produced by acetylene alone on nickel (914).⁸¹

425. α -Heptine, or cenanthylidene, is readily hydrogenated over nickel to n.heptane.*

4. The Triple Bond Between Carbon and Nitrogen

426. The direct hydrogenation of nitriles, R.C: N, easily carried out with nickel, leads to the formation of the primary amines, R.CH₂.NH₂, which on account of secondary reactions caused by the metal, are accompanied by secondary and tertiary amines. These reactions correspond to the formation of ammonia which is eliminated, and consequently the secondary amine constitutes the major portion of the product. We have:

and
$$R \cdot CH_2 \cdot NH_2 + (R \cdot CH_2)_2NH = NH_3 + (R \cdot CH_2)_2NH = NH_3 + (R \cdot CH_2)_3N$$
.

427. Aliphatic Nitriles. Formic-nitrile or hydrocyanic acid is not

⁸¹ SABATIER and SENDERENS, Compt. rend., 128, 1173 (1899).

⁸² Sabatier and Senderens, Compt. rend., 135, 87 (1902).

affected by hydrogenation except above 250° but then gives the three methyl-amines and ammonia.

Acetonitrile is readily hydrogenated at 200° and gives a mixture containing 60% diethyl-amine, and 20% each of the mono- and triamines.

With ethyl cyanide, the dipropyl-amine forms nearly 80% of the product.

Isoamyl cyanide likewise gives chiefly secondary amine, the primary being formed in least amount. The amines are accompanied by a little isopentane.

It is evident that the hydrogenation of aliphatic nitriles gives us a valuable and convenient method of preparing secondary amines.³⁸

428. Aromatic Nitriles. The results are not nearly as good with aromatic nitriles from which the hydrocarbons and ammonia are formed.

However, the hydrogenation of benzonitrile at 250° gives a certain proportion of benzyl-amine and dibenzyl-amine and the same is true of p.toluo-nitrile which gives a mixture of the primary and secondary amines.²⁴

429. Dicyanides. Ethylene dicyanide, when hydrogenated over nickel, gives a certain proportion of tetramethylene-diamine resulting from the regular hydrogenation:

$$CN \cdot CH_{\bullet} \cdot CH_{\bullet} \cdot CN + 4H_{\bullet} = NH_{\bullet} \cdot CH_{\bullet} \cdot CH_{\bullet} \cdot CH_{\bullet} \cdot CH_{\bullet} \cdot NH_{\bullet}$$

This is accompanied by a little ammonia and pyrrolidine,

5. Quadruple Bond Between Carbon and Nitrogen

430. Carbylamines. The aliphatic isocyanides, or carbylamines, R.N.C, which former wet reduction methods were unable to hydrogenate because they were decomposed by hydration, can add 4H over nickel at 160–180° to form the secondary amines, R.NH.CH₃

They are accompanied by a small amount of the primary amine, R.CH₂.NH₂, and the secondary amine, (R.CH₂)₂NH, resulting from the hydrogenation of the nitrile, R.C:N, produced by partial isomerization of the isocyanide.

Methyl carbylamine gives a yield of 80% of dimethyl-amine.

⁸⁸ SABATIER and SENDERENS, Compt. rend., 140, 482 (1905).

⁸⁴ FRÉBAULT, Compt. rend., 140, 1036 (1905).

⁸⁵ GAUDION, Bull. Soc. Chim. (4), 7, 824 (1910).

The metal is gradually coated with tarry material which diminishes its activity.

Ethyl carbylamine gives chiefly methyl-ethyl-amine with a little mono- and di-propyl amines.

Tertiary-butyl-isocyanide, (CH₃)₈C. N : C, hydrogenated at 170-180°, gives methyl-tert.butyl-amine, which has never been obtained by other methods.

If the reaction is carried on at 220–250°, the secondary amine molecule is broken up with the formation of ammonia and hydrocarbon.²⁶

431. Aliphatic Carbimides. It is convenient to consider along with the carbylamines the aliphatic carbimides, or isocyanates, R.N:CO (although the hydrogenation is not simply the addition of hydrogen but also its substitution for the oxygen atom), because the result is the same for both classes.

Over nickel at 180-190°, the chief reaction is:

$$R.N:CO + 3H_2 - H_2O + R.NH.CH_3.$$

But a disturbance is caused by the water produced which reacts immediately with a part of the carbimide to form a disubstituted urea, (R.NH)₂CO, and carbon dioxide. The alkyl urea is immediately hydrogenated, giving:

$$(R.NH)_2CO + 3H_2 - H_2O + NH_2.R + R.NH.CH_3.$$

Hence there is a certain amount of the primary amine, R. NH₂, and on account of secondary actions of the metal, the secondary, R₂NH, and tertiary, R₃N, also.

Thus ethyl isocyanate gives a considerable amount of methylethyl-amine, a little diethyl-amine, and traces of ethyl-amine and triethyl-amine.³⁷

6. Double Bond between Carbon and Oxygen

432. The carbonyl group, CO, which frequently occurs in organic compounds, is readily hydrogenated over nickel to the alcohol group, CHOH.

Aliphatic Aldehydes. Hydrogenated over nickel below 180°, these are regularly transformed into the primary alcohols without the production of di-secondary glycols or acetals as by-products.

Formaldehyde vapors at 90° are readily transformed and methyl

³⁶ SABATIER and MAILHE, Compt. rend., 144, 955 (1907).

⁸⁷ Sabatter and Mailhe, Compt. rend., 144, 824 (1907).

alcohol is condensed along with water which is due to the formation of methane according to the reaction:

$$H \cdot CO \cdot H + 2H_2 - CH_4 + H_2O$$
.

But the covering over of the metal surface with a thin coating of trioxy-methylene soon suppresses its activity. If the temperature is raised, this trouble disappears, but the formation of methane increases as well as the decomposition of the formaldehyde itself (508).

Acetaldehyde is readily transformed into alcohol around 140°, but at 200° the destruction of the aldehyde is already apparent.

Propionaldehyde is regularly hydrogenated to propyl alcohol between 100 and 145°.

It is the same with isobutyric and isovaleric aldehydes at 135-160° which yield about 70% of the alcohols, the rest of the product being unchanged aldehyde with a little acetal.

433. Aromatic Aldehydes. These do not give this reaction but tend to form the hydrocarbons; thus benzaldehyde at 210-235° gives benzene and toluene accompanied by a certain proportion of the corresponding cyclohexane compounds. The reaction which takes place is:

$$C_{e}H_{s}$$
. $CHO + 2H_{2} - C_{e}H_{s}$. $CH_{s} + H_{2}O$

along with the decomposition of benzaldehyde by nickel:

$$C_6H_6$$
. CO. $H - C_6H_6 + CO$

followed by a partial hydrogenation of the carbon monoxide to methane.²⁸

434. Pyromucic Aldehyde. Furfural, or pyromucic aldehyde, C₄H₂O. CHO, hydrogenated over nickel at 190°, gives furfuryl alcohol, C₄H₂O. CH₂OH, accompanied by some secondary products (see 371 and 487).²⁹

435. Aliphatic Ketones. Aliphatic ketones, being more stable towards nickel than the aldehydes, are hydrogenated regularly into the secondary alcohols and, unlike their conduct in the classic reduction by sodium amalgam, they do not form any secondary products such as pinacones. The method is an excellent one for the preparation of many secondary alcohols, which are produced almost quantitatively.

This process is readily applied to acetone which forms isopropyl alcohol at 115-125° which is thus prepared quite cheaply. It is no less good for butanone, diethyl-ketone, methyl-isopropyl-ketone,

as Sabatter and Senderens, Compt. rend., 137, 301 (1903).

⁸⁹ PADOA and PONTI, Lincei, 15 (2), 610 (1906), C., 1907 (1), 570.

methyl-propyl-ketone, and methyl-butyl-ketone. It is only above 200° that decompositions of the molecules begin to take place. 40

Disopropyl 41 and disobutyl 42 ketones are readily transformed into the secondary alcohols under the same conditions.

When the hydrogenation of ketones is carried out above 200° different results are obtained. Acetone hydrogenated between 200 and 300° gives neither isopropyl alcohol nor its pinacone, but chiefly methyl-isobutyl-ketone (boiling at 114°) accompanied by diisobutyl-ketone (b.165°).48

Methyl-nonyl-ketone, hydrogenated at 300°, does not give the corresponding alcohol but various products, one of them a ketone, $C_{22}H_{44}O$.

436. Alicyclic Ketones. The method is readily applied to these. Cyclopentanone is hydrogenated over nickel at 125° to give 50% of cyclopentanol, a little cyclopentane, and 40% of a complex ketone formed by the joining of two rings, the cyclopentyl-cyclopentanone.

The α - and β -methylcyclopentanones are hydrogenated at 150° to the corresponding alcohols, accompanied by greater quantities of the dimethylcyclopentyl-pentanones formed by the union of two rings.⁴⁶

Cyclohexanone and the three methyl-cyclohexanones are regularly hydrogenated below 180° to the corresponding alcohols with small amounts of the hydrocarbons.⁴⁷

genated by an active nickel at 140-160°, gives a mixture of menthol and pulegomenthol.48

437. Keto-acids. Laevulinic acid, CH₃. CO.CH₂. CH₂. COOH, hydrogenated over nickel around 250°, gives the hydroxy-acid, which loses water to form valerolactone, CH₃. CH. CH₂. CO. CO.

438. Diketones. The results of the hydrogenation of these depend on the nature of the compounds.⁵⁰

- 40 SABATTER and SENDERENS, Compt. rend., 137, 302 (1903).
- 41 AMOUROUX, Bull. Soc. Chim. (4), 7, 154 (1910).
- 42 MAILHE, Bull. Soc. Chim. (4), 15, 327 (1914).
- 48 LASSIEUR, Compt. rend., 156, 795 (1913).
- 44 HALLER and LASSIEUR, Compt. rend., 150, 1017 (1910).
- 45 Godchot and Taboury, Compt. rend., 152, 881 (1911).
- 46 Godchor and Taboury, Bull. Soc. Chim. (4), 13, 591 (1913).
- 47 SABATTER and SENDERENS, Ann. Chim. Phys. (8), 4, 402 (1905).
- 48 HALLER and MARTINE, Compt. rend., 140, 1298 (1905).
- 49 SABATIER and MAILHE, Ann. Chim. Phys. (8), 16, 78 (1909).
- ⁵⁰ Sabatier and Mailier, Compt. rend., 144, 1086 (1907).

α-Aliphatic Diketones. Diacetyl, or butanedione, CH₃.CO.-CO.CH₃, by hydrogenation at 140–150°, is totally transformed into a mixture of butanol-one (2,3), CH₃.CH(OH).CO.CH₃, and butanediol (2,3), CH₃.CH(OH).CH(OH).CH₃.

439. β -Aliphatic Diketones. Acetyl-acetone, CH₃. CO.CH₂.-CO.CH₃, when hydrogenated at 150°, gives two simultaneous reactions. One part is normally hydrogenated to form pentol(2) one (4), CH₃. CH(OH). CH₂. CO.CH₃, while the larger portion is split into two fragments:

$$CH_3 \cdot CO \cdot CH_2 \cdot CO \cdot CH_3 + H_2 = CH_3 \cdot CO \cdot H + CH_3 \cdot CO \cdot CH_3$$

The acetaldehyde and acetone thus formed are then reduced to ethyl and isopropyl alcohols.

Methyl-acetyl-acetone, CH₃. CO.CH(CH₃). CO.CH₃, forms hardly anything but the decomposition products.

440. γ -Aliphatic Diketones. Acetonyl-acetone, CH₃.CO.CH₂.CH₂.CO.CH₃, when hydrogenated at 190°, is totally transformed, not into the corresponding diol but into the ether, CH₃.CH₂.CH₂.CH₃.CH₃, produced by its dehydration.

441. Aromatic Ketones. Aromatic ketones and diketones give the corresponding alcohols by hydrogenation but go chiefly into the

hydrocarbons (389 et seq.).

442. Quinones. We can consider quinones as unsaturated alicyclic diketones. They are readily hydrogenated by nickel at 200°, and add H₂ to form the corresponding diphenols in excellent yields.

This is the case with ordinary quinone which gives hydroquinone quantitatively, with toluquinone, with p.xyloquinone, and with thymoquinone.

But if the operation is carried on at a higher temperature, 220 to 250°, the diphenol is no longer obtained, but water, the monophenol, and even the hydrocarbon.⁵¹

443. Ethylene Oxides. The direct hydrogenation of these oxides is doubtless readily carried out in all cases.

In the particular case of the ether of cyclohexanediol(1,2), hydrogen is added at 160° to give a quantitative yield of cyclohexanol: 52

$$\begin{array}{c} \mathrm{CH_2.CH_2.CH} \\ \cdot \\ \mathrm{CH_2.CH_2.CH} \end{array} \rightarrow \begin{array}{c} \mathrm{CH_2.CH_2.CHOH} \\ \cdot \\ \mathrm{CH_2.CH_2.CH_2} \end{array}$$

⁵¹ SABATIER and MAILHE, Compt. rend., 146, 457 (1908).

⁵² BRUNEL, Ann. Chim. Phys. (8), 6, 237 (1905).

7. The Aromatic Nucleus

444. The direct hydrogenation of the aromatic nucleus has long been considered very difficult to accomplish. When benzene is reduced by concentrated hydriodic acid at 250°, cyclohexane, C_eH₁₂, is not produced as was hoped, but its isomer, methyl-pentamethylene, boiling at 69°, is formed by a molecular rearrangement.⁵² However, this method of reduction has been successfully used with toluene and m.xylene which give certain amounts of the corresponding saturated cyclic compounds. But this formation is very difficult and most of the aromatic hydrocarbons can not be hydrogenated in this way. The hydroaromatic hydrocarbons might be separated from Baku petroleum by laborious fractionations or prepared by tedious synthetic processes.

The direct hydrogenation of phenol and of its homologs had never been accomplished, nor had that of aniline and related aromatic amines.

On the contrary, benzoic and the phthalic acids had been hydrogenated by sodium amalgam.

445. In 1900, Lunge and Akunoff showed that combination takes place when a mixture of benzene vapor and hydrogen is passed over platinum black in the cold, or better, at 100°, and calculated from the decrease in volume of the mixture that cyclohexane, C₆H₁₂, must have been formed although the same reaction with platinum sponge gave only cyclohexene, C₆H₁₀. But the activity of the catalyst was quickly exhausted, and they were not able to isolate any product of the hydrogenation.⁵⁴

The use of reduced nickel enables us to hydrogenate the aromatic nucleus regularly in most cases. This hydrogenation ordinarily takes place around 180° without isomerization and usually without side reactions, hence with good yields. This is without doubt the most important service rendered by reduced nickel.

446. Aromatic Hydrocarbons. The direct hydrogenation of benzene to cyclohexane, C₆H₁₂, takes place with nickel above 70°. Its speed increases with the temperature up to 170–190°, where it is rapid without any side reaction. Above that, and particularly above 300°, a part of the benzene is reduced to methane and carbon is deposited on the nickel.

Cyclohexane is sometimes obtained at once, but usually it contains some benzene which has escaped the reaction and which is more

⁵⁸ KISHNER, J. prakt. Chem. (2), 56, 364 (1897).

⁵⁴ Lungs and Akunoff, Zeit. anorgan. Chem., 24, 191 (1900).

abundant the more worn out the nickel is. Treatment with a mixture of 1 volume fuming nitric acid to 2 volumes concentrated sulphuric acid easily removes the benzene.⁵⁵

447. All of the homologs of benzene are hydrogenated over nickel at 150 to 180°, being transformed into the homologs of cyclohexane.

Below 250° the hydrogenation takes place without any complications with the methyl derivatives of benzene, toluene, ortho, meta, and para xylene, mesitylene and pseudocumene, the yields of the corresponding methyl cyclohexane derivatives being practically quantitative, though traces of the aromatic hydrocarbons remain. These may be readily eliminated by shaking with the nitric-sulphuric acid mixture which has little effect on the saturated hydrocarbons in the cold.

448. But if we start with substituted benzenes containing long side chains, ethyl, propyl, isopropyl, and butyl, while the corresponding derivative of cyclohexane is always the chief product, there is always more or less of the saturated hydrocarbon resulting from the shortening of the long side chain. Thus ethyl-benzene gives, along with ethyl-cyclohexane, a little methyl-cyclohexane with correlative formation of methane. Propyl-benzene gives a little ethyl- and methyl-cyclohexane. This disturbance is more serious when the long side chain is a branched one, e. g., isopropyl. Thus with p.cymene which is p.methyl-isopropyl-benzene, along with about 66% of p.methyl-isopropyl-cyclohexane, about 16% each of p.dimethyl- and p.methyl-ethyl-cyclohexane are obtained.

This formation of by-products which is due to the power that the nickel has of dissociating the molecules, is greater with higher temperature, and for that reason it is best not to go above 180°.

- 449. By this method, methyl-cyclohexane, the three dimethyl-cyclohexanes, 1,3,5— and 1,3,4—trimethyl-cyclohexanes, propyl-cyclohexane, p.methyl-ethyl-cyclohexane, isopropyl-cyclohexane, the three methyl-isopropyl-cyclohexanes or menthanes, and dimethyl-isobutyl-cyclohexane have been prepared.⁵⁶
- 450. Above 250°, and particularly above 300°, the production of the cyclohexane hydrocarbons diminishes and then disappears altogether since the inverse dehydrogenation begins and becomes more and more rapid (641).
 - 451. Phenyl-ethylene, styrene, or cinnamine, C₆H₅. CH: CH₂, is

⁵⁵ SABATIER and SENDERENS, Compt. rend., 132, 210 (1901).

⁵⁶ SABATIER and SENDERENS, Compt. rend., 132, 566 and 1254 (1901). SABATIER and MURAT, Compt. rend., 156, 184 (1913), and Ann. Chim. (9), 4, 271 (1915).

hydrogenated at 160° by an active nickel to ethyl-cyclohexane. With a slightly active nickel around 200° hardly anything but ethylbenzene is obtained.⁵⁷

Phenyl-acetylene, C_eH₅. C: CH, hydrogenated over nickel at 180°, gives almost exclusively ethyl-cyclohexane.⁵⁸

452. Polycyclic Aromatic Hydrocarbons. Hydrogenation over active nickel at about 170° permits the addition of 6 atoms of hydrogen to each aromatic nucleus. The low volatility of the polyphenyl hydrocarbons, which do not boil except at temperatures above 250°, makes it necessary to carry their vapors along by a large excess of hydrogen. A single passage over the nickel under the conditions used for benzene does not effect complete hydrogenation and it is usually necessary to repeat the process with the product.

However, a single operation is all that is required to transform diphenyl-methane, C₆H₅. CH₂. C₆H₅, into dicyclohexyl-methane, C₆H₁₁. CH₂. C₆H₁₁. S⁵⁰

With diphenyl, C_0H_5 . C_0H_5 , Eijkman ⁵⁰ obtained only phenyl-cyclohexane, C_0H_5 . C_0H_{11} , boiling at 240°, but Sabatier and Murat have succeeded in transforming it into dicyclohexyl, C_0H_{11} . C_0H_{11} , melting at 4° and boiling at 233° and almost unattacked by the mixture of nitric and sulphuric acids. ⁶⁰

Likewise symmetrical diphenyl-ethane, or dibenzyl, C₆H₅. CH₂.-CH₂.C₆H₅, has been completely transformed into 1,2-dicyclohexylethane, C₆H₁₁. CH₂. CH₂. C₆H₁₁, boiling at 270°. The 1,1-diphenylethane, (C₆H₅)₂CH. CH₃, is changed with greater difficulty into the 1,1-dicyclohexyl-ethane boiling at 256°.

The four diphenyl-propanes are more or less readily transformed into the dicyclohexyl-propanes over nickel at around 170°.

Only in the case of dimethyl-diphenyl-methane, $(C_6H_5)_2C(CH_8)_2$, a quaternary hydrocarbon, is there any notable breaking up of the molecule into isopropyl-cyclohexane, ethyl-cyclohexane, methyl-cyclohexane and even cyclohexane.

Five diphenyl-butanes have been easily hydrogenated over active nickel at 170°, to the corresponding dicyclohexyl-butanes, and like results have been obtained with three diphenyl-pentanes which do not boil below about 300°.51

453. According to whether the temperature is higher or lower, triphenyl-methane, CH(C₆H₅)₈, gives first dicyclohexyl-phenyl-

- 57 SABATTER and SENDERENS, Compt. rend., 132, 1255 (1901).
- 58 SABATIER and SENDERENS, Compt. rend., 135, 88 (1902).
- ⁵⁹ EIJEMAN, Chem. Weekblad, 1, 7 (1903), C., 1903 (2), 989.
- 60 SABATIER and MURAT, Compt. rend., 154, 1390 (1912).
- 61 SABATIER and MURAT, Ann. Chim. (9), 4, 303 (1915).

methane, C_6H_5 . $CH(C_6H_{11})_2$, and then tricyclohexyl-methane, $CH(C_6H_{11})_3$.

On the contrary, the hydrogenation of symmetrical tetraphenylethane, $(C_0H_5)_2CH \cdot CH(C_0H_5)_2$, has miscarried, since under the influence of very active nickel at 230–240°, it yields only dicyclohexylmethane produced by the hydrogenation of the two halves of the molecule.⁶⁸

454. Hydrindene, which can be regarded as benzene with a satu-

to form dicyclononane, C₂H₁₆, boiling at 163°.64

decahydro-fluorene boiling at 258°.65

455. Aromatic Ketones. With an active nickel at a moderate temperature, the -CO- group is changed to -CH₂- and the aromatic rings are hydrogenated (389).

Thus acetophenone gives ethyl-cyclohexane.

Dibenzyl-ketone, C₆H₅. CH₂. CO. CH₂. C₆H₅, with active nickel at 175°, can give immediately symmetrical dicyclohexyl-propane, C₆H₁₁. CH₂. CH₂. CH₃. C₆H₁₁.

456. Phenols. The direct hydrogenation of the aromatic nucleus can be readily accomplished in phenol and its homologs by the use of nickel.

Phenol, hydrogenated at 180°, gives immediately cyclohexanol, C₆H₁₁. OH, containing 5 to 10% unchanged phenol, small quantities cyclohexanone and cyclohexene, C₆H₁₀. The mixture boiling between 155 and 165° can be purified by a second passage over the nickel at 150–170° which changes the phenol and cyclohexanone completely into cyclohexanol.⁶⁷

457. o.Cresol is regularly transformed by nickel at 200-220° into o.methyl-cyclohexanol with a yield of better than 90%. There is a little of the ketone which can be extracted with sodium bisulphite.

m. Cresol, under the same conditions, gives a mixture of the alcohol

- 62 Godchot, Compt. rend., 147, 1057 (1908).
- 68 Sabather and Murat, Compt. rend., 157, 1497 (1913).
- 64 EIJKMAN, Chem. Weekblad, 1, 7 (1903), C., 1903 (2), 989.
- 65 SCHMIDT and METZGER, Berichte, 40, 4566 (1907).
- 86 SABATIER and MURAT, Compt. rend., 155, 385 (1912).
- 67 SABATUR and SENDERENS, Compt. rend., 137, 1025 (1903).

and ketone which can be rehydrogenated at 180° to give practically pure m.methyl-cyclohexanol.

p.Cresol is readily hydrogenated at 200-230° to form p.methyl-cyclohexanol containing only traces of the ketone which are readily eliminated by bisulphite.⁶⁸

458. The xylenols, or dimethyl-phenols, are hydrogenated over nickel with varying degrees of success. 1,3-Dimethyl-phenol(4), at 190-200°, changes almost completely to the corresponding dimethyl-cyclohexanol with a little ketone and m.xylene.

The same may be said of the 1,4-dimethyl-phenol(2), which gives the corresponding cyclohexanol with about 10% of the ketone.

- 1,2-Dimethyl-phenol(4), hydrogenated under the same conditions, gives only about 25% of the desired cyclohexanol, about 8% of the ketone, while about 67% is reduced to o.xylene.⁶⁹
- 459. In the same manner with an active nickel at below 160° the regular hydrogenation of p.butyl-phenol, methyl-butyl-phenol, one dimethyl-butyl-phenol, one diethyl phenol 11 have been hydrogenated into the corresponding cyclo-aliphatic alcohols.

Thymol is satisfactorily hydrogenated to hexahydrothymol at 180-185°.

The same may be said of its isomer carvacrol, which is hydrogenated at 195-200° to hexahydrocarvacrol. 72

- 460. Polyphenols. The addition of 6H to the nucleus in polyphenols is difficult to realize by the use of nickel, doubtless because the desired reaction can be effected only between narrow temperature limits. At 200° the hydrogenation leads to phenol and benzene, mixed with cyclohexanol and cyclohexane, without any appreciable amount of the desired cycloaliphatic diols or triols.⁷³
- 461. On the contrary, by lowering the temperature to around 130°, the normal addition of hydrogen can be accomplished in some cases.

Hydroquinone at 130°, gives exclusively cyclohexadiol (1,4), or quinite, as the cis form, but if the hydrogenation is carried on at 160°, a mixture of the cis and trans forms is obtained with some phenol and cyclohexanol.

Pyrocatechin, at 130°, gives exclusively cyclohexadiol(1,2), melting at 75°.

- 68 SABATTER and MAILHE, Compt. rend., 140, 350 (1905).
- 69 SABATIER and MAILHE, Compt. rend., 142, 553 (1906).
- ⁷⁰ Darzens and Rost, Compt. rend., 152, 607 (1911).
- 71 HENDERSON and BOYD, J. Chem. Soc., 99, 2159 (1911).
- 72 BRUNEL, Compt. rend., 137, 1268 (1903).
- 78 SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 428 (1905).

Resorcine is difficult to hydrogenate at low temperatures on account of its slight volatility, but small amounts of cyclohexadiol(1,3), melting at 65°, have been isolated.⁷⁴

- 462. Pyrogallol, at 120-130°, gives cyclohexatriol(1,2,3), melting at 67°.74
- 463. Thymoquinol, C₂H₇(CH₂)C₆H₂(OH)₂, is hydrogenated by nickel to menthane-diol(2,5), melting at 112°.75
- 464. Ethers of Phenol. By means of nickel, below 150°, we may accomplish the direct hydrogenation of phenol ethers without breaking up the molecules.

Thus anisol, C_0H_0 . OCH₀, gives methoxy-cyclohexane, C_0H_{11} . OCH₀. The methyl ethers of the cresols are transformed into the corresponding methoxy-methyl-cyclohexanols. Phenetol gives ethoxy-cyclohexanol.⁷⁶

- 465. Aromatic Alcohols. Up to the present, the hydrogenation over nickel has not been accomplished without eliminating the hydroxyl group. Thus benzyl alcohol gives toluene and methyl-cyclohexane.
- p.Tolyl-dimethyl-carbinol, CH₂. C₆H₄. C(OH) (CH₂)₂, changes to hexahydrocymene, identical with menthane, when hydrogenated at 150°."
- 466. Aromatic Amines. On hydrogenating aniline at 190°, ammonia is evolved and a nearly colorless liquid with strong ammoniacal odor is obtained which gives on fractionation:
 - 1. A little benzene and cyclohexane, going over around 80°,
 - 2. About 30% of cyclohexyl-amine, C₆H₁₁. NH₂, boiling at 134°,
 - 3. A small amount of unchanged aniline, boiling at 182°,
- 4. A portion boiling above 190°, consisting of about 30% dicyclohexyl-amine, boiling at 252°, and about 30% of cyclohexyl-aniline, boiling at 279°, and a little diphenyl-amine, boiling at 311°.

The dicyclohexyl-amine comes from the decomposition of the cyclohexyl-amine by the nickel, with elimination of ammonia, similar to what has been mentioned in connection with the hydrogenation of nitriles (426). The cyclohexyl-aniline and the diphenyl-amine can be regarded as produced by the partial dehydrogenation of the dicyclohexyl-amine.⁷⁸

- 74 Sabatuse and Mailies, Compt. rend., 146, 1193 (1908).
- 75 HENDERSON and SUTHERLAND, J. Chem. Soc., 97, 1616 (1910).
- ⁷⁶ BRUNEL, Ann. Chim. Phys. (8), 6, 205 (1905). SABATIER and SENDERENS, Bull. Soc. Chim. (3), 33, 616 (1905).
 - ¹⁷ SMIRNOV, J. Russian Phys. Chem. Soc., 41, 1374 (1909).
 - 78 SARATIER and SENDERENS, Compt, rend., 138, 457 (1906).

467. The toluidines, CH₂. C₆H₄. NH₂, are more difficult to hydrogenate than aniline, but appear to give similar results.

By operating with m.toluidine (boiling at 197°) over nickel at 200°, we obtain, along with a little methyl-cyclohexane, boiling at 101°, and unchanged m.toluidine, a considerable amount of methocyclohexyl-amine, CH₃. C₆H₁₀. NH₂, boiling around 150° and having an intensely alkaline reaction, and higher alkaline products boiling at 145 and 175° respectively under 20 mm. pressure, which are doubtless dimethocyclohexyl-amine and methocyclohexyl-aniline. But the activity of the nickel falls off rapidly to nothing. This effect is even more marked with ortho and para toluidines and with the xylidines, whether these amines contain toxic substances or whether slightly volatile products of the reaction remain on the surface of the nickel and suppress its activity.⁷⁹

468. The hydrogenation of the nucleus by nickel at 160-180° is more readily accomplished for anilines substituted in the NH₂- group. The most difficult of these is *methyl-aniline* which gives a rather moderate yield of cyclohexyl-methyl-amine. A secondary reaction, which becomes more and more important as the temperature is raised, tends to produce the aliphatic amine, with the simultaneous liberation of cyclohexane or benzene.

Much more satisfactory results are obtained with ethyl-aniline, which gives cyclohexyl-ethyl-amine, boiling at 164°, with dimethyl-aniline, which leads to cyclohexyl-dimethyl-amine, boiling at 165°, and with diethyl-aniline which yields cyclohexyl-diethyl-amine, boiling at 193°. ••

469. Diphenyl-amine, (C_eH₅)₂NH, when submitted to hydrogenation over nickel at 250°, is decomposed into ammonia and cyclohexane. But by working at 190-210° with vapors of diphenyl-amine carried along by a large excess of hydrogen, it is possible to accomplish a regular hydrogenation, producing cyclohexyl-aniline and dicyclohexyl-amine, accompanied by certain amounts of cyclohexane, cyclohexyl-amine, and even aniline, resulting from the breaking up of the molecule by nickel.⁵¹

470. Benzyl-amine, such as is usually obtained by various methods of preparation, can not be hydrogenated over nickel without breaking up of the molecule into ammonia and toluene, even below 100°. The cause must be the presence of foreign substances which injure the catalyst, since the normal hydrogenation can be realized with

⁷⁹ SABATTER and SENDERENS, Ann. Chim. Phys. (8), 4, 387 (1905).

⁸⁰ SABATIER and SENDERENS, Compt. rend., 138, 1257 (1904).

⁸¹ SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 483 (1905).

benzyl-amine obtained by the catalytic action of thoria on a mixture of ammonia and benzyl alcohol vapors, and hexahydrobenzyl-amine is obtained, accompanied by dihexahydrobenzyl-amine.⁸²

471. Aromatic Acids. Direct hydrogenation over nickel fails when it is applied to benzoic acid or its homologs. When the vapors of benzoic acid, carried along by an excess of hydrogen, are passed over a very active nickel at 180-200°, the production of a little cyclohexane and traces of hexahydrobenzoic acid is observed at the start, but after a very short time the benzoic acid passes on unchanged, the surface of the nickel having doubtless become coated with a stable benzoate.*

Sabatier and Senderens failed likewise in the hydrogenation of the esters of benzoic acid, as the nickel rapidly became inactive. But by operating with the metal block at a perfectly regulated temperature below 170°, Sabatier and Murat have succeeded in accomplishing the regular hydrogenation of methyl benzoate, and even more readily, the hydrogenation of the esters of higher alcohols, and have thus obtained methyl, ethyl, isobutyl, and isoamyl hexahydrobenzoates, the isoamyl ester in 80% yield. The saponification of these esters yields the hexahydrobenzoic acid immediately.*

By the same method, they realized the complete hydrogenation of esters of phenyl-acetic acid to those of cyclohexyl-acetic at 170–185°, so of esters of hydrocinnamic acid to esters of β -cyclohexyl-propionic, so and finally of esters of ortho, meta, and para toluic acids to those of the corresponding hexahydrotoluic acids.

8. Various Ring Compounds

472. Trimethylene Ring. Cyclopropane, or trimethylene, CH₂. CH₂, is hydrogenated by nickel above 80°, and rapidly at CH₂. to form propane.³⁷

Likewise ethyl-trimethylene is hydrogenated by nickel to isopentane:

$$\begin{array}{c} \mathrm{CH_{5}} \\ \cdot \\ \mathrm{CH_{2}}.\mathrm{CH_{2}}.\mathrm{CH_{2}} \\ \end{array} \rightarrow \begin{array}{c} \mathrm{CH_{5}} \\ \mathrm{CH_{2}}.\mathrm{CH_{3}}.\mathrm{CH_{3}}.\mathrm{SS} \\ \end{array}$$

- 82 SABATHER and MAILHE, Compt. rend., 153, 160 (1911).
- 88 SABATIER and MURAT, Compt. rend., 154, 923 (1912).
- 84 SABATIER and MURAT, Compt. rend., 154, 924 (1912).
- 85 SABATIER and MURAT, Compt. rend., 156, 424 (1913).
- 86 SABATIER and MURAT, Compt. rend., 156, 751 (1913).
- 87 WILLSTÄTTER and KAMETAKA, Berichte, 41, 1480 (1908).
- 88 Rozanov, J. Russian Phys. Chem. Soc., 48, 168 (1916), C. A., 11, 454.

Methyl-cyclopropene yields isobutane at 170-180°: **

$$\begin{array}{c} CH \\ \cdot \\ CH_{\bullet} \end{array} C.CH_{\bullet} \rightarrow \begin{array}{c} CH_{\bullet} \\ CH_{\bullet} \end{array} CH.CH_{\bullet}.$$

Dimethylmethylene-cyclo-propane gives isohexane at 160°: **

$$\begin{array}{c} \mathrm{CH_2} \\ \cdot \\ \mathrm{CH_2} \end{array} \leftarrow \mathrm{C} : \mathrm{C}(\mathrm{CH_2})_2 \ \rightarrow \ \mathrm{CH_2} \cdot \mathrm{CH_2}\mathrm{CH_2}\mathrm{CH}(\mathrm{CH_3})_2$$

473. Tetramethylene Ring. Cyclobutane furnishes butane, while cyclobutene, at 180°, passes first into cyclobutane and then into butane.⁹¹

474. Pentamethylene Ring. Cyclopentadiene is regularly hydrogenated to cyclopentane. 92

475. Hexamethylene Ring. Cyclohexene, C₆H₁₀, is readily reduced to the cyclohexane condition by nickel below 180°. The same is true of the cyclohexadienes.

All the cyclohexene hydrocarbons are readily hydrogenated by nickel to the cyclohexane hydrocarbons. Thus the ethylene hydrocarbons formed from the three dimethyl-cyclohexanols readily furnish the three dimethyl-cyclohexanes. Methyl-ethyl-1,2-cyclohexene regularly passes into the corresponding saturated derivative. 4

Menthene, CH₂. C₂H₃. C₂H₇, submits to regular hydrogenation at 175° to give p.methyl-isopropyl-cyclohexane, or menthane, identical with that formed from cymene and accompanied by certain amounts of the same secondary products ⁹⁵ (448).

Phenyl-cyclohexene (1,1) is readily changed to phenyl-cyclohexane by a slightly active nickel. The same is true of cyclohexyl-cyclohexene (1,1), which furnishes dicyclohexyl.**

476. Acetyl-cyclohexane, CH₆. CO. C₆H₁₁, is hydrogenated by nickel at 160°, without affecting the ketone group, to give hexahydroacetophenone.⁹⁷

Ethyl tetrahydrobenzoate, C₆H₉.CO₂C₂H₅, is transformed into

- 89 MERRSHKOWSKI, J. Russian Phys. Chem. Soc., 46, 97 (1914), C. A., 8, 1965.
- 90 ZELINSKY, Berichte, 40, 4743 (1907).
- 91 WILLSTÄTTER and BRUCE, Berichte, 40, 4456 (1907).
- 92 EIJEMAN, Chem. Weekblad, 1, 7 (1903).
- 98 SABATIER and MAILHE, Ann. Chim. Phys. (8), 10, 552, 555 and 559 (1907).
- 94 MURAT, Bull. Soc. Chim. (4), 1, 774 (1907).
- 95 SABATIER and SENDERENS, Compt. rend., 132, 1256 (1901).
- 96 SABATHER and MURAT, Compt. rend., 154, 1390 (1912).
- 97 DARZENS and ROST, Compt. rend., 151, 758 (1910).

ethyl hexahydrobenzoate, and the ester of cyclohexene-acetic acid, C_aH₂. CH₂. CO₂H, into that of hexahydrophenyl-acetic acid.**

Carvone adds hydrogen to its double bond and its ketone group passes into the alcohol, forming a mixture of hydrocarvols.**

477. Terpenes. The terpenes with two double bonds add 2H₂ with nickel at 180°, while the terpenes with one double bond usually add only H₂.

Limonene gives menthane, identical with that from menthene and cymene with the same secondary products. The same is true of sulvestrene and terpinene.

Pinene is readily transformed at 170-180° into dihydropinene, C₁₀H₁₈, boiling at 166°, identical with that prepared by the action of hydroiodic acid (Berthelot).

The camphene (from an unknown source), melting at 41°, studied by Sabatier and Senderens, added H₂ with difficulty at 165–175° to furnish a camphane, C₁₀H₁₈, boiling at 164° and appearing to be identical with that which Berthelot had previously isolated.¹⁰⁰

The camphene from pinene hydrochloride gave a mixture of a solid *camphane*, melting at 65-67°, and liquid *camphane*, boiling at 160°. ¹⁰¹

An inactive camphene melting at 47-49° was transformed into a solid *camphane*, melting at 60°, by a single hydrogenation over nickel.¹⁰²

478. Terpineol, hydrogenated over nickel, even at a low temperature, around 125°, is changed to hexahydrocymene. 108

479. Heptamethylene Ring. Cycloheptadiene, C_7H_{10} , hydrogenated over nickel at 180°; yields only cycloheptane, stable even with prolonged hydrogenation at 200°, but at 235° it seems to isomerize into methyl-cyclohexane. 105

- 480. Octamethylene Ring. Cyclo-octadiene, C₈H₁₂, hydroge-
- 98 DARZENS, Compt. rend., 144, 328 (1907).
- 99 HALLER and MARTINE, Compt. rend., 140, 1302 (1905).
- 100 Sabatier and Senderens, Compt. rend., 132, 1256 (1901).
- ¹⁰¹ Lipp, Annalen, 382, 265 (1911).
- 102 NAMETRIN and MISS ABAUMOVSKAYA, J. Russian Phys. Chem. Soc., 47, 414 (1915), C. A., 10, 45.
 - 108 HALLER and MARTINE, Compt. rend., 140, 1393 (1905).
 - 104 ZELINSKY, J. Russian Phys. Chem. Soc., 36, 768 (1904).
 - 105 WILLSTÄTTER and KAMEKATA, Berichte, 41, 1480 (1908).

nated very slowly over nickel at 180°, gives cyclo-octane, C₈H₁₆,¹⁰⁶ which further hydrogenation at 200–250° appears simply to isomerize into dimethyl-cyclohexane.¹⁰⁷

Bicyclo-octene, at 150°, furnishes bicyclo-octane, boiling at 140°. 108

481. Naphthalene Nucleus. Naphthalene is transformed at 200° by nickel into tetrahydronaphthalene, 100 boiling at 205°, 110 while at 175°, decahydronaphthalene, or naphthane, boiling at 187°, is formed. 111

 α -Naphthol, by means of two successive hydrogenations at 170° and 135°, respectively, is transformed into decahydro- α -naphthol, melting at 62°.

Likewise by hydrogenation at 170° and then at 150°, β -naphthol yields decahydro- β -naphthol, melting at 75°. 113

in constitution, is transformed by nickel at 210°, as well as at 250°, into the tetrahydro-, C₁₂H₁₄, boiling at 254°. 118

- 483. Anthracene Nucleus. Anthracene is hydrogenated in steps, more hydrogen being taken up at lower temperatures. At 260° tetrahydroanthracene, C₁₄H₁₄, melting at 89°, is formed, while at 200°, octohydroanthracene, melting at 71°, is obtained. By using a very active recently prepared nickel, it is possible to transform the octohydro—into perhydroanthracene, C₁₄H₂₄, melting at 88°. 114
- 484. Phenanthrene Nucleus. Phenanthrene, C₁₄H₁₀, hydrogenated at 160° over a very active nickel, gave a mixture of the hexa-
 - 106 WILLSTÄTTER and VERAGUTH, Berichte, 40, 957 (1907).
 - 107 WILLSTÄTTER and WASER, Berichte, 44, 3444 (1911).
 - 108 WILLSTÄTTER and VERAGUTH, Berichte, 41, 1480 (1908).
- 100 The tetrahydro has d. 0.91520 and boils at 205-207° and is known as tetralin while the dekahydro is known as dekalin and has d. 0.882720 and boils at 189-191°. Tetralin spirits is a mixture of the two. These are coming to be important as turpentine substitutes, particularly in Europe. See DE KECHEL, Rev. chim. ind., 29, 173-178 (1920), C. A., 14, 3803; also Shecker, Annalen, 426, 1 (1922).— E. E. R.
 - 110 SABATIER and SENDERENS, Compt. rend., 132, 1257 (1901).
 - 111 LEBOUX, Compt. rend., 139, 672 (1904).
- ¹¹² LEBOUX, Compt. rend., 141, 953 (1905). Ann. Chim. Phys. (8), 21, 483 (1910).
- ¹¹⁸ SABATIER and SENDERENS, Compt. rend., 132, 1257 (1901). GODCHOT, Bull. Soc. Chim. (4), 3, 529 (1908).
 - 114 GODCHOT, Ann. Chim. Phys. (8), 12, 468 (1907).

hydro-, boiling at 305°, and the octohydro-, C₁₄H₁₈, boiling at 280°. ¹¹⁸ These results are different from those obtained by Schmidt and Metzger, who got only dihydrophenanthrene at 150°, ¹¹⁶ and from those of Padoa and Fabris, who obtained a mixture of the solid dihydro- and the liquid tetrahydro- at 200°, but were able to get the dodecahydro- at 175°. ¹¹⁷

485. Complex Rings. Pyrrol, when hydrogenated over nickel at 180-190°, gives 25% of pyrrolidine, C₄H₂N, with a small quantity of a substance which appears to be hexahydro-indoline.¹¹⁸

486. Pyridine is only slowly attacked by hydrogenation over nickel between 120 and 220°, and does not yield any piperidine; there is opening of the ring with the formation of some amyl-amine.¹¹⁸

487. Furfuryl-ethyl-carbinol yields tetrahydrofurfuryl-ethyl-carbinol on hydrogenation at 175°. 120

Methyl-a-fufurane adds 2H, at 190° to give tetrahydro-methyl-

$$\alpha$$
-furfurane, $CH_2 \cdot CH_2 \cdot CH_2 \cdot CH_3 \cdot CH_4 \cdot CH_4 \cdot CH_5 \cdot$

the ring is opened and methyl-propyl-ketone is formed, finally methyl-propyl-carbinol, or pentanol(2).¹²¹

488. Quinoline, when hydrogenated over a very active nickel at 160-190°, adds 2H₂ to the pyridine ring to form tetrahydroquinoline in excellent yield.

Likewise 6-methyl-quinoline is readily hydrogenated to the corresponding methyl-tetrahydroquinoline. 122

By carrying out the hydrogenation at 130-140°, over a very active nickel, decahydroquinoline may be obtained. Likewise quinaldine furnishes decahydroquinaldine in excellent yield.¹²⁸

489. By hydrogenating quinoline at a higher temperature, the normal addition of hydrogen does not take place, but the ring is opened to yield *ethyl-o.toluidine*, which does not remain as such but closes the ring, with loss of hydrogen to give α -methyl-indol: ¹²⁴

```
<sup>115</sup> Breteau, Compt. rend., 140, 942 (1905).
```

¹¹⁶ SCHMIDT and METZGER, Berichte, 40, 4240 (1907).

¹¹⁷ Padoa and Fabris, Gaz. Chim. Ital., 39 (1), 333 (1909).

¹¹⁸ Padoa, Gaz. Chim. Ital., 36 (2), 317 (1906).

¹¹⁹ SABATIER and MAILHE, Compt. rend., 144, 784 (1907).

¹²⁰ Douris, Compt. rend., 157, 722 (1913).

¹²¹ PADOA and PONTI, Lincei, 15 (2), 610 (1906), C., 1907 (1), 570.

¹²² DARZENS, Compt. rend., 149, 1001 (1909).

¹²⁸ SABATIER and MURAT, Compt. rend., 158, 309 (1914).

¹²⁴ PADOA and CARUGHI, Lincei, 15, 113 (1906), C., 1906 (2), 1011.

490. Carbazol, diphenyl-imide, when hydrogenated over nickel at 200° under 10 atmospheres pressure. gives $\alpha\beta$ -dimethyl-indol: 125

491. Acridine is slowly hydrogenated over nickel at 250-270° to αβ-dimethyl-quinoline: 126

g. Carbon Disulphide

492. When carbon disulphide vapors are carried by an excess of hydrogen over nickel at 180°, a volatile, extremely ill-smelling substance is produced which gives a yellow mercury salt, a white cadmium salt, and brown lead and copper salts, and which appears to be methylene-dithiol, H₂C(SH)...¹²⁷

HYDROGENATIONS WITH DECOMPOSITIONS

493. Catalytic nickel quite frequently exercises a more or less intense decomposing action on the molecules: in such cases not only the original compound but also the fragments resulting from its decomposition are hydrogenated.

Hydrocarbons. We shall study in Chapter XXI the decompositions that hydrocarbons undergo at high temperatures in the presence of nickel and other catalysts. The study of the simultaneous hydrogenations can not be separated from that of the decompositions and molecular condensations resulting therefrom.

- 125 Padoa and Chiaves, Lincei, 16 (2), 762 (1907), C., 1908 (1), 649.
- 126 Padoa and Fabris, Lincei, 16 (1), 921 (1907), C., 1907 (2), 612.
- 127 SABATTER and Espil, Bull. Soc. Chim. (4), 15, 228 (1914).

494. Aliphatic and Aromatic Ethers. Aliphatic ethers resist hydrogenation over nickel quite well, but when it is carried out above 250°, there is decomposition into hydrocarbon and alcohol which is then attacked, furnishing the products of the hydrogenation of its debris.

Thus ethyl ether gives ethane and alcohol which gives the fragments of acetaldehyde, of which the carbon monoxide is partly changed to methane: 128

then

$$(C_2H_5)_2O + H_2 - C_2H_6 + CH_3 \cdot CH_2OH$$

 $CH_3 \cdot CH_2OH - CH_4 + CO + H_2$
 $CO + 3H_2 - CH_4 + H_2O$.

Aromatic ethers undergo an analogous decomposition with nickel, this taking place at moderate temperatures with the mixed alkyl phenyl ethers and greatly diminishing the yields of the mixed alkylcyclo-aliphatic ethers which are made by their hydrogenation.

In the hydrogenation of anisol to methoxy-cyclohexane (464), there is the production of certain amounts of methyl alcohol and cyclohexane. 128 If the operation is carried on above 300°, there is no hydrogenation of the nucleus and scission is rapid in the same manner as with aliphatic ethers.

We have two reactions:

$$C_eH_s \cdot O \cdot R + H_2 = RH + C_eH_s \cdot OH$$

and

$$C_6H_8 \cdot O \cdot R + H_2 - C_6H_6 + R \cdot OH$$

the alcohol itself being more or less broken down by the hydrogenation.

This is the case with the *methyl* ethers of *phenol*, of the three *cresols*, of α -naphthol, etc., and also with *phenyl oxide* which is the most resistant to decomposition.¹²⁹

495. Phenyl Isocyanate. Phenyl isocyanate, when hydrogenated over nickel at 190°, breaks up into two portions which are hydrogenated separately:

$$C_6H_5$$
. $N : CO - CO + C_6H_5$. $N-$.

We obtain aniline and carbon monoxide which yields methane with the formation of water. This reacts quantitatively with the original compound to give carbon dioxide and solid diphenyl-urea. 120

- 128 SABATTER and SENDERENS, Bull. Soc. Chim. (3), 33, 616 (1905).
- 129 MAILHE and MURAT, Bull. Soc. Chim. (4), 11, 122 (1912).
- 180 SABATIER and MAILHE, Compt. rend., 144, 825 (1907).

496. Amines. Various amines hydrogenated over nickel at above 300-350°, tend to form ammonia and a hydrocarbon. This reaction which takes place readily with aliphatic amines has already been mentioned with aniline (378). It takes place with the homologs of aniline, with benzyl-amine and with the naphthyl-amines.

Hexamethylene-tetramine is completely decomposed yielding ammonia, trimethyl-amine and methane: 181

$$N(CH_2.N:CH_2)_2 + 9H_2 - N(CH_2)_2 + 3NH_2 + 3CH_4$$

497. Compounds Containing -N.N-. Phenylhydrazine, hydrogenated above 210°, is split into ammonia and aniline, accompanied by cyclohexyl-amine, dicyclohexyl-amine, and even by benzene and cyclohexane.¹⁸²

The main reaction is:

$$C_6H_5$$
. NH. NH₂ + H₂ - NH₂ + C_6H_5 . NH₂.

Azobenzene, C₆H₅. N: N. C₆H₅, hydrogenated at 290°, yields aniline chiefly.¹⁸²

Indol. On hydrogenation over nickel at 200°, indol is split into o.toluidine and methane: 188

$$C_{\bullet}H$$
 CH
 CH
 CH
 CH_{\bullet}
 CH_{\bullet}
 CH_{\bullet}
 CH_{\bullet}
 CH_{\bullet}
 CH_{\bullet}
 CH_{\bullet}

- 181 GRASSI, Gas. Chim. Ital., 36 (2), 505 (1906).
- 182 SABATIER and SENDERBNS, Bull. Soc. Chim. (3), 35, 259 (1906).
- 188 CARRASCO and PADOA, Lincei, 14 (2), 699 (1906), C., 1906 (2), 683.

CHAPTER X

HYDROGENATIONS (Continued)

HYDROGENATIONS IN GASEOUS SYSTEM (Continued)

I.—USE OF VARIOUS CATALYSTS

498. Nickel as a hydrogenation catalyst can be replaced by various finely divided metals, such as cobalt, iron, copper, platinum, and the platinum metals, particularly palladium.

Cobalt

499. Finely divided cobalt such as is produced by the reduction of the oxide in the hydrogenation tube itself, seems to be able to take the place of nickel in all the various reactions which nickel can catalyze.

But its use is disadvantageous because its activity is less and more easily destroyed than that of nickel; because higher temperatures are required when using it; and also because the reduction of its oxide is practicable only in the neighborhood of 400°, and hence the oxide resulting from spontaneous oxidation during the time the apparatus is cold and out of use, can not be reduced at temperatures below 250° such as are commonly used in hydrogenations.

500. Ethylene Hydrocarbons. When a mixture of ethylene and an excess of hydrogen is passed over cold reduced cobalt, immediate action takes place with the production of ethane, and the end of the cobalt layer becomes hot. The heated portion moves slowly along the metal and the evolution of heat finally ceases and the production of ethane stops also, doubtless because the cobalt is slightly carbonized in the course of the reaction and its activity so diminished that it is unable to continue the reaction without the aid of external heat.

At 150°, the hydrogenation of ethylene continues indefinitely, but the cobalt is slowly weakened, more rapidly than nickel.

Above 300°, the disturbance due to the action of the cobalt on the ethylene (910) appears and the issuing gases contain methane and carry along small amounts of liquid hydrocarbons.¹

¹ Sabatier and Senderens, Ann. Chim. Phys. (8), 4, 344 (1905).

The action of cobalt on the homologs of ethylene is similar to that of nickel but weaker.

- 501. Acetylene. Reduced cobalt, entirely free from nickel, can serve to hydrogenate acetylene, but there is no reaction in the cold. The fixation of hydrogen begins at about 180°, and the ethane produced is accompanied by a small amount of liquid hydrocarbons, which are more abundant if the reaction is carried on at 250°.
- 502. Benzene and its Homologs. Reduced cobalt can effect the direct hydrogenation of benzene and its homologs at 180°, but its activity falls off rather rapidly.
- 503. Aliphatic Aldehydes and Ketones. Cobalt can transform aliphatic aldehydes and ketones into the alcohols below 180°, but is less active than nickel. Under identical conditions, with the same apparatus, the same temperature, the same velocity of hydrogen, and the same rate of admission of acetone, the yield of isopropyl alcohol was about 83% with nickel as catalyst but slightly less than 50% with cobalt.
- 504. Carbon Monoxide and Dioxide. Reduced cobalt can cause the transformation of carbon monoxide into methane, as does nickel, but the reaction does not begin till about 270°. It is rapid at 300°, but is opposed more strongly, than is the case with nickel, by the decomposition of carbon monoxide into carbon and the dioxide (615). This decomposition is as rapid with cobalt as with nickel, while the hydrogenation is slower with the cobalt.

The hydrogenation of carbon dioxide is effected by cobalt from 300° up. It is rapid at 360° and even more so at 400° and is accomplished without any complications.

Iron

505. Finely divided iron, obtained by the reduction of its oxides, can be substituted for nickel as a hydrogenation catalyst in certain cases, but is less active than nickel and even less active than cobalt. Besides, it has the marked disadvantage of being much more difficult to prepare from its oxide. Between 400 and 500° it is necessary to continue the action of hydrogen from six to seven hours to obtain complete reduction. When the metal is reduced more rapidly at higher temperatures, it is no longer pyrophoric and has only slight activity.

- ² Sabatier and Senderens, Ann. Chim. Phys. (8), 4, 352 (1905).
- ³ Sabatier and Senderens, Ann. Chim. Phys. (8), 4, 368 (1905).
- 4 SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 400 and 403 (1905).
- ⁵ Sabatier and Senderens, Ann. Chim. Phys. (8), 4, 424 (1905).

506. Ethylene Hydrocarbons. Iron causes the hydrogenation of ethylene only above 180°, and its activity decreases with the slow carbonizing of the metal.

Acetylene. The hydrogenation of acetylene does not commence till above 180°, and always gives rise to the formation of rather large amounts of colored hydrocarbons, containing higher ethylene hydrocarbons soluble in sulphuric acid, aromatic hydrocarbons, and only a small amount of saturated hydrocarbons. The odor and appearance of the product suggest certain natural petroleums of Canada.

To a certain extent, iron can cause the hydrogenation of aldehydes, ketones and nitro compounds, but is incapable of transforming carbon monoxide and dioxide into methane or of adding hydrogen to the benzene nucleus.

Copper

507. Reduced copper is a useful catalyst for certain hydrogenations. For such its use is advantageous on account of its ease of preparation, the low temperature, below 180°, at which its oxide can be reduced, and its resistance to poisons which is more marked than with other metal catalysts.

508. Reduction of Carbon Dioxide. Copper, even in its most active form (59), is incapable of causing the direct hydrogenation of carbon monoxide to methane and does not show any action on mixtures of carbon monoxide and hydrogen below 450°.

It is the same way with mixtures of hydrogen and carbon dioxide below 300°, but between 350 and 400° a special reaction appears gradually and is quite definite at 420-450°. There is reduction of the carbon dioxide into carbon monoxide and water, according to the equation:

$$CO_{\bullet} + H_{\bullet} - CO + H_{\bullet}O.$$

Thus with a mixture of one part carbon dioxide to about three parts of hydrogen, a gas was obtained containing:

Carbon monoxide	10.0%	by	volume
Carbon dioxide	17.2%	"	"
Hydrogen	72.8%	"	"

More than a third of the carbon dioxide had been reduced to the monoxide. The proportion reduced is less when the concentration of hydrogen in the mixture is less.

⁶ Sabatter and Senderens. Ann. Chim. Phys. (8), 4, 345, 353, 368, 425, and 428 (1905).

In no case is even a trace of methane formed.

509. Nitro Compounds. Copper gives results analogous to those with nickel (373 to 378) only at higher temperatures.

Nitrous oxide is reduced to nitrogen at 180° and nitric oxide is changed into ammonia at the same temperature. Nitrogen peroxide gives copper nitride in the cold, and it is only towards 180° that ammonia is produced. If the proportion of nitrogen peroxide becomes too great, there is incandescence followed by an explosion.

510. Nitromethane, hydrogenated between 300 and 400°, gives, along with methyl-amine, a liquid of a more or less brown color with a disgusting odor in which appear crystals which are the methyl-amine salt of nitromethane.

Between 300 and 400°, nitro-ethane gives ethyl-amine without notable complications.¹⁰

511. Copper is the best of all the finely divided metals for transforming aromatic nitro derivatives into the amines, since its very regular hydrogenating action affects only the -NO₂ group and does not touch the aromatic nucleus. Nitrobenzene is thus changed to aniline from 230° up, the reaction being rapid and very regular between 300 and 400°, and so long as the hydrogen is in excess, aniline is obtained in 98% yield containing only traces of nitrobenzene and the red azobenzene. The same metal can be used for a long time. The hydrogen can, without inconvenience, be replaced by water gas, the carbon monoxide of which acts usefully as a reducing agent to some extent since a part of it is transformed into carbon dioxide. The manufacture carried out with copper, a metal which is not costly and which serves for a long time and is easily regenerated without loss, and by means of a very cheap gas, can be carried on continuously and is very economical.¹¹

Coppered pumice at 200-210° has been proposed as a substitute for copper.¹²

512. The manufacture of the toluidines from the nitrotoluenes is also advantageously carried on by copper at 300-400°, and likewise

- ⁷ SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 426 (1905).
- 8 SABATTER and SENDERENS, Ann. Chim. Phys. (7), 7, 401 (1896).
- * SABATTER and SENDERENS, Compt. rend., 135, 278 (1902).
- 19 SABATIER and SENDERENS, Compt. rend., 135, 227 (1902).
- ¹¹ Sabatier and Senderens, Compt. rend., 133, 321 (1901). Sabatier, Vth. Cong. Pure and Appl. Chem., Berlin, 1903, II, 617. Senderens, French Patent, 312,615 (1901).
- ¹⁵ Badisches, English patent, 6,409 of 1915. J. Soc. Chem. Ind., 35, 920 (1916).

 α -naphthyl-amine is readily obtained from α -nitronaphthalene at 330-350°. 12

The chlornitrobenzenes are regularly transformed by copper into the chloranilines at 360-380°. On the contrary, copper gives poor results with the dinitrobenzenes and the bromnitrobenzenes.¹⁴

At 265° the results are excellent with the nitrophenols and the nitronilines.18

- 513. Esters of Nitrous Acid. Nitrous esters are regularly hydrogenated into the amines, over copper as well as over nickel, but at a higher temperature, 330–350°, the results are satisfactory for nitrites with heavy hydrocarbon chains, but are less so for methyl nitrite which gives brown products analogous to those obtained from nitromethane.¹⁶
- 514. Oximes. Copper accomplishes the regular hydrogenation of aliphatic aldoximes and ketoximes between 200 and 300° into primary and secondary amines without complications, 17 and the same may be said about aliphatic amides. 18
- 515. Ethylene Compounds. Most often copper serves to add hydrogen to the ethylene double bond.

Ethylene, propylene and α -octene are changed to the corresponding saturated compounds at above 180°. However, trimethylethylene and β -hexene are not hydrogenated by copper, and it has been concluded that copper does not cause the hydrogenation of any except α -ethylene compounds, that is to say, those in which one of the CH₂ groups of the ethylene is not substituted.¹⁹

This limitation is not general since the vapors of oleic acid are readily hydrogenated into stearic acid at around 300°. Water gas can be substituted for the pure hydrogen in this preparation and it has industrial possibilities.²⁰

It may be noted that copper does not cause the hydrogenation of symmetrical diphenyl-ethylene, or stilbene, C_6H_5 . CH: CH. C_6H_5 , of cyclohexene, C_6H_{10} , or of the methyl-cyclohexenes.²¹

516. The use of copper, which acts on the ethylene double bond

- ¹³ SABATTER and SENDERENS, Compt. rend., 135, 225 (1902).
- ¹⁴ MIGNONAC, Bull. Soc. Chim., (4), 7, 154, 270 and 504 (1910).
- ¹⁵ Brown and Carrick, J. Amer. Chem. Soc., 41, 436 (1919).
- 16 GAUDION, Ann. Chim. Phys. (8), 25, 136 (1912).
- ¹⁷ Mailhe, Compt. rend., 140, 1691 (1905) and 141, 113 (1905).
- ¹⁸ MAILHE, Bull. Soc. Chim. (3), 35, 614 (1906).
- ¹⁹ Sabatier and Senderens, Compt. rend., 134, 1127 (1902).
- ²⁰ SABATIER, French patent, 394,957 (1907).
- ²¹ Sabatier, 50th. Cong. des Soc. Sav. (1912). Journ. Offic., 3628: April 11, 1912.

without attacking the aromatic nucleus, permits us to effect certain hydrogenations distinct from those obtained with nickel. *Phenylethylene*, or *styrene*, C₆H₈. CH: CH₂, which nickel changes into *ethyl-cyclohexane*, is transformed quantitatively at 180° by copper into *ethyl-benzene*.³²

changes into menthane (477), gives only dihydrolimonene, C₁₀H₁₈, isomeric with menthene.²²

518. Acetylene Hydrocarbons. Copper can not hydrogenate acetylene in the cold, the reaction being around 130° over copper with a light purple color and around 180° over copper of a clear red. Carried on with excess of hydrogen, the reaction always gives a certain proportion of liquid hydrocarbons along with the ethane.

When the amount of acetylene equals or surpasses the amount of hydrogen, the special condensing action of copper on acetylene (914) becomes evident: the copper swells up gradually on account of the formation of solid *cuprene*, $(C_7H_6)_x$ the gases evolved contain higher ethylene hydrocarbons and a mixture of liquid ethylene and aromatic hydrocarbons (benzene, and homologs and styrene) is collected.

A gas mixture containing 21 H₂ to 19 C₂H₂ gave, at 150° over violet copper, a condensation of materials containing 25 C with about 65% carbon, one third as cuprene and the other two thirds as liquid hydrocarbons.²³

519. The hydrogenation of α-heptine over copper at below 200°, gave a little heptane, but chiefly heptene, diheptene, and triheptene.²⁴

520. Phenyl acetylene, C_eH₅. C: CH, which nickel transforms easily into ethyl-cyclohexane (451), gave by hydrogenation over copper between 190 and 250°, ethyl-benzene, C_eH₅. CH₂. CH₂, accompanied by a little phenyl-ethylene and a nearly equal amount of symmetrical diphenyl-butane, C_eH₅. CH₂. CH₂. CH₂. CH₂. C_eH₅, a well crystallized solid.²⁵

521. Nitriles. Copper can transform nitriles into primary and secondary amines 26 in the same manner that nickel does. It acts

- * SABATIER and SENDERENS, Compt. rend., 132, 1255 (1901).
- ²⁸ SABATIER and SENDERENS, Compt. rend., 130, 1559 (1900).
- 24 SABATIER and SENDERENS, Compt. rend., 135, 87 (1902).
- SABATIER and SENDERENS, Compt. rend., 135, 88 (1902).
- ²⁶ Sabatter and Senderens, Compt. rend., 140, 482 (1905) and Bull. Soc. Chim. (3), 33, 371 (1905).

similarly on the carbyl-amines,²⁷ but its action is less rapid than that of nickel.

522. Aliphatic Aldehydes and Ketones. Below 200°, copper can transform these slowly into the alcohols, but the inverse action usually preponderates and this makes the use of copper less advantageous.

Furthermore, copper is incapable of transforming the oxides of carbon into methane or of hydrogenating the aromatic nucleus.

523. Aromatic Ketones. When benzophenone is hydrogenated at 350° over copper with a violet tint, prepared by the reduction of the hydroxide (59), diphenyl-methane is formed directly.²⁸

Platinum

524. Platinum black can be used for direct hydrogenation in quite a large number of cases and its activity is greater than that of copper though less than that of nickel. Its activity is greater, the more tenuous the black and the more recently it has been prepared. It is rapidly exhausted and this fact taken together with the high cost of the metal renders its use generally less advantageous.

Platinum moss, or sponge, behaves the same way but with less activity, which is usually not manifested except at a higher temperature.

525. Union of Carbon and Hydrogen. The presence of finely divided platinum on the carbon accelerates its direct combination with hydrogen to form *methane* at 1200°, the limit of the combination, 0.53%, not being altered.²⁹

526. Ethylene Compounds. A mixture of ethylene and hydrogen is transformed into ethane in the cold in the presence of platinum black.³⁰ But after some time the slight carbonization of the metal prevents the reaction from proceeding at the ordinary temperature and it is necessary to heat to 120°, or even to 180°, to obtain a rapid formation of ethane.³¹

Analogous results are obtained with propylene.

The vapors of amyl oleate can be hydrogenated over platinized asbestos to amyl stearate.³²

527. Acetylene Hydrocarbons. Acetylene combines with hydro-

- 27 SABATTER and MAILHE, Ann. Chim. Phys., (8) 16, 95 (1909).
- * SABATIER and MURAT, Compt. rend., 158, 761 (1914).
- 20 PRING, J. Chem. Soc., 97, 498 (1910).
- ⁸⁰ VON WILDE, Berichte, 7, 352 (1874).
- M SABATIER and SENDERENS, Compt. rend., 131, 40 (1900).
- * FOKIN, J. Russian Phys. Chem. Soc., 38, 419 (1906), C., 1906 (2), 758.

gen in the cold in the presence of platinum black, giving first ethylene and then ethane.²⁰

In presence of an excess of hydrogen, acetylene is entirely transformed into pure ethane without any side reactions.

At 180° the same reaction takes place more rapidly but there is the formation of a certain amount of higher liquid hydrocarbons. By augmenting the proportion of acetylene in the mixture, *ethylene* becomes the main product but some ethane is always formed even though unchanged acetylene remains.

If the proportion of acetylene becomes great enough, with the platinum black at 180°, a certain amount of smoky decomposition of the gas is observed and this ends with incandescence, as is the case with nickel (914).

Platinum sponge is not active in the cold and does not effect the hydrogenation of acetylene except above 180°.33

- 528. Hydrocyanic Acid. Platinum black can hydrogenate hydrocyanic acid to *methyl-amine* at 116°, but the cyanidation of the metal soon diminishes its activity and stops the reaction.³⁴
- 529. Nitro Compounds. Nitrogen oxides, either nitric oxide or the dioxide, are readily reduced to ammonia with the aid of platinum sponge which is thereby heated to incandescence.³⁵
- 530. Nitromethane is hydrogenated over platinum sponge at 300°, more slowly than over copper but with analogous results (510).³⁶
- 531. Various forms of platinum, black, sponge, and platinized asbestos, can cause the transformation of *nitrobenzene* into aniline, but their catalytic power is low and, if the hydrogen is not in large excess, there is incomplete reduction with the formation of crystallized hydrazobenzene.⁸⁷
- 532. Aliphatic Aldehydes and Ketones. Finely divided platinum is unsuitable for the regular transformation of these into the alcohols, since at the temperatures which must be used, which are above 200°, the metal acts powerfully to break up the aldehyde molecule into carbon monoxide and hydrocarbon (622).
- 533. Finely divided platinum, even in the form of highly active black, has proved powerless to effect the direct hydrogenation of carbon monoxide or dioxide to methane. There is no action even up to 450°.**
 - * SABATIER and SENDERENS, Compt. rend., 131, 40 (1900).
 - ²⁴ DEBUS, J. Chem. Soc., 16, 249 (1863).
 - * KUHLMANN, Compt. rend., 7, 1107 (1838).
 - ³⁴ Sabatier and Senderens, Compt. rend., 135, 226 (1902).
 - 87 SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 414 (1905).
 - * SABATIER and SENDERENS, Compt. rend., 134, 514 and 689 (1902).

534. Aromatic Nucleus. Recently prepared platinum black can transform benzene into cyclohexane at 180° for a time, but its activity diminishes rapidly and soon disappears.

Platinum sponge has not this power.89

According to Zelinsky, platinum is as well able to hydrogenate benzene, toluene, the three xylenes and ethyl-benzene, as is nickel.40

He states the same about palladium.

535. Polymethylene Rings. Spirocyclane, with the aid of platinum, first adds H₂ to form ethyl trimethylene, which passes to pentane by a second addition: ⁴¹

Cyclo-octatetrene adds 4H₂ with the aid of platinum sponge to form cyclooctane.⁴²

Palladium

536. Palladium, previously charged with hydrogen, is able to effect varied hydrogenations, such as the transformation of nitrobenzene into aniline, nitromethane into methyl-amine, and nitrophenols into aminophenols (Graham). It is easy to foresee that it can serve equally well as a hydrogenation catalyst, the intermediate hydride which enables it to accomplish these results being notably stable in this case.

The formation of aniline by the action of hydrogen on nitrobenzene in the presence of palladium was shown by Saytzeff.⁴²

Carbon monoxide can be reduced in the cold, or better, at 400°, to methane in the presence of palladium sponge.44

Phenanthrene, carried over palladium sponge at 150-160° by a current of hydrogen, gives a mixture of tetrahydro- and octohydro-phenanthrene.⁴⁵

Unfortunately the excessive price of palladium restricts its useful applications.

- 88 SABATTER and SENDERENS, Ann. Chim. Phys. (8), 4, 368 (1905).
- 40 ZELINSKY, J. Russian Phys. Chem. Soc., 44, 274 (1912).
- 41 ZELINSKY, J. Russian Phys. Chem. Soc., 44, 275 (1912).
- 4 WILLSTÄTTER and WASER, Berichte, 44, 3423 (1911).
- KOLBE and SAYTZEFF, J. prakt. Chem. (2), 4, 418 (1871).
- 44 BRETEAU, Etude sur les meth. d'hydrogenation, 1911, p. 22.
- 46 BRETEAU, Ibid., p. 24.

II. - HYDROGENATION BY NASCENT HYDROGEN

537. Certain catalyses yield hydrogen and the gas so produced can be immediately employed for hydrogenation purposes. We can thus use as sources of active hydrogen, alcohol vapors, formic acid, and even a mixture of water and carbon monoxide.

Hydrogenation by Alcohol Vapors

538. Primary and secondary alcohols can, under the influence of various catalysts, be decomposed into aldehydes and ketones and hydrogen (653): the hydrogen thus set free can act in the nascent state on substances the vapors of which are mixed with the alcohols.

Copper can easily realize such reactions, but we can attribute to its action the hydrogenation correlative to the decomposition.

We can use mixed oxide catalysts (675) and even dehydrating catalysts, such as thoria, the presence of the substance that can be hydrogenated orienting the decomposition of the alcohol in the direction of the separation of hydrogen and greatly diminishing the extent of the dehydration reaction.

Thus over thoria at 420°, benzhydrol, with ethyl alcohol in excess, gives much diphenyl-methane accompanied by a little benzophenone and tetraphenyl-ethane (720): acetaldehyde is evolved and the gases arising from its decomposition.

The alcohol most suitable for this sort of hydrogenation is methyl alcohol on account of its great tendency to produce formaldehyde and particularly the products of its decomposition, carbon monoxide and hydrogen (693):

$$H.CH.OH = 2H. + CO.$$

The vapors of the substance to be hydrogenated are passed over thoria at 420°, with an excess of methyl alcohol, the hydrogenation is advantageously accomplished in all cases in which the product is stable at that temperature. Thus benzophenone and benzhydrol are changed almost completely into diphenyl-methane, while benzyl alcohol and benzaldehyde give toluene, acetophenone furnishes ethylbenzene, and nitrobenzene yields aniline.⁴⁶

SARATIER and MURAT, Compt. rend., 157, 1499 (1913). — Bull. Soc. Chim. (4), 15, 227 (1914).

⁴⁷ By using 2.5 moles of ethyl alcohol to 1 of bensaldehyde, and passing the mixed vapors over ceria on asbestos at 300-300°, bensyl alcohol is obtained along with acetaldehyde. Similarly citronellol is formed from citronellal and phenylethyl alcohol from phenylacetaldehyde. The yields are variable and the catalyst is rapidly fouled, probably on account of the formation of condensation products of the aldehydes either alone or with each other. See article by Milligan and myself, Jour. Amer. Chem. Soc., 44, 202 (1922).— E. E. R.

Hydrogenation by the Vapors of Formic Acid

539. The vapors of *formic acid* passing over various catalysts, finely divided platinum, copper or nickel reduced from their oxides, cadmium, stannous oxide or zinc oxide, are decomposed below 300° into carbon dioxide and hydrogen (824):

$$HCO_2H - H_2 + CO_2$$

This hydrogen can be used to hydrogenate substances the vapors of which are present in the system. Under these conditions, using nickel at 300°, acetophenone is changed to ethyl-benzene, phenylethyl-ketone into propyl-benzene, and benzophenone into diphenylmethane.

Thoria, alumina and zirconia effect the same hydrogenations above 300°, but the oxides of manganese appear to be inactive. 48

Hydrogenation by the Mixture of Carbon Monoxide and Water

540. The mixture $CO + H_2O$ can be transformed into $CO_2 + H_2$, the reaction being favored by the temporary combination of the carbon dioxide with the catalyst or by the immediate utilization, thanks to the catalyst, of the hydrogen to hydrogenate the carbon monoxide into methane. The reaction then becomes:

$$4CO + 2H_2O - 3CO_2 + CH_4$$

It is found, in fact, that a mixture of steam and carbon monoxide passing over *lime* at above 1000° gives the above reaction and we have the following reaction at the same time:

$$CO + H_2O - CO_2 + H_2$$

As calcium carbonate is entirely decomposed at this temperature, the lime acts as a true catalyst. By separating the carbon dioxide, we can obtain a mixture containing:

Hydrogen	 88%	by	volume
Methane 49	 12%		

The same mixture passing over iron wool likewise gives methane in varying amounts:

At	250°	 7.3%	by	volume
At	950°	 11.2%		
At	1250°	 7.1%		

By the use of fine nickel turnings a maximum content of 12.5% of

- 44 MAILHE and DE GODON, Bull. Soc. Chim. (4),"21, 61 (1917).
- 49 VIGNON, Compt., rend., 156, 1995 (1913).

methane is obtained at 400°. With copper turnings, almost no result is obtained at 500°, and the maximum, 6.3%, is obtained at 700°.

Precipitated silica gives a maximum of 8.4% at 700°, while for alumina, obtained by calcining the hydroxide, the maximum, 3.8%, is obtained at 950°, and for magnesia, a maximum of 6.7% at 900°. 50

⁵⁰ Vignon, Compt. rend., 157, 131 (1913). — Bull. Soc. Chim. (4), 13, 889 (1913).

CHAPTER XI

HYDROGENATIONS (Continued)

DIRECT HYDROGENATIONS OF LIQUIDS IN CONTACT WITH METAL CATALYSTS

541. We have explained the phenomena of direct hydrogenation as accomplished by various finely divided metals when the substance to be hydrogenated is brought in contact with the metal in the gaseous form, by assuming a sort of hydride of the metal, an unstable compound formed rapidly and decomposed rapidly in the act of hydrogenating the substance (165). This explanation does not necessarily require that the substance to be hydrogenated be in the gaseous form as we can see that the same reaction can be accomplished with a liquid material intimately mixed with a finely divided metal capable of taking up hydrogen. In order that the hydrogen may come into contact with the metal it is necessary that its solubility in the liquid be made sufficiently great by using low temperatures at the ordinary pressure, or a high pressure of hydrogen if it is necessary to heat.

An energetic and continuous agitation, constantly renewing the contact of the catalyst with the unchanged portions of the liquid will be most useful.

Furthermore, in order for the metal to be able to preserve its activity, it must not be oxidisable at the working temperature, or this temperature must be high enough to assure the reduction by the hydrogen in the system of any oxide formed.

542. From these conditions may be derived several methods which give results in general identical with those obtained by the method of Sabatier and Senderens of hydrogenating vapors over nickel, and which may offer great advantages in some cases.

The first attempt to hydrogenate substances directly in the liquid state had for its object the hydrogenation of liquid fats and was made in 1902–1903.¹ Then followed the method of Ipatief based on the use of nickel at 250 to 400° in the presence of hydrogen compressed to more than 100 atmospheres, and at almost the same time the method

¹ LEPRINCE and SIEVKE, German patent, 141,029 (1903). — NORMAN, English patent, 1515 of 1903. Chem. Cent., 1903 (1), 1199.

of Paal, relying on the use of colloidal metals (platinum or palladium) acting at near the ordinary temperature, and then in 1908, the method of Willstätter which depends on the use of platinum black.

We shall take up first the methods using the precious metals, then those employing the common metals whether at high pressures of hydrogen or at pressures near the atmospheric.

543. Except the process of Ipatief, which, on account of the high pressures used, demands an entirely special outfit, the various methods of hydrogenation in liquid medium employ apparatus of the same kind, though they may vary much in forms and dimensions. The main thing is a working vessel containing the liquid to be hydrogenated, either alone or dissolved in a suitable solvent and mixed with the solid catalyst. This recipient, which must be capable of being kept at known temperatures, is mounted on a mechanical shaker capable of assuring the best possible contact between liquid, catalyzer, and hydrogen. It is kept in communication with a cylinder of compressed hydrogen which can be introduced from time to time under known pressures, or if the hydrogenation is to be carried on at atmospheric pressure, the recipient communicates continuously with a hydrogen gasometer, the graduations of which enable us to follow the course of the reaction and to determine its end.

I. - METHOD OF PAAL

544. The methods of preparing colloidal platinum and palladium, such as are used in the method of Paal, have been given above (67 to 71). The amounts of these metals to be used are not over 16 to 50% of the weight of the substance to be hydrogenated, and can, according to Paal, be reduced to from 0.5 to 1% for colloidal palladium or to 1 to 2% for colloidal platinum.²

Use of Colloidal Palladium

545. Reductions with Simultaneous Fixation of Hydrogen. Nitro compounds are readily changed into amino compounds. Thus nitrobenzene is easily transformed into aniline, particularly at 65-85°.

Nitroacetophenone gives aminoacetophenone.4

The halogen of chlorine or bromine derivatives may be readily replaced by hydrogen when a current of hydrogen is passed through

- ² Paal, German patent, 298,193, 1913, Chem. Cent., 1917 (2), 145.
- ³ PAAL and AMBERGER, Berichte, 38, 1406 (1906).
- 4 SKITA and MEYER, Berichte, 45, 3579 (1912).

the compound containing colloidal palladium and boiling under reflux. Thus we obtain benzene from brombenzene. This reduction works well with o.chlor-benzoic acid, o.chlorcinnamic acid, chlorcrotonic acid, and chlorcaffeine, etc., without any other change in the molecule.

546. Fixation of Hydrogen by Addition. The ethylene double bond is readily hydrogenated.

Ethylene is easily transformed into ethane.

Sturene gives ethul-benzene.

Bromstyrene is simultaneously saturated and dehalogenated to ethyl-benzene.

1,10-Diphenyl-decadiene (1,9) furnishes 1,10-diphenyl-decane.3

Mesityl oxide, treated in alcohol solution with the metal prepared by means of gum arabic, changes into methyl-isobutyl-ketone.

 α -Methyl- β -ethyl-propenal, hydrogenated under the same conditions under 2 atmospheres pressure of hydrogen, gives chiefly the saturated aldehyde, α -methyl-valeric aldehyde, accompanied by a small amount of the unsaturated alcohol α -methyl-pentenyl alcohol.¹⁰

Crotonic, isocrotonic, and tetrolic acids are transformed into the corresponding saturated acids.¹¹

Fumaric acid in an hour and a half, and maleïc acid in seven hours are changed into succinic acid. Oleïc acid gives a 60% yield of stearic acid in 43 minutes.

Cinnamic acid is changed into phenylpropionic acid.12

Cinnamic aldehyde, dissolved in 20 parts of alcohol, is transformed into phenylpropionic aldehyde. 18

Isopropylidene-cyclopentanone adds H₂ to form isopropyl-cyclopentanone: ¹⁴

- ⁵ ROSENMUND and ZETSCHE, Berichte, 51, 579 (1918).
- ⁶ Paal and Hartmann, Berichte, 48, 984 (1915).
- 7 Borsche and Heimbürger, Berichte, 48, 452 (1915).
- BORSCHE and WOLLEMANN, Berichte, 44, 3185 (1911).
- WALLACH, Nach. Ges. der Wiss. Gottingen, 1910, 517. SKITA, Berichte, 48
 1486 (1915).
 - 10 SKITA, Berichte, 48, 1486 (1915).
- ¹¹ Börseken, van der Weide and Mom, Rec. Trav. Chim. Pays Bas., 35, 260 (1915).
 - ¹² Paal and Gerum, Berichte, 41, 2273 and 2277 (1908).
- ¹⁸ SKITA, Berichte, 48, 1691 (1915). BÖESEKEN, VAN DER WEIDE and MOM, Rec. Trav. Chim. Paye-Bas, 35, 260 (1916).
 - ¹⁴ WALLACH, Annalen, 394, 362 (1912).

547. In the case of diethylene compounds, if the double bonds are consecutive, both are hydrogenated simultaneously but if they are separated by more than one carbon atom, they are hydrogenated successively.

Thus phorone gives first dihydrophorone and then valerone.

Dibenzylidene-acetone, C_0H_5 . $CH:CH:CO:CH:CH:C_0H_5$, can give first benzyl-benzylidene-acetone, C_0H_5 . $CH:CH:CO:CH_2$.- CH_2 . C_0H_3 , and then dibenzyl-acetone. C_0H_3 .

548. The acetylene triple bond can be saturated in two steps. Acetylene itself gives ethylene chiefly, up to 80%.16

Phenyl-acetylene in acetic acid solution gives styrene and then ethyl-benzene.17

Tolane yields stilbene and then dibenzyl. Diphenyl-diacetylene passes into αγ-diphenyl-butadiene αγ, then into αγ-diphenyl-butane.

Phenylpropiolic acid, C₆H₅. C: C. COOH, gives a poor yield of cinnamic acid, C₆H₅. CH: CH. CO₂H, and does not go into phenylpropionic.¹⁸

2,5-Dimethyl-hexine (3)-diol (2,5) adds only H₂ to give the ethylene-diol, and the same is true of 1,4-diphenyl-butine (2)-diol (1,4) ¹⁹ and of dimethyl-diethyl-butine-diol,²⁰ while dimethyl-diphenyl-butine-diol gives, in succession, the ethylene glycol and the saturated glycol.²¹ On the contrary, 2-methyl-4-phenyl-butine (3)-ol(2), (CH₂)₂C(OH). C C.C₆H₅, adds 2H₂ immediately to give the saturated alcohol.²²

hydrogenated in alcohol solution, adds H₂ to form dimethyl-octenediol.²⁸

In the hydrogenations of these acetylene glycols, the speed of the reaction is usually proportional to the amount of catalyst present,

- ¹⁵ PAAL, Berichte, 45, 2221 (1912).
- ¹⁶ Paal and Hohenegger, Berichte, 48, 275 (1915). Paal and Schware, Berichte, 48, 1202 (1915).
 - 17 Kelber and Schwarz, Berichte, 45, 1951 (1912).
 - 18 PAAL and SCHWARZ, Berichte, 51, 640 (1918).
 - ¹⁹ Zalkind, J. Russian Phys. Chem. Soc., 45, 1875 (1914), C. A., 8, 1419.
- ²⁶ Zalkind and Miss Markaryan, J. Russian Phys. Chem. Soc., 48, 538 (1916), C. A., 11, 584.
- ²¹ Zalkind and Kvapishevskii, J. Russian Phys. Chem. Soc., 47, 688 (1915), C. A., 9, 2511.
 - ²² ZALKIND, J. Russian Phys. Chem. Soc., 47, 2045 (1915), C. A., 10, 1355.
- ZALKIND and MISS MARKARYAN, J. Russian Phys. Chem. Soc., 48, 538 (1916), C. A., 11, 584.

but sometimes it is independent of the amount of catalyst, contrary to all predictions.

549. The transformation of aldehydes and ketones into alcohols can be effected, but with difficulty.

Benzaldehyde is partially reduced to benzyl alcohol.24

Phenyl-acetaldehyde is regularly hydrogenated to the corresponding alcohol.

With hydrogen at one atmosphere pressure, phorone is hydrogenated to di-isobutyl-carbinol, but under half an atmosphere, the reduction stops at valerone.²⁴

In acetic acid solution, *mesityl oxide* is hydrogenated to methylisobutyl-carbinol, but in alcohol, as stated above, the reaction stops at the ketone.²⁸ The saturated alcohol is also obtained by working under 5 atmospheres pressure.²⁴

- 550. Hydroxy-methylene derivatives containing the group >C: CHOH, are changed into methyl derivatives >CH. CH₂. 26
 - 551. Benzoic acid furnishes hexahydrobenzoic.27
- 552. Carvone is transformed into tetrahydrocarvone. There is addition of hydrogen to the double bonds of pinene, which, under 2 atmospheres pressure, gives pinane, of camphene which passes to camphane, melting at 53° , of eucarvone, of α and β -terpineols, of thujone, of isothujone, of methylheptenone, of cyclohexenone, etc. Likewise pulegone is changed to menthone.
 - 553. Naphthalene is reduced to decahydronaphthalene. 30
- 554. Azobenzene, in alcohol solution under 2 atmospheres pressure of hydrogen, is reduced to hydrazobenzene in five minutes and then into aniline in 4.5 hours. Orange No. 3 is immediately decolorized under these conditions.⁸⁰

The α - and β -ionones are transformed into the odorless dihydroand then into the tetrahydroionones.³¹

555. Quinidine gives dihydroquinidine, melting at 165°. Cinchonidine adds H₂ to form the dihydro- melting at 229°. Cinchonine adds H₂ to form cinchotine. Cinchotine.

- № SKITA and RITTER, Berichte 43, 3393 (1910).
- ²⁵ SKITA, Berichte, 48, 1486 (1915).
- ²⁶ Kötz and Schaeffer, J. prakt. Chem. (2), 88, 604 (1913).
- 27 SKITA and MEYER, Berichte, 45, 3587 (1912).
- * SKITA and MEYER, Berichte, 45, 3579 (1912).
- 29 WALLACH, Annalen, 336, 37 (1904).
- 30 SKITA, Berichte, 45, 3312 (1912).
- MEYER and BERGEN, Berichte, 45, 3312 (1912).
- ²² SKITA and Nord, Berichte, 45, 3316 (1912).
- PAAL, German patent, 223,413.

Pyridine is changed to piperidine and quinoline to decahydro-quinoline.²⁷ Diacetyl-morphine furnishes the dihydro- and piperine, tetrahydropiperine.²⁸

Strychnine, dissolved in dilute nitric acid under 2 atmospheres pressure of hydrogen, gives the dihydro-, but under 3 atmospheres, tetrahydrostrychnine, while brucine always gives its dihydro-.**

Colchicine furnishes tetrahydrocolchicine. 25

Egg lecithine, dissolved in absolute alcohol, gives hydrolecithine. 86

Use of Colloidal Platinum

556. Colloidal platinum, prepared according to one of the methods given in Chapter II (67 to 71), can be substituted for colloidal palladium and gives results but little different.

According to Paal and Gerum its activity is less.⁸⁷ According to Fokin, on the contrary, the platinum is three times as active and much more apt to hydrogenate the aromatic nucleus.⁸⁸ The velocity of the hydrogenation increases rapidly with the amount of the metal employed.⁸⁹

557. The reduction of nitro-derivatives into amino- is readily carried out with nitrobenzens which gives aniline and with nitroacetophenone which yields aminoacetophenone.⁴⁰

558. The addition of hydrogen to double and triple bonds takes place easily even with many complex rings.

Ethylens is transformed to ethane but less rapidly than by colloidal palladium, the action being proportional to the amount of platinum used.⁴¹

Amylene is changed to pentane, oleic and linoleic acids into stearic and crotonic, maleic, aconitic, sorbic, citraconic, and itaconic acids are changed into the corresponding saturated acids, while allyl alcohol gives propyl alcohol.³⁸

Acetylens is reduced to a mixture of ethylene and ethane.

559. Heptaldehyde, hydrogenated by the aid of colloidal platinum prepared by the germ method, is changed to heptyl alcohol.48

- M SKITA and PAAL, German patent, 230,724, C., 1911 (1), 522.
- * HOFFMANN LA ROCHE & Co., German patent, 279,999, C., 1914 (2),1214.
- ⁸⁶ PAAL and OEHME, Berichte, 46, 1297 (1913).
- ²⁷ Paal and Gerum, Berichte, 40, 2209 (1907) and 41, 2273 (1908).
- * FOKIN, J. Russian Phys. Chem. Soc., 40, 276 (1908).
- 39 FOKIN, Z. Angew. Chem., 22, 1492 (1909).
- 46 SKITA and MEYER, Berichte, 45, 3579 (1912).
- 41 PAAL and SCHWARZ, Berichte, 48, 994 (1915).
- ⁴⁸ Paal and Schwarz, Berichte, 48, 1202 (1915).
- ⁴⁸ SEITA and MEYER, Berichte, 45, 3589 (1912).

a-Methyl-β-ethyl-propenal, treated in acetic acid solution, is changed completely into a-methyl-pentanol.

Mesityl oxide, in water solution, goes to methyl-isobutyl-ketone, but in acetic acid solution, into methyl-isobutyl-carbinol.44

560. The aromatic nucleus is hydrogenated more or less readily. With the metal prepared by the germ method, benzene is transformed into cyclohexane.

Toluene, in acetic acid solution under 2 atmospheres pressure, is changed to methyl-cyclohexane and benzoic acid into hexahydrobenzoic. 45

Cinnamic aldshyds is transformed into phenylpropionic aldshyds in the cold. In acetic acid solution phenylpropyl alcohol is obtained mixed with a little propyl-benzens, while with a larger amount of the catalyst and a pressure of 3 atmospheres, cyclohexyl-propyl alcohol is obtained. Under the same conditions, in acetic acid solution, benzyl-anilins furnishes hexahydrobenzyl-anilins accompanied by cyclohexyl-amins and methyl-cyclohexans.⁴⁶

Phenylacetaldehyde gives the corresponding alcohol with a little ethyl-benzene, cyclohexanol, cyclohexanone, and cyclohexane.

Benzaldshyds gives toluene and methyl-cyclohexane along with benzyl alcohol.

Benzophenone yields dicyclohexyl-methane at 60°.

a and β-Ionones, in acetic acid solution, furnish trimethyl-hydroxybutylcyclohexane.47

Caryophyllene, C15H24, adds H2 in methyl alcohol solution.48

561. With colloidal platinum, prepared with gum arabic, we can obtain piperidine from pyridine.45

The addition of 3H₂ takes place with various homologs of pyridine, hydrogenated in acetic acid solution under atmospheric pressure or under 2 or 3 atmospheres.⁴⁹

The pyridine-carbonic acids are transformed into piperidinic acids. 50

Quinoline gives, in turn, tetrahydro- and then decahydroquinoline. 45

Diacetyl-morphine adds H₂ and cinchonine yields hexahydrocinchonine. 51

- 44 SKITA, Berichte, 48, 1486 (1915).
- 45 SEITA and MEYER, Berichte, 45, 3589 (1912).
- 46 SKITA, Berichte, 48, 1685 (1915).
- 47 SEITA, Berichte, 48, 1486 (1915).
- ⁴⁸ DEUSSEN, Annalen, 388, 136 (1912).
- 49 SKITA and BRUNNER, Berichte, 49, 1597 (1916).
- 50 HESS and LIEBBRANDT, Berichie, 50, 385 (1917).
- SKITA and BRUNNER, Berichte, 49, 1597 (1916).

II. MRTHOD OF WILLSTÄTTER

562. The process consists in submitting to a current or to an unlimited amount of an atmosphere of hydrogen gas, the substance dissolved in a suitable vehicle and intimately mixed by means of constant agitation with the *platinum* or *palladium black*. It was employed first by Fokin, who transformed in this way *oleic acid* dissolved in ether into stearic acid by a current of hydrogen in the cold with palladium or platinum black as catalyst. ¹²

But Willstätter is the one who has generalized this method by applying it to various uses.

Platinum black prepared according to the method of Loew (62) serves best.⁵² Palladium black can also be used: it is prepared by reducing palladous chlorids with formaldehyde in the presence of caustic soda.⁵⁴ But it is not so desirable as platinum black.

The substance dissolved in ether or in any other inert solvent is treated with the *platinum black* and is put into a flask which is continually agitated by a mechanical shaking machine and which communicates with a gasometer filled with hydrogen. According to circumstances, quite different amounts of platinum black are used, varying from 3 to 33% of the weight of the substance.

Dilution of the material is not indispensable to the success of the method.

Use of Platinum Black

563. Willstätter has called attention to this quite unexpected fact, that in certain cases hydrogenation by means of platinum black is not possible unless it has been previously charged with a certain proportion of oxygen.

In most cases, platinum black containing oxygen or free from oxygen may be used indifferently, as in the hydrogenation of benzene to cyclohexane; on the contrary, the hydrogenation of pyrrol requires platinum black absolutely free from oxygen. On the other hand, the decomposition of hydrazine demands that the platinum black that is to be used be previously aerated.⁵⁵

The aeration of the platinum black is indispensable for the hydrogenation in acetic acid solution of phthalic and naphthalic anhydrides and the reaction does not continue unless the apparatus is opened

- 22 FOKIN, J. Russian Phys. Chem. Soc., 39, 607 (1907).
- * Somewhat improved method WILLSTÄTTER and WALDSCHMIDT-LEITZ, Berichte. 54, 121 (1921). E. E. R.
 - BRETEAU, Div. meth. d'hydr. app. au Phénant, Paris, 1911, p. 25.
 - * PURGOTTI and ZANICHELLI, Gas. Chim. Ital., 34 (1), 57 (1904).

from time to time for the aeration of the black. Oxygen appears to play an active part in the hydrogenation which is indicated by the products obtained. For phthalic anhydride the products are, hexahy-

acids, and for naphthalic acid, tetrahydronaphthalid, hexahydronaphthalid, decahydroacenaphthene, C₁₂H₂₀, and tetrahydro-methyl(1) naphthalene-carbonic acid(8).

The influence of these anhydrides on the conditions of hydrogenation can effect even the hydrogenation of the dibasic acids themselves; the presence of the anhydrides prevents this from taking place unless the platinum be aerated. *Isophthalic* acid, which usually contains traces of the anhydride, can not be hydrogenated except with aerated platinum.⁵⁶ ⁵⁷

564. Nitro Compounds. The reduction of nitro or nitroso compounds to amino is readily effected by 1 cg. of platinum black to 1 g. of the material dissolved in ether or acetone. A few minutes are sufficient for complete reduction; thus p.nitrotoluens is changed to p.toluidins, 1-nitrosonaphthol(2), into aminonaphthol. But the nitrosoterpenes are changed quantitatively into the corresponding hydrox-vlamines. 58

565. Ethylene Double Bonds. These are readily saturated. Amylene is changed to pentane.

 ω -Nitrostyrens, dissolved in absolute alcohol or in glacial acetic acid, adds a single atom of hydrogen, two molecules combining ⁵⁰:

$$\begin{array}{cccc} C_6H_5.CH:CH.NO_2 & \longrightarrow & C_6H_5.CH.CH_2.NO_2 \\ C_6H_5.CH:CH.NO_2 & \longrightarrow & C_6H_5.CH.CH_2.NO_2 \end{array}$$

Olete alcohol is readily transformed into octadecyl alcohol, ethyl oleate quantitatively into ethyl stearate, and erucic alcohol into docosyl alcohol.

MILLSTÄTTER and JACQUET, Berichte, 51, 767 (1918).

^{54, 113 (1920)]} show that the presence of oxygen in the platinum black is necessary in all cases. This oxygen is gradually used up by the hydrogen during the process of hydrogenation. With ethylene compounds the addition of the hydrogen is so extremely rapid that the desired hydrogenation may be accomplished before the catalyst becomes inactive by loss of its oxygen but if the hydrogenation is slow, the catalyst may require revivification by aeration at intervals during the process. In this respect palladium black and even nickel act similarly to platinum black. — E. E. R.

⁵⁸ Cusmano, Lincei, 26 (2), 87 (1917).

⁵⁰ Sonn and Schnellemberg, Berichte, 50, 1913 (1917).

Phytens, C₂₀H₄₀, gives phytans, C₂₀H₄₂; phytol, C₂₀H₃₀OH, dihydrophytol, C₂₀H₄₁OH, slowly but with a good yield. Geraniol (416) is hydrogenated only slowly and gives the corresponding saturated alcohol at the end of several days.⁶⁰

Linalool furnishes 2, 6-dimethyl-octanol (6).61

Safrol and isosafrol are hydrogenated in two hours to dihydrosafrol. Likewise eugenol and isosugenol pass into isopropul-quaiacol. **

Piperonal-acetone and dipiperonal-acetone are transformed into the saturated ketones.

Cholesterine, in ether solution with one third its weight of platinum black, is changed into dihydrocholesterine in two days.⁶⁴

Oletc acid gives stearic and ethyl oleate yields ethyl stearate. 65

566. Acetylene Triple Bonds. The acetylene glycols of the formula, RR': C(OH).C:C.C(OH): RR', give the corresponding saturated glycols and also certain amounts of the alcohols, RR': C(OH)CH₂.-CH₂.CH₂: RR'.⁶⁴ Thus 2, 5-dimethyl-hexine(3)diol(2, 4) furnishes the saturated glycol.⁶⁶

Dimethyl-diethyl-butine-diol, which adds only H₂ with colloidal palladium (548), takes up 2H₂ with platinum black.⁶⁷

Dimethyl-diphenyl-butine-diol can add H₂ and then 2H₂ by steps. ⁶⁸
Octadi-ine-diol(1, 8), HOH₂C.C. C.CH₂.CH₂.C. C.CH₂OH, hydrogenated at 70° in alcohol solution, gives a mixture of octane-diol (1, 8) and n.octyl alcohol. ⁶⁹

Octadi-ine-dioic acid, HO₂C.C : C.CH₂.CH₂.C : C.CO₂H, dissolved in a mixture of alcohol and ether, furnishes substic acid in four days.⁷⁰

- 567. Aldehydes and Ketones. Aldehyde and ketone groups can be regularly transformed into the corresponding alcohol groups. Crotonic aldehyde, in anhydrous ether, is changed in eleven hours into a mixture of 70% butyric aldehyde and 30% butyl alcohol.⁷¹
 - WILLSTÄTTER and MAYER, Berichte, 41, 1475 (1908).
 - EL BARBIER and Locquin, Compt. rend., 158, 1555 (1914).
 - FOURNIER, Bull. Soc. Chim. (4), 7, 23 (1910).
 - [∞] VAVON and FAILLEBIN, Compt. rend., 169, 65 (1919).
 - 4 WILLSTÄTTER and MAYER, Berichte, 41, 2199 (1908).
 - DUPONT, Compt. rend., 156, 1623 (1913).
 - ⁶⁴ ZALKIND, J. Russian Phys. Chem. Soc., 45, 1875 (1914), C. A., 8, 1419.
- ⁶⁷ Zalkind and Miss Markaryan, J. Russian Phys. Chem. Soc., 48, 538 (1916), C. A., 11, 584.
- ⁶⁸ ZALKIND and KVAPISHEVSKII, J. Russian Phys. Chem. Soc., 47, 688 (1915), C. A., 9, 2511.
 - 69 LESPIEAU, Compt. rend., 158, 1187 (1914).
 - 76 LESPIEAU and VAVON, Compt. rend., 148, 1335 (1909).
 - ¹¹ FOURNIER, Bull. Soc. Chim. (4), 7, 23 (1910).

Actors is changed to isopropyl alcohol in water solution and methylsthyl-ketons is changed into methyl-ethyl-carbinol in 12 hours. Disthyl and dipropyl-ketonss are similarly reduced.

The transformation into the alcohol is more readily effected with cyclopentanons dissolved in 5 volumes of ether, with cyclohexanons and with the methyl-cyclohexanones.

Mesityl oxide gives first methyl-isobutyl-ketone and then methyl-isobutyl-carbinol.

In ether solution, phorone yields dissobutyl-ketons, while in acetic acid it gives dissobutyl-carbinol.

Citral in ether solution gives a mixture of 2, 6-dimethyl-octane and 2, 6-dimethyl-octanol (8).72

Menthone yields menthol and pulegone gives pulegomenthol.⁷² Carrons with 20% of platinum black takes up in succession, H₂, 2H₂, 3H₂ to form carvotanacetone, tetrahydrocarvone and finally carvomenthol slowly.⁷⁴

568. Aromatic aldehydes are transformed almost quantitatively into alcohols, which is a valuable reaction since other methods give hydrocarbons (388). With 10 g. of black a gram molecule can be hydrogenated in a few hours. This can be done with benzaldshyde, methylsalicylic, benzoyl-salicylic, and anisic aldshydes, vanilline and its methyl, ethyl, acetyl, and benzoyl derivatives, piperonal, which gives the alcohol melting at 54°, and cinnamic aldshyde, which yields phenyl-propyl alcohol.⁷⁵

At 70° anisaldshyds gives anisalcohol but at 97° it is polymerized. On the contrary, acetophenous takes up 10 atoms of hydrogen at once to form sthyl-cyclohexans. 76

569. Aromatic Nucleus. Aromatic compounds are completely hydrogenated to cyclohexane derivatives on the condition that they are perfectly pure. Traces of impurities, particularly sulphur compounds, hinder the reaction.⁷⁷

Toluens and the xylenes are hydrogenated more readily than benzene, and higher homologs still more readily. Butyl-benzene, amylbenzene, hexyl-benzene, octyl-benzene, etc., up to pentadecyl-benzene are readily changed in acetic acid solution into the corresponding cyclohexane derivatives.⁷⁸

Durene furnishes hexahydrodurens.

```
<sup>72</sup> VAVON, Ann. Chim. (9), 1, 144 (1914).
```

⁷⁸ VAVON, Compi. rend., 155, 287 (1912).

⁷⁴ VAVON, Compt. rend., 153, 68 (1911).

⁷⁵ Vavon, Compt. rend., 154, 359 (1912).

⁷⁶ VAVON, Compt. rend., 155, 287 (1912).

⁷⁷ WILLSTÄTTER and HATT, Berichte, 45, 1471 (1912).

⁷⁸ HALSE, J. prakt. Chem. (2), 92, 40 (1915).

Styrene gives ethyl-cyclohexane; and phenol, cyclohexanol. Lugenol adds 4H, to form propul-methoxu-cyclohexanol.

Aniline produces chiefly dicyclohexyl-amine with only 10% of cyclohexyl-amine. Chlortoluene is transformed into methyl-chlorcyclohexane.

In ether solution, benzoic acid is slowly changed to hexahydrobenzoic acid without intermediate products.⁸¹

In acetic acid solution, p.aminobenzoic acid is quantitatively reduced to p.aminocyclohexane-carbonic acid and hydroxybenzoic acid is similarly hydrogenated.³²

We have seen above (563) that phthalic anhydride can be hydrogenated by means of platinum black aerated from time to time. The ordinary method serves well for *phthalimids* which gives as the sole product, *hexahydrophthalimids*: **

570. Terpenes. Limonens, in ether solution with 25% of its weight of platinum, adds H₂ in 30 minutes in the cold to form carromenthens, boiling at 175°, and then an additional H₂ in 65 minutes to form menthans.⁸⁴

Pinene, 500 g. with 15 g. platinum, absorbs hydrogen rapidly, 60 l. per hour at the start, and at the end of 24 hours is entirely transformed into dihydropinene, boiling at 166° (477). Camphene gives a solid dihydrocamphene melting at 87°.85

 α -Thujens, C₁₆H₁₆, which by the method of Sabatier and Senderens yields menthane (478), is totally transformed by platinum black and hydrogen under 25 to 50 atmospheres in two days into thujans, C₁₀H₁₈, boiling at 157°, the inner ring remaining intact. Similar transformations take place with β -thujens and with sabinens.⁸⁶

Isoamyl-carvol adds 2H₂ to give the corresponding saturated alcohol.⁸⁷

The sesquiterpenes, C₁₅H₂₄, as well as their ketone and alcohol derivatives, add 4 or 6 atoms of hydrogen.

- 79 WILLSTÄTTER and KING, Berichte, 46, 527 (1913).
- ³⁰ Madinaveitia and Blanes, Soc. Espan. Fis. Quim., 10, 381 (1913), C. A., 7, 3500.
 - WILLSTÄTTER and MAYER, Berichte, 41, 1475 (1908).
 - * HOUBEN and PEAU, Berichte, 49, 2294 (1916).
 - * WILLSTÄTTER AND JACQUET, Berichte, 51, 767 (1918).
 - M VAVON, Bull. Soc. Chim. (4), 15, 282 (1914).
 - * VAVON, Compt. rend., 149, 997 (1909) and 152, 1675 (1911).
 - TCHOUGAEFF and FOMIN, Compt. rend., 151, 1058 (1910).
 - 87 SEMMLER, JONAS and OELSNER, Berichte, 50, 1838 (1917).

Thus isozingiberens, 88 sudesmens, 80 and ferulens, 90 take up 2H₂. The same is true of doremons, C₁₅H₂₆O, which gives tetrahydrodoremons without alteration of the ketone group and of doremol which forms the saturated alcohol. Farnesol, C₁₅H₂₆O adds 3H₂. 90

Betulol, C₁₅H₂₄O, adds 2H₂ to form the alcohol, C₁₅H₂₈O, when it is hydrogenated in anhydrous ether solution.⁹¹

571. Complex Rings. Cyclo-octenone is changed to cyclo-octanone by 10% of its weight of black. Cyclo-octatriens and cyclo-octatetrens, CH: CH.CH: CH

CH . CH CH . CH

In the hydrogenation of the latter, the first three H₂ are fixed with about the same velocity, while the last H₂ is added only about half so fast. ²⁸

Naphthalens adds hydrogen rapidly to form the dihydro- and then the tetrahydro- and finally, more slowly, decahydro-naphthalens.⁹⁴

Phenanthrens, dissolved in ether, gives dihydro-phenanthrens (melting at 94°), in two days in the cold, or in 8 hours at the boiling point of the ether. However, Breteau failed to obtain any hydrogenation in cyclohexane solution. 66

Santonine yields tetrahydro-santonine when hydrogenated in glacial acetic acid.⁹⁷ Sodium santonate takes up the same amount of hydrogen to form sodium tetrahydrosantonate.⁹⁸

Pyrrol adds 2H₂ to form pyrrolidins.99

Indol, in glacial acetic acid, yields octahydro-indol, an alkaline liquid with a disagreeable odor boiling at 182°, accompanied by a little dihydro-indol. 100

- 572. Quinine sulphate is completely hydrogenated in dilute sulphuric acid solution by hydrogen under a pressure of more than an atmosphere to dihydroquinine sulphate, the hydrogenation being con-
 - ³² SEMMLER and BECKER, Berichte, 46, 1814 (1913).
 - ** SEMMLER and RISSE, Berichte, 46, 2303 (1913).
 - ⁸⁰ SEMMLER, JONAS and ROENISCH, Berichte, 50, 1823 (1917).
 - ⁸¹ SEMMLER, JONAS and RICHTER, Berichte, 51, 417 (1918).
 - WILLSTÄTTER and WASER, Berichte, 44, 3434 (1911).
 - WILLSTÄTTER and HEIDELBERGER, Berichte, 46, 517 (1913).
- ** WILLSTÄTTER and HATT, Berichte, 45, 1471 (1912). WILLSTÄTTER and King, Ibid., 46, 527 (1913).
 - SCHMIDT and FISCHER, Berichte, 41, 4225 (1908).
 - BRETEAU, Méth. d'hydrog. app. au Phénant, Paris, 1911, p. 20.
 - 97 ASAHINA, Berichte, 46, 1775 (1913).
 - 98 CUSMANO, Lincei, 22, 507 (1913).
 - •• WILLSTÄTTER and HATT, Berichte, 45, 1371 (1912).
 - 100 WILLSTÄTTER and JACQUET, Berichte, 51, 767 (1918).

tinued till the solution does not decolorize potassium permanganate. Dihydromorphine and dihydrocodeine can be obtained in the same way. 102

Use of Palladium Black

- 573. The use of palladium black ¹⁰⁸ immersed in the liquid appears to be usually less advantageous than the use of platinum black. However, it has led to some remarkable results, such as the reduction of carbonates to formates.
- 574. Reduction without Addition of Hydrogen. The most important reaction is the synthesis of formates by the reduction of bicarbonates:

$$KHCO_3 + H_2 = HCO_2K + H_2O.$$

This requires a high pressure and a temperature around 70°.

In a silver plated bomb, 10 g. potassium bicarbonate, 200 cc. water, and 1.5 g. palladium black are placed with hydrogen at 60 atmospheres. After heating for 24 hours to 70°, 74.7% of the salt is changed to formate.

The reaction takes place without catalyst, but extremely slowly, only 0.6% of formate being produced in 24 hours.

The potassium bicarbonate can be replaced by sodium borate, the bomb then being filled with equal volumes of carbon dioxide and hydrogen under 60 atmospheres.¹⁰⁴

The reaction can be carried out without the presence of the alkali salt, by maintaining a mixture of carbon dioxide and hydrogen under high pressure in the presence of water and palladium black. By working at 20° and under a pressure of 110 atmospheres a 1% solution of formic acid is obtained.¹⁰⁵

575. Reduction of Acid Chlorides. Another reaction which is peculiar to palladium black is the reduction of acid chlorides to aldehydes:

$$R.COCl + H_2 = R.CHO + HCl.$$

The acid chloride, dissolved in a hydrocarbon, is submitted to hydrogenation in the presence of palladium black precipitated on barium sulphate.

Benzoyl chloride gives benzaldehyde with a yield of 97%; butyryl

- 101 VEREIN, CHININFABR. ZIMMER & Co., English patent 3,948 of 1912.
- 102 German patent 260,233.
- ¹⁰⁶ Preparation WILLSTÄTTER and WALDSCHMIDT-LEITZ, Berichte, 54, 123 (1921). E. E. R.
 - 104 Bredig and Carter, Berichte, 47, 541 (1914).
- ¹⁰⁶ Bredig and Carter, English patent, 9,762 of 1915; J. S. C. I., 34, 1207 (1915).

chlorids furnishes 50% of the aldehyde and stearyl chlorids is reduced to its aldehyde. 106

576. Nitro Compounds. The reduction of nitro to amino compounds is difficult to carry out with palladium, but nitrobenzens does give aniline on prolonged contact with an excess of hydrogen and palladium black in alcohol solution.¹⁰⁷

577. Ethylene and Acetylene Bonds. Olsic acid, in ether solution, is slowly transformed to stearic acid, the reduction being rapid when it is carried on at a higher temperature and with hydrogen under pressure. The same is true for the ssters of olsic acid and this is the basis for the industrial use of palladium black in the hardening of liquid fats (946).

hydrogen under 35 atmospheres in the presence of palladium chloride, which is reduced, yields ethyl-trimethylene. 108

The acetylene glycols of the type, RR': C(OH).C:C.C(OH): RR', yield mainly the saturated hydrocarbons, RR': CH.CH₂.CH₂-CH: RR'. 109

Eugenol stops with the formation of dihydroeugenol, 110 the ring not being hydrogenated as with platinum black (569).

- 578. Aromatic Nucleus. The hydrogenation of the aromatic nucleus is not usually effected by palladium black, but the hydrogenation of hexahydroxybenzens to inosite at 50-55° may be mentioned. The inosite formed melts at 218° as does natural inosite. 111
- 579. Phenanthrens is hydrogenated, in cyclohexane solution, by half its weight of the black to tetrahydrophenanthrens.¹¹²

Use of other Metals of Platinum Group

580. Ruthenium Black. The black prepared by formaldehyde and ruthenium chloride solution has a catalytic activity inferior to that of platinum.

If 0.05 g. of this black is added to 0.5 g. cinnamic acid in 2 cc. glacial acetic acid, phenyl-propionic acid is formed in 8 hours without

- 166 ROSENMUND, Berichte, 51, 585 (1918).
- 107 GERUM, Inaug. Dissertation, Erlangen, 1908.
- 100 FILIPPOV, J. Russian Phys. Chem. Soc., 44, 469 (1912).
- 100 DUPONT, Compt. rend., 156, 1623 (1913).
- 110 MADINAVEITIA and BLANES, Soc. Espan. Fis. Quim., 10, 381 (1913), C. A., 7, 3500.
 - 111 WIELAND and WISHORT, Berichte, 47, 2082 (1914).
 - 112 Breteau, Div. meth. hydrog., Paris, 1911, p. 26.

the ring being attacked. *Toluene*, dissolved in acetic acid and subjected to hydrogenation for 8 hours, is not affected. 113

- 581. Rhodium Black. Rhodium black is more active than ruthenium. Under the conditions given above, cinnamic acid is transformed into phenyl-propionic in 3 hours and into cyclohexyl-propionic in 15 hours. Toluens can be hydrogenated to hexahydrotoluens in 12 hours by 10% of its weight of the black.¹¹³
- 582. Iridium Black. This black prepared by reducing the *chlorids* by sodium formate, has an activity entirely analogous to that of ruthenium black (580).¹¹³
- 583. Osmium Black. This black, prepared by reducing osmic anhydride by formic acid, does not effect any hydrogenation of cinnamic acid in 5 hours.¹¹⁸

Osmium dioxids has been mentioned as able to hydrogenate oils when used to the amount of 0.5%, 114 but it is certain that it acts after it is reduced to the metal which is the true catalyst. 115

- 113 Madinaveitia, Soc. Espan. Fis. Quim., 11, 328 (1913).
- 114 LEHMANN, Arch. Pharm., 251, 152 (1913), C. A., 8, 586 (1914).
- 115 NORMAN and Schick, Arch. Pharm., 252, 208 (1914).

CHAPTER XII

HYDROGENATIONS (Continued)

DIRECT HYDROGENATION OF LIQUIDS IN CONTACT WITH METAL CATALYSTS (Continued)

III. METHOD OF IPATIEF

584. This method consists in warming the substance to be hydrogenated in contact with nickel or nickel oxide and hydrogen compressed to at least 100 atmospheres in a very strong container. The hydrogenation velocity is greater when the oxide is used, which Ipatief attributes to the real catalytic power of the oxide. As we have seen above (80), catalytic power appears to belong exclusively to the metal; since the temperature is always above 250°, the nickel oxide must at least in part be reduced to the metal which is more active than the metal prepared in advance and which has been subjected to incandescence more or less intense while being introduced into the container, being thereby agglomerated and reduced in catalytic power.

The nickel is frequently replaced by copper, copper oxide, iron or palladium, or even by zinc powder.

585. Apparatus. The apparatus used for all of this work consists of a soft steel tube lined with copper, holding 250 to 275 cc. and capable of sustaining 600 atmospheres at 600°.¹ It is heated electrically by a nickel resistance wire. Changes of pressure are shown by a manometer. If the apparatus has been filled with hydrogen at a certain pressure, the pressure increases according to the rise of temperature, if there is no absorption of hydrogen or evolution of gas, but less rapidly if there is absorption of hydrogen, while if there is decomposition with evolution of gas, the pressure increases more rapidly and this increase measures the rate of decomposition.

The material of which the apparatus is constructed appears to influence the results in some way. Thus in a bronze tube the use of reduced copper as catalyst did not effect the complete hydrogenation of the aromatic nucleus, while this was realized in an iron apparatus.²

¹ IPATIEF, Berichte, 37, 2961 (1904). — J. Russian Phys. Chem. Soc., 36, 786 (1904), C., 1904 (2), 1020.

³ IPATIEF, J. Russian Phys. Chem. Soc., 42, 1557 (1910).

Use of Nickel

In order to carry out a hydrogenation, about 25 g. of the material to be hydrogenated is placed in the apparatus with 2 to 3 g. nickel oxide (NiO or Ni₂O₃) and hydrogen is admitted to 100 atmospheres at which pressure it holds about one gram molecule of hydrogen. The temperature of heating may reach 400° or even 600° and the resulting pressure may be 2.5 or 3 times the original, i.e. 250 to 300 atmospheres. The necessity of having an expensive apparatus and the real dangers of its use are against the general employment of the method of Ipatief, which is not superior to the method of Sabatier and Senderens except in special cases where the slowness of hydrogenation or the need of high pressures requires its use. Most organic substances give the same products by both methods.

586. Formation of Methane. The direct hydrogenation of carbon in the presence of nickel, oxide of nickel, or nickel oxide and alumina, does not take place below 600° under moderate pressures of hydrogen, but under very high pressures, *methane* is produced above 500°, the amount increasing with rise of temperature.

The reduction of carbon dioxide to methans which takes place incompletely at 450° under ordinary pressure, is not more complete at high pressures even with an excess of hydrogen.³

587. Ethylene Double Bonds. The hydrogenation of substances containing ethylene bonds is readily effected.

Olsic acid heated a long time at 100° with finely divided nickel and hydrogen at 25 atmospheres is not affected, but under 60 atmospheres pressure it is changed to stearic acid in 12 hours. Liquid fats are transformed to solid.⁴

50 g. cottonseed oil with 3 g. nickel oxide at 220–230° with hydrogen at 60 atmospheres gave in 4 hours a fat with iodine number (938) of only 11, while at ordinary pressures this result was obtained only at 255.° ⁶

Dimethyl-allyl-carbinol is changed to dimethyl-propyl-carbinol under the same conditions.

At 140 to 150°, mesityl oxide gives methyl-isobutyl-ketone mixed with a little of the corresponding alcohol.

Cuclohexene is reduced to cuclohexane.

- ⁸ IPATIEF, J. prakt. Chem. (2), 87, 479 (1913).
- ⁴ FOKIN, J. Russian Phys. Chem. Soc., 38, 419 and 855 (1906).
- ⁵ IPATIEF, J. Russian Phys. Chem. Soc., 46, 302 (1914).
- ⁶ With high speed stirring this reduction can be accomplished in about the same time with 0.1 g. nickel on infusorial earth with hydrogen at atmospheric pressure at 180°. E. E. R.

588. Aldehydes and Ketones. The transformation of aliphatic aldehydes and ketones into the alcohols can be accomplished, but it is limited by the inverse reaction of dehydrogenation especially when the temperature exceeds 200 to 250°.

Isobutyric and isovaleric aldshydes are partly reduced to the corresponding alcohols at 250° and 100 atmospheres.

At 250° acstons is completely changed to isopropyl alcohol and the same is true of various aliphatic ketones at around 200°. At about 280° the hydrogenation is limited by the inverse reaction which increases with elevation of temperature. From 300 to 325° acetone no longer gives any alcohol since isopropyl alcohol is decomposed into water, propane and lower saturated hydrocarbons, especially methane.

Lastruloss in solution is transformed, at 130° under 100 atmospheres, to α -mannits, glucoss into sorbits, and galactoss into dulcits.

589. Aromatic Nucleus. The hydrogenation of the aromatic nucleus is realized in all cases.

Benzens is totally changed to cyclohexans in 1.5 hours at 250° with 8% nickel oxide. Nickel sesquioxide gives better results than the monoxide. At 300° the cyclohexane produced does not remain but is decomposed into benzene, methane and carbon.⁸

At 250° diphenyl is reduced to dicyclohexyl and dibenzyl to dicyclohexylethane.

At 245° phenol is transformed in 14 hours to cyclohexanol accompanied by some cyclohexans. At 200° hydroquinons gives quinits. The product is a mixture of the cis and trans forms, but the yield is poor, as most of the diphenol goes into resinous products. 10

At 230° under 100 atmospheres, phenyl oxide gives in 12 hours a mixture of cyclohexyl oxide, cyclohexanol and cyclohexans.¹¹

Anisol, C₆H₅OCH₂, in 24 hours at 240° under 100 atmospheres, gives 40% hexahydroanisol accompanied by cyclohexanol and cyclohexane.

Guaiacol, o.HO.C₆H₄.OCH₃, in 12 to 15 hours at 220 to 240° and 100 atmospheres, yields hexahydroguaiacol with cyclohexanol and a little cyclohexane.¹²

- ⁷ IPATIEF, J. Russian Phys. Chem. Soc., 38, 75 (1906) and 39, 681 (1907), C. A., 1, 2877. Berichte, 40, 1270 (1907).
- ² IPATIEF, J. Russian Phys. Chem. Soc., 39, 681-693 (1907), C. A., 1, 2877 and 2878
- IPATIEF, J. Russian Phys. Chem. Soc., 38, 75 (1906) C., 1906, (2), 86.—Berichte, 40, 1281 (1907).
 - 10 IPATIEF and LOUVOGOI, J. Russian Phys. Chem. Soc., 46, 470 (1914).
- ¹¹ IPATIEF and PHILIPOW, J. Russian Phys. Chem. Soc., 40, 501 (1908), C., 1908 (2), 1098. IPATIEF, Berichte, 41, 993 (1908).
 - 12 IPATTEF and Louvogoi, J. Russian Phys. Chem. Soc., 46, 470 (1914).

590. The hydrogenation of phenols having unsaturated side chains is accomplished in two steps. At 95° and 30 to 50 atmospheres, only the side chain is attacked but by raising the temperature to 185 to 200°, the nucleus is also hydrogenated.

Thus, anethol. p.CH₂O.C₆H₄.CH: CH.CH₂, with 10% nickel at 95° and 50 atmospheres is transformed completely in 4 hours to methoxy-propyl-benzens, but 20 hours at 200° produce propyl-cyclohexans, the methoxy group being reduced to water and methane.

Likewise eugenol, H₃C(HO)C₆H₃.CH₂CH: CH₂, and isoeugenol, H₃C(HO)C₆H₃.CH: CH.CH₃, furnish methoxy-propyl-phenol in 2 or 3 hours at 29°, while at 195° in 7 hours, the chief product is methoxy-propyl-cyclohexane, H₃CO·C₆H₁₀·C₃H₇, the phenol group being eliminated.

The methyl ether of eugenol adds only H₂ at 95°, but in 10 hours at 210°, the same product is obtained as from eugenol.

Safrol and isosafrol are not hydrogenated at ordinary pressure at 140 to 160° with constant agitation for 5 hours, but under 50 atmospheres at 93°, dihydrosafrol, boiling at 228°, is obtained in 2 hours. In 10 to 12 hours at 180°, a product is obtained boiling at 207° which appears to be methoxy-propyl-cyclohexans.¹³

By 50 hours heating at 220° under 115 atmospheres, aniline gives 40 to 50% of cyclohexyl-amine, about 10% dicyclohexyl-amine and some cyclohexyl-aniline. 14

Diphenyl-amine vields dicyclohexyl-amine. 15

Benzaldehyde, at 200°, gives toluene and methyl-cyclohexans, 16 while at 280° in 12 hours, toluene, dibenzyl and resinous products are obtained. 14

Aromatic ketones act as they do in Sabatier's process (389) and yield hydrocarbons, benzophenone going into diphenyl-methane and benzoine into dibenzyl.¹⁵

Ipatief's process is useful for the hydrogenation of aromatic acids, but it is not well to use the free acids which attack the nickel nor the esters which give poor results (sthyl terephthalats is decomposed into ethyl p.toluats, methane and carbon dioxide), but the alkaline salts. Thus potassium benzoate gives 40% of the hexahydrobenzoats at 280° in 9 hours and sodium benzoats is even more readily hydrogenated.

- ¹⁸ IPATIEF, Berichte, 46, 3589 (1913).
- 14 IPATIEF, Berichte, 41, 993-1001 (1908).
- 15 IPATTEF, J. Russian Phys. Chem. Soc. 40, 491 (1908), C., 1908 (2), 1098.
- ¹⁶ IPATIEF, J. Russian Phys. Chem. Soc., 38, 75 (1906), C., 1906 (2), 86.
- ¹⁷ IPATIEF, J. Russian Phys. Chem. Soc., 38, 75 (1906) and 39, 693 (1907), C. A., 1, 2877.

Potassium phthalats gives the hexahydrophthalats at 300° in good vield. 18

Sodium cinnamate gives the cyclohexyl-propionate at 300° under 100 atmospheres. 19

591. Terpenes. Terpene compounds undergo the regular transformations.²⁰

Limonens is transformed into dihydrolimonene and then into menthans at 300-320° under 120 atmospheres.

At 265° pinene gives pinane and menthane at 300°.

At 240°, in 10 to 15 hours, camphene furnishes an isocamphane melting at 57° and boiling at 162.5°.

At 280°, under 120-130 atmospheres, carrons passes into carromenthons. At 220°, pulsyons gives menthons, which at 280° is mixed with menthans.

Camphor is completely changed into borneol at 350°.

592. Various Rings. At 250° under 120 atmospheres, naphthalene gives, in turn, tstrahydro- and decahydro-naphthalene.

The α - and β -naphthols are changed to α - and β -decahydronaphthols, melting at 57° and 99° respectively.²¹

Anthracens, submitted to repeated hydrogenations at 260-270° under 100 to 125 atmospheres for 10 to 16 hours, gives in succession, tetrahydro-, decahydro- (m.73°) and finally perhydroanthracene (m.88°) and at the same time is partially destroyed.

At 400°, phenanthrens gives better results, the dihydro- and then the tetrahydro- being obtained and, by a second operation, the octahydro- and perhydrophenanthrens with the odor of caoutchouc.²²

Quinoline first yields tetrahydroquinoline and then, almost quantitatively. decahydroquinoline.22

Use of Iron

- 593. At 350-400°, iron transforms aliphatic aldehydes and ketones into the alcohols. Acetons, at 400° and 103 atmospheres in 20 hours yields 25% of isopropyl alcohol. Isobutyric aldehyde gives 75% of the corresponding alcohol at 350°, but acetaldehyde is partly resinified and partly decomposed into carbon monoxide and methane.
- ¹⁸ IPATIEF and PHILIPOW, J. Russian Phys. Chem. Soc., 40, 501 (1908), C., 1908
 (2), 1098. IPATIEF, Berichte, 41, 993 (1908).
 - ¹⁹ IPATIEF, J. Russian Phys. Chem. Soc., 41, 1414 (1909).
- ²⁰ IPATIEF, Berichte, 43, 3546 (1910). IPATIEF and MATOW, Berichte, 45, 3205 (1912).
- ²¹ IPATIEF, J. Russian Phys. Chem. Soc., 39, 693 (1907), C. A., 1, 2877.— Berichts, 40, 1281 (1907).
 - 21 IPATIEF, JAKOWLEW and RAKITIN, Berichte, 41, 996 (1908).
 - * IPATIEF, J. Russian Phys. Chem. Soc., 40, 491 (1908), C., 1908 (2), 1098.

The hydrogenation of the aromatic nucleus does not take place, even at 420°, but cyclohexane is brought back to benzene.²⁴ At 280° benzaldehyde gives a mixture of toluene and dibenzyl. The same result is obtained when benzyl alcohol is hydrogenated at 350° and 96 atmospheres.²⁵

Use of Copper

594. Copper, or copper oxide (certainly reduced to the metal), readily permits the hydrogenation of ethylene bonds at 300 to 350° under 100 to 200 atmospheres, but when used alone does not effect the hydrogenation of the benzene ring.**

Sodium cinnamate is changed to the phenyl-propionate.27

Unsaturated side chains of phenols are saturated at 270 to 300° without modification of the nucleus.²⁸

Acetone yields 65% of isopropyl alcohol at 280-300°.

Pinene is transformed to pinane, while camphene gives two hydrides, a solid melting at 66° and a liquid boiling at 162°.29

The sodium salts of the two naphthalic acids act differently when hydrogenated with copper at 300° under 100 atmospheres. The α acid furnishes tetrahydronaphthalens directly, while the β leads first to the tetrahydro-naphthalic acid and then to decahydronaphthalens.

Use of Other Metals

595. Zinc powder can cause the reduction of acetone to the alcohol with a yield of 50%.

By using palladium, reduced from the chloride by formates, in the proportion of 1 g. to 30 g. of the substance to be hydrogenated under 110 atmospheres at 110°, methyl-ethyl-acroleine, C₂H₅.CH: CH(CH₂)-CHO, is transformed in 2 or 3 days to methyl-pentanol.

Mesityl oxide is changed in 2 days at 110° to methyl-isobutyl-ketone. By continuous shaking at 110°, citral is reduced to the decanol with a little of the decane. The same may be said of geraniol.

Acetyl-acetone, under 116 atmospheres at 109° is changed to pentanediol in six hours.

Carbohydrates dissolved in aqueous alcohol are changed to the

- ²⁴ IPATIEF, J. Russian Phys. Chem. Soc., 38, 75 (1906) and 39, 681 (1907), C. A., 1, 2877.
 - * IPATIEF, J. Russian Phys. Chem. Soc., 40, 489 (1908), C., 1908 (2), 1098.
 - 26 IPATIEF, Berichte, 43, 3387 (1910).
 - 27 IPATIEF, J. Russian Phys. Chem. Soc., 41, 1414 (1909).
 - ™ IPATIEF, Berichte, 46, 3589 (1913).
- ²⁹ IPATIEF and DRACHUSSOF, J. Russian Phys. Chem. Soc., 42, 1563 (1911), C., 1911 (1), 1292.

corresponding hexites at 110° under 100 atmospheres. Lasruloss yields mannite, glucoss goes into sorbits and galactoss into dulcits.³⁰

IV. HYDROGENATIONS BY NICKEL IN LIQUID SYSTEMS UNDER LOW PRESSURES

596. Very extensive use has been made of the common metals, particularly nickel, for hydrogenation in liquid medium in the case of liquid fats the molecules of which contain ethylene bonds. The description of the methods followed and the results obtained is the special object of the last chapter but the same process can be generalized and extended to a large number of cases. The fundamental condition of success is a sufficiently energetic agitation in the hydrogen. A pressure of several atmospheres is useful but not indispensable, the hydrogenation being capable of being carried out with even reduced pressure. Simply bubbling the hydrogen through the liquid is not sufficient.

Brochet has tried to define exactly the conditions for using this method.³¹

597. Apparatus. Different forms of apparatus may be used according to the amount of the work to be done and the magnitude of the pressure to be used. The pressures run from 1 to 50 atmospheres, being usually around 10 to 15.

A red copper autoclave of 1200 cc. capacity, which can operate satisfactorily with 700 to 800 cc. of liquid, may be used. The bronze cover is fitted accurately and made tight with lead foil packing, being held in place by screw clamps. It is fitted with a thermometer-well dipping into the liquid, a pressure gauge, and a valve for the introduction of the hydrogen. The apparatus is heated electrically by a ferro-nickel coil insulated by asbestos and surrounded by sheet asbestos to keep the heat in. After the introduction of the liquid to be hydrogenated, either alone or in solution, and the addition of the catalyst, the autoclave is closed and connected with the hydrogen tank which is placed along side on the platform of a mechanical shaker. When the operation is finished, the catalyst is filtered off and may frequently be used immediately for another hydrogenation.

Brochet uses a 500 cc. glass cylinder connected with a hydrogen tank by means of a bubble counter which measures the amount of hydrogen absorbed, and enables one to follow the course of the reaction.

⁸⁰ IPATIEF, J. Russian Phys. Chem. Soc., 44, 1002, and 1710 (1912); C. A., 7, 1171, and Berichte, 45, 3218 (1913).

²¹ Brochet, Bull. Soc. Chim. (4), 13, 197 (1913) and 15, 554 (1914).

⁸⁵ A convenient laboratory apparatus with high speed stirring has been described by Reid, J. Amer. Chem. Soc., 37, 2112 (1915).—E. E. R.

213

598. Catalysts. The nickel used is prepared by reducing at about 300° the oxide prepared by calcining the carbonate, nitrate or oxalate. After cooling in a current of hydrogen, the reduced metal is plunged quickly into the liquid to be hydrogenated, avoiding contact with the air as much as possible.

The nickel may be used alone as a metal powder or incorporated with inert materials such as infusorial earth, pumice, or charcoal (126). This incorporation with a carrier is advantageous and gives, on reduction at 450°, a catalyst which is more active than the metal alone reduced at 350°, and a fortiori more active than the metal alone reduced at 450°.

Nickel on a carrier is much less sensitive to toxic agents than nickel alone. Thus for the metal alone, the amount of hydrogen sulphide required to kill the catalyst is 0.02-0.005 g. to 0.5 g. of the catalyst, according to the method of preparation, but may be as high as 0.1 g. for the metal on a porous support.³⁴

We have seen (584) that Ipatief has found it advantageous with his method to use an oxide of nickel, such as NiO or Ni₂O₃, in place of the metal, and that he considers the oxide more active. The same substitution has been proposed for the hydrogenation of oils (943), in which the oxides should show a greater activity and should be less susceptible to the action of poisons, particularly sulphur.²⁵ But in all cases the activity of the oxide may be explained by assuming that it is partially reduced to the metallic state, the metal being more active on account of being formed within the liquid and in a better state of subdivision. This is the opinion of Brochet, who considers the presence of the free metal necessary for hydrogenation but thinks that it is activated by the presence of foreign substances, such as its oxide, or salts or even other metals.²⁶

The presence of metallic nickel in the oxide which is used as catalyst has been denied by Erdmann, who bases his conclusion on the absence of conductivity in the catalyst after it has been freed from fatty material.

At any rate, it is well established that at the temperature at which the hydrogenation of oils is carried on, nickel oxide is reduced to the suboxide, Ni₄O, which is necessarily slowly reduced at these same temperatures to the free metal, the presence of which is easily shown by the direct formation of nickel carbonyl by the action of carbon

^{**} KELBER, Berichte, 49, 55, (1916).

²⁴ Kelber, Berichte, 49, 1868 (1916).

²⁶ BEDFORD and ERDMANN, J. prakt. Chem. (2), 87, 425 (1913).

⁸⁴ Brochet, Bull. Soc. Chim. (4), 15, 770 (1914).

monoxide below 100°. Meigen and Bartels, 8 Norman and Pungs, and later Frerichs, who found an appreciable conductivity in the oxide which had served for the hydrogenation of oil, 40 have come to the same conclusion, that is, that the oxide is inactive in hydrogenation, the activity belonging only to the free metal.

Erdmann has claimed that the most active factor in hydrogenation is a suboxide, such as Ni₂O, which would form an unstable hydride

with hydrogen, e.g. ONiH, which is capable of transferring hydro-

gen to the molecules which can take it up. This special aptitude of the suboxide has been claimed by Senderens and Aboulenc, according to whom acetone can be hydrogenated at 110° under 30 atmospheres pressure by the suboxide but not by the metal.⁴¹

The amount of catalyst may be as low as 0.5% of the liquid to be hydrogenated, but it is better to use larger amounts in order to hasten the reaction.

599. Method of Work. It is best to operate at least 20° below the boiling point of the liquid used as solvent so that its vapor will not dilute the hydrogen too much. If substances are hydrogenated without solvent, 100 to 150° is the usual range of temperatures but sometimes from 150 to 200°.

Alcohol, more or less diluted, and acetic acid are the most favorable solvents. Benzene, acetone, ether, and ethyl acetate are not so good, while chloroform is rather harmful.⁴²

The course of the reaction is easily followed, either by the pressure gauge or by the bubble counter, which shows directly the volume absorbed. This enables one to see at what temperature the reaction goes best.

600. Results Obtained. Nitro derivatives are readily changed to the corresponding amines. Azo and hydroazo compounds are split into two amines; but by operating in the presence of caustic soda which moderates the action of the catalyst, it is possible to obtain azoxy, azo, hydrazo and finally amino from aromatic nitro compounds.

- ³⁷ SABATIER and ESPIL, Compt. rend., 158, 674 (1914).
- MEIGEN and BARTELS, J. prakt. Chem. (2), 89, 296 (1914).
- » Norman and Pungs, Chem. Zeit., 39, 29 (1915), C. A., 9, 1552.
- 40 FRERICHS, Arch. Pharm., 253, 512 (1915).
- SENDERENS and ABOULENC, Bull. Soc. Chim. (4), 17, 14 (1915).
- ⁴⁸ In hydrogenating cotton seed oil, 0.1% nickel on a carrier is ample and even 0.01% gives fair results.— E. E. R.
 - 4 KELBER, Berichte, 49, 55 (1916).
 - 44 BROCHET, Bull. Soc. Chim. (4), 15, 554 (1914).

601. Ethylene Double Bonds. These are easily saturated at low temperatures, even in the cold, with the evolution of heat.

A mixture of *ethylens*, with hydrogen in excess, is changed to *ethane* by being passed at atmospheric pressure through a saturated hydrocarbon in which a nickel catalyst is kept in suspension by rapid stirring.⁴⁶

 α -Octens, treated in alcohol solution with 20% of active nickel and hydrogen at 15 atmospheres, is completely changed to octans in the cold. This can be accomplished under atmospheric pressure but takes much longer.⁴⁶

Olsic acid is reduced to stearic acid at 250° with a velocity which is nearly proportional to the pressure of the hydrogen. 47

The aliphatic esters of *oleic acid* are transformed into stearic esters.

The salt formed by combining hot oleic acid with *anilins* is rapidly hydrogenated to a brittle solid melting at 76°.48

Cinnamic acid, in twice its weight of amyl alcohol, is completely changed to phenylpropionic in 45 minutes by 10% of nickel at 100° under 15 atmospheres. The fact that the acid attacks the nickel does not hinder the reaction. However, it is better to use sodium cinnamate in 4 parts of water, which is hydrogenated in the cold.

Methyl cinnamate, dissolved in methyl alcohol, is changed to methyl phenylpropionate 40 in the cold in 3 hours under 15 atmospheres pressure. Under ordinary pressure the action is much slower, the reduction of ethyl cinnamate requiring 7 hours at 70°.50

Ansthol, CH₂O.C₅H₄.CH: CH.CH₂, is rapidly transformed into methoxy-propyl-benzene when treated without solvent with 10% of nickel at 60-80° under 15 atmospheres, but requires 5 times as long at 1 atmosphere.

Isosafrol, dissolved in 3 parts of alcohol with 19% of nickel, adds H₂ in an hour at 65°.

Geraniol and linalool saturate their double bonds, but allyl alcohol does not at 70° under 15 atmospheres, neither does allyl sulphocyanate. Piperonyl-acrilic acid gives piperonyl-propionic acid in the cold under 15 atmospheres.⁵¹

The acetylene triple bond is also saturated without difficulty.

- 45 RATHER and REID, J. Amer. Chem. Soc., 37, 2115 (1915).
- ⁴⁶ Brochet and Bauer, Bull. Soc. Chim. (4), 17, 50 (1915), and Compt. rend., 150, 190 (1914).
 - 47 SHAW, J. Soc. Chem. Ind., 33, 771 (1914).
 - 48 ELLIS and RABINOVITZ, J. Ind. Eng. Chem., 8, 1105 (1916).
 - 49 BROCHET and BAUER, Loc. cit.
 - 50 BROCHET and CABARET, Compt. rend., 159, 326 (1914).
 - BROCHET and BAUER, Bull. Soc. Chim. (4), 17, 50 (1915).

602. Aldshydes and ketones. Aldehydes and ketones are not appreciably hydrogenated under atmospheric pressure. Thus the allyl-ketones dissolved in 5 parts of alcohol and treated at 60° with hydrogen under atmospheric pressure are hydrogenated in several hours to the saturated ketones without affecting the ketone group.

On the contrary, by working under pressure it is possible to change aldehydes and ketones to the corresponding alcohols.⁵²

603. Various Rings. The hydrogenation of the benzene ring or of similar rings is much more difficult to attain and is scarcely realizable except in the case of phenols and of compounds directly related to them.⁵⁴

With ordinary phenol the addition of hydrogen takes place slowly from 50° up and rapidly between 100 and 150° under 15 atmospheres, with complete transformation into cyclohexanol without the simultaneous production of cyclohexanons.

Likewise several hours are sufficient for the hydrogenation of α -and β -naphthols at 150° under 15 atmospheres.

Eugenol, CH₂O(OH)C₆H₂.CH₂.CH: CH₂, adds H₂ rapidly at 60° and 15 atmospheres to form propyl-methoxy-phenol but the ring is not hydrogenated unless the operation is carried on at 150°.

Indigotine. Indigo, dry or in paste, suspended in water containing a little caustic soda (10 g. indigo to 250 c. dilute caustic soda) is reduced at 70° by 5 g. nickel to indigo white in 40 minutes. The same reaction applies to thio-indigo and to malachite green which is reduced to the leuco base. **

Hydrogenations by nascent Hydrogen in Liquid Systems in contact with Metals

- 604. The decomposition of formic acid by the catalytic action of metals of the platinum group provides hydrogen (824) which can be used in the liquid itself to effect hydrogenations. By the use of spongy or colloidal palladium, cinnamic acid can be transformed into phenylacstic or quining into hydroquining. 56
 - ⁵⁶ CORNUBERT, Compt. rend., 159, 78 (1914).
 - BROCHET and CABARET, Compt. rend., 159, 326 (1914).
- ⁵⁴ The hydrogenation of naphthalene is thoroughly described by Shroeter, Annalen, 426, 1, (1922).—E. E. R.
 - ⁵⁵ Brochet, Compt. rend., 160, 306 (1915).
- ⁵⁶ Verein. Chininfabr. Zimmer & Co., German patent, 267,306, 1914, C., 1914 (1), 88.

CHAPTER XIII

VARIOUS ELIMINATIONS

§ I. — ELIMINATION OF HALOGENS

605. The classical method for the elimination of halogens from chlorine, bromine or iodine compounds is treatment with sodium. The presence of benzene or petroleum ether retards this reaction greatly, but ordinary ether and ethyl acetate usually accelerate it. The use of small amounts of acetonitrile greatly facilitates the reaction. Thus sodium does not act on methyl iodide in the cold but the addition of one or two drops of acetonitrile causes an immediate and abundant evolution of ethane, CH₂, CH₂.

The same is true with ethyl, propyl, isopropyl and allyl iodides, trimethylens bromide and benzyl chloride. Ethyl cyanide produces a similar
catalytic effect and propyl cyanide is less effective while benzonitrile
and benzyl cyanide have no such effect.

§ 2. — ELIMINATION OF NITROGEN

606. Diazo Compounds. In many important reactions of aromatic diazo compounds, a molecule of nitrogen is eliminated. Cuprous salts are frequently useful or indispensable catalysts for these decompositions. Copper powder can produce the same effects, doubtless through the initial formation of cuprous compounds.

Diazobenzene hydroxide, C₆H₆.N: N.OH decomposes immediately even at 0° in the presence of copper powder to form phenol and nitrogen. The copper for this purpose is precipitated by zinc dust in a saturated solution of copper sulphate, washed with water and then with a very dilute solution of hydrochloric acid and preserved wet and protected from the air.⁴

607. Hydrochloric acid reacts with diazo chlorides, on boiling, to give the corresponding aromatic chloride, on condition that the de-

¹ Wurtz, Ann. Chim. Phys. (3), 44, 275 (1855).

² Elbs, Synth. Darstel. d. Kohlenst., Leipzig, 1889, 2, 59.

³ Michael, Amer. Chem. Jour., 25, 419 (1901).

⁴ GATTERMANN, Berichte, 23, 1220 (1890).

composition takes place in the presence of copper powder or cuprous chloride. We have:

$$C_0H_6$$
, N_2 , $C_1 + HC_1 = N_2 + C_0H_6$, $C_1 + HC_1$.

The cuprous chloride is used in hydrochloric acid solution.

This action of cuprous chloride has been explained by assuming that it acts in the presence of hydrochloric acid as a reducing agent giving cupric chloride and hydrogen:

and
$$2CuCl + 2HCl = 2CuCl2 + 2H$$
$$2H + C6H5 . N : NCl = C6H5 . NH . NHCl.$$

The hydrazine compound thus formed reduces the cupric chloride:

$$2CuCl2 + C6H5.NH.NHCl = 2CuCl + 2HCl + C6H5.Cl + N2.$$
resensated

The regenerated cuprous chloride repeats the same effects.

- 608. Hydrobromic acid reacts in a similar way on diazonium bromides in the presence of *cuprous bromids*. The cuprous bromide is prepared by warming 20 g. copper turnings with a solution of 12.5 g. copper sulphate and 36 g. potassium bromide in 80 cc. water containing 11 g. sulphuric acid.⁶
- 609. Diazonium salts in water solution with sodium nitrite, in the presence of copper powder or moist cuprous oxide, are transformed into *nitro* compounds (Sandmeyer reaction):

$$C_0H_5 \cdot N_2 \cdot Cl + N_0NO_2 = C_0H_5 \cdot NO_2 + N_0Cl + N_2$$

- 610. Diazonium salts yield the corresponding aromatic isocyanates, C₅H₅.NCO, when treated with potassium isocyanate in presence of copper powder.⁷
- 611. Hydrazine Compounds. Phenylhydrazine is decomposed at 150° into aniline, nitrogen and ammonia, on contact with cuprous chloride, bromide, or iodide:

$$3C_6H_5.NH.NH_2 = 3C_6H_5.NH_2 + N_2 + NH_3$$

The chloride acts more rapidly than the bromide and this more rapidly than the iodide. When more than 1% of the chloride is added, the decomposition is violent and almost explosive. The crystallized compound, CuI.2C₆H₅.NH.NH₂, which may serve as an intermediate step in the catalysis, has been isolated.⁸

- ³ SANDMEYER, Berichte, 17, 1635 (1884).
- Sandmeyer, Berichte, 17, 2652 (1884).
- ⁷ GATTERMANN, Berichte, 23, 1220 (1890).
- ² Arbusow and Tichwinsky, Berichte, 43, 2295 (1910) and J. Russian Phys. Chem. Soc., 45, 69 (1913), C. A., 7, 2225.

The hydrazones derived from hydrazine and saturated cyclic ketones are decomposed, with the evolution of nitrogen, on contact with a small fragment of solid potash.

Cyclohexone hydrazone gives cyclohexane in a violent reaction:

$$CH_{2}$$
. CH_{2} . $CH_{$

In every case the hydrocarbon obtained contains CH_2 in place of the CO of the ketone. Thus hydrazones from the methyl cyclohexones yield methyl cyclohexane, that from camphor furnishes camphane, $C_{10}H_{18}$ melting at 158°, and that from fenchone leads to fenchane, boiling at 151°.

612. 3, 5. — Diphenyl-pyrazoline heated with fragments of potash and platinized porous porcelain decomposes into nitrogen and diphenyl-cyclopropans.¹⁰

$$C_6H_5$$
. CH
 CH_2
 CH_5
 CH_5

§ 3. — SEPARATION OF FREE CARBON

- 613. In many cases the dehydrogenation of hydrocarbons leads to the separation of free carbon and we shall see (Chapter XXI) that various finely divided metals frequently provoke this decomposition. But it is well to consider here a very important reaction which takes place with the separation of carbon from carbon monoxide in contact with certain substances.
- 614. Decarbonization of Carbon Monoxide. In the reduction of the oxides of iron, nickel and cobalt carried on above 400° by carbon monoxide, it has long been known that carbon is deposited and this continues at the expense of the carbon monoxide according to the equation:

$$2CO = CO_2 + C.$$

Mond found that nickel can produce this effect between 350 and 450°.11

Sabatier and Senderens have shown that the reaction takes place with *reduced nickel* above 230°, elevation of temperature accelerating the decomposition of the carbon monoxide. With a layer of nickel

⁹ Kizhner, J. Russian Phys. Chem. Soc., 43, 582 (1911), C. A., 6, 347.

¹⁰ KIZHNER, J. Russian Phys. Chem. Soc., 47, 1102 (1915), C. A., 9, 3051.

¹¹ Mond, Langer and Quincke, Chem. News, 62, 95 (1890).

35 cm. in length and a flow of gas of 25 cc. per minute, the amounts of carbon dioxide formed from 100 cc. of the monoxide were:

At 238	•	1.2	cc.	
250	•	3.8		
275	•	17.9		
285	•	23.2		
300	•	40.5		
320	•	49.0		
349	o and above	50.0	complete	transformation.

The reaction may be complete as can be shown by experiment; and besides, the inverse formation of carbon monoxide from carbon and the dioxide does not begin below 400°. We do not have to any extent:

$$C + CO_2 = 2CO$$
,
nor: $Ni + CO_2 = NiO + CO$.

This would take place no more at higher temperatures, such as 650° and 800°.12

615. Reduced cobalt gives rise to the identical reaction at above 300°.

Finely divided iron, kept at 445° with carbon monoxide for several hours, transforms it completely into carbon dioxide with the deposition of carbon.¹³

Finely divided platinum, reduced copper, and finely divided silver, do not produce a similar effect on carbon monoxide below 450°.

616. The separation of carbon can be explained by assuming the temporary formation of *nickel* or *cobalt carbonyl* which the high temperature decomposes into metal, carbon, and carbon dioxide.¹⁴

But we can explain the phenomenon equally well by the mechanism which is apparent in the case of iron. At low temperatures, iron tends to reduce carbon monoxide to carbon with the formation of ferrous oxide:

$$Fe + CO = FeO + C$$
,

but at a higher temperature, there is the formation of carbon dioxide and iron:

$$FeO + CO = CO_2 + Fe$$
.

The iron thus regenerated can repeat the first reaction. These two successive steps may take place likewise with nickel and cobalt without our being able to perceive the intermediate compound, the oxide, since

¹² SABATIER and SENDERENS, Bull. Soc. Chim. (3), 29, 294 (1903).

¹³ BOUDOUARD, Ann. Chim. Phys. (7), 24, 5 (1901).

¹⁴ BERTHELOT, Ann. Chim. Phys. (6), 26, 560 (1892).

the reduction of the oxide by the carbon monoxide takes place at a temperature lower than that at which the metal reduces the gas, the oxide of the metal can remain only in inappreciable amount. From this it can be seen that the reaction will take place better with nickel than with iron, since a considerable proportion of the iron is actually transformed into the oxide.¹⁵

617. Manganous oxide, which dehydrogenates alcohols after the manner of metals (701), appears to give, doubtless by a mechanism analogous to that which has just been described, a certain amount of decomposition of carbon monoxide into carbon and carbon dioxide, but it is always small below 350°.16

§ 4. — ELIMINATION OF CARBON MONOXIDE

618. The decomposition of aldehydes and ketones can take place as a consequence of the elimination of carbon monoxide under the influence of catalysts, either finely divided metals or anhydrous oxides acting at higher temperatures.

With aldehydes the reaction goes more readily and yields chiefly:

$$R.CO.H = CO + RH.$$

619. Reduced nickel acts energetically above 200°. The vapors of propionic aldehyde are rapidly dissociated at 235° into carbon monoxide and ethane. Benzaldehyde is largely decomposed at 220° into benzene and pure carbon monoxide. 17

Furfural is changed by nickel at 270° into furfurane13:

620. With ketones the result is more difficult to obtain. Starting with a ketone R.CO.R' a certain amount of the hydrocarbon R.R' may be formed but the débris resulting from the groups R and R' are the chief products.

Actions is decomposed by nickel, slowly at 240° and rapidly at 270°, yielding carbon monoxide and the CH_s radicals which give a little ethane and ethylene but chiefly methane, hydrogen and carbon. ¹⁷

621. Reduced copper has less effect: at 310° its action on propionic aldehyds is negligible and it is only at 350° or better at 400° that

¹⁵ SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 485 (1905).

¹⁶ SABATIER and MAILEE, Ann. Chim. Phys. (8), 20, 315 (1910).

¹⁷ Sabatier and Senderens, Ann. Chim. Phys. (8), 4, 474 (1905).

¹³ Padoa and Ponti, Lincei, 15 (2), 610 (1906), C., 1907 (1), 570.

carbon monoxide and a mixture of ethane, hydrogen and butane is obtained.¹⁷ Its action is energetic on formaldshyds which it decomposes almost completely into carbon monoxide and hydrogen.¹⁹ The resulting carbon monoxide can be absorbed by caustic soda present in the mixture and furnish, according to a well known reaction, sodium formate.²⁰

Copper has no appreciable effect on ketones below 400°.

622. Platinum sponge and particularly platinum black have an intense destructive action on aldehydes. Propionic aldehyds is attacked at 225°, and at 275° decomposes rapidly into the same gaseous products as are obtained with copper.²¹

The action on ketones is less intense.

- 623. At 300° palladium black decomposes formaldehyde completely into carbon monoxide and hydrogen with traces of carbon dioxide and methane. Likewise acetaldehyde, propionic aldehyde, butyric aldehyde, benzaldehyde, and the toluic aldehydes are more or less split at temperatures around 300° into carbon monoxide and the corresponding hydrocarbons.²²
- 624. The decomposition of formic acid into carbon monoxide and water which is effected by certain oxides, titania, blue oxide of tungsten, alumina, silica, and zirconia, and which can be regarded as an elimination of carbon monoxide, will be studied later (825), as also the decomposition of formic esters, which is chiefly according to this reaction (866):

$$H.CO_2C_nH_{2n+1} = CO + \underbrace{C_nH_{2n+1}.OH.}_{alcohol}$$

625. Anhydrous aluminum chloride can decompose acid chlorides with the elimination of carbon monoxide. This takes place with dichloracetyl chloride which is split into carbon monoxide and chloroform with heptachlorpropans as a by-product, resulting from the action of the chloroform on the original product.²³

§ 5. — ELIMINATION OF HYDROGEN SULPHIDE

626. Mercaptans. Cadmium sulphide catalyzes the decomposition of mercaptans according to two consecutive reactions exactly analogous to those according to which a primary alcohol is dehydrated to an ether and then to an unsaturated hydrocarbon (701).

- 19 SABATIER and MAILHE, Ann. Chim. Phys. (8), 20, 345 (1910).
- ²⁰ LOEW, Berichte, 20, 145 (1887).
- 21 SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 475 (1905).
- ** KUENEZOV, J. Russian Phys. Chem. Soc., 45, 557 (1913).
- ²⁸ PRINS, J. prakt. Chem. (2), 89, 414 (1914).

At a moderate temperature, we have:

$$2C_nH_{2n+1}.SH = H_2S + \frac{(C_nH_{2n+1})_2S}{\text{sulphide}}$$

At a higher temperature, a more rapid decomposition yields hydrogen sulphide and the ethylene hydrocarbon:

$$C_nH_{2n+1}.SH = H_2S + C_nH_{2n}.$$

Thus ethyl mercaptan, C₂H₆.SH, passed over cadmium sulphide at 320°, is almost completely transformed into the neutral sulphide, (C₂H₆)₂S, while at 380° it is completely decomposed into hydrogen sulphide and ethylene.

Isoamyl mercaptan is changed into isoamyl sulphide at 360°, but above 400° gives hardly anything but amylene.

The decomposition of *primary aliphatic mercaptans* over cadmium sulphide at regulated temperatures constitutes a regular method of preparing primary sulphides from the mercaptans.

627. The mechanism of the decomposition is altogether analogous to that of alcohols (169). We can assume the formation of a cadmium mercaptide from the mercaptan and cadmium sulphide. This would decompose, according to the temperature, either into the neutral sulphide or into the ethylene hydrocarbon with the regeneration of the metal sulphide which would then repeat the reaction, thus playing the part of a catalyst.

We have at first:

$$CdS + 2C_nH_{2n+1}.SH = \underbrace{(C_nH_{2n+1}S)_2Cd}_{mercaptide} + H_2S$$

then:

$$(C_nH_{2n+1}S)_2Cd = CdS + \underbrace{(C_nH_{2n+1})_2S}_{\text{sulphide}}$$

and at a higher temperature:

$$(C_nH_{2n+1}S)_2Cd = CdS + H_2S + 2C_nH_{2n}.$$

The transitory formation of the cadmium mercaptide is further indicated by the change of color of the sulphide, which takes on an orange tint quite different from the bright yellow of the original sulphide and retains that color after cooling in consequence of the persistence of a certain amount of the mercaptide.

628. Secondary mercaptans have a stronger tendency to decompose into the ethylene hydrocarbons but can, nevertheless, furnish some neutral sulphide.

Cyclohexyl mercaptan, passed over cadmium sulphide at 300°, gives

12 to 15% of the sulphide but the major portion is decomposed to cyclohexene, while at 350° all of it goes into cyclohexene.*

629. Thiophenols. Aluminum chlorids acts on a warm solution of thiophenol in petroleum ether, eliminating hydrogen sulphide and forming the diphenyl sulphide. At the same time some thianthrene,

630. Formation of Thioureas. The thioureas can be obtained by the reaction of primary aromatic amines on carbon disulphide in the presence of a little sulphur as catalyst.

Thus 1 part each of aniline, alcohol, and carbon disulphide and 0.005 part crystallized sulphur are warmed for several hours on the steam bath to obtain symmetrical diphenyl-thiourea:

$$CS_2 + 2C_0H_5NH_2 = H_2S + CS NH.C_0H_5$$

NH.C₀H₅

The ortho and para toluidines, the naphthyl-amines and even p. aminophenol give the same reaction.²⁶

§ 6. — ELIMINATION OF AMMONIA

631. Reduced nickel has various effects on primary, secondary, and tertiary amines, and among these effects one is the elimination of ammonia.²⁷

Above 300° this is a clean cut reaction with aliphatic amines containing less than 5 carbon atoms. Thus *ethylamins* splits up into ammonia and *ethylens*, which at that temperature is in turn decomposed into carbon, methane, hydrogen and ethane (910). We have:

$$C_2H_5.NH_2 = NH_3 + C_2H_4.$$

Amines containing five or more carbon atoms, e.g. amyl-amine, undergo this reaction and are simultaneously dehydrogenated to nitriles. This is true for benzyl-amine also.²⁸

Aromatic amines, aniline and the toluidines, are much more resistant, being hardly attacked by nickel at 350°, but towards 500° there is elimination of ammonia with complete destruction of the molecule, according to a complicated reaction.²⁹

- ²⁴ Sabatier and Mailhe, Compt. rend., 150, 1570 (1910).
- * DEUSS, Rec. Trav. Chim. Pays-Bas, 27, 145 (1908).
- ²⁴ Hugershoff, Berichte, 32, 2245 (1899).
- ²⁷ Sabatier and Gaudion, Compt. rend., 165, 309 (1917).
- ³⁸ Sabatier and Gaudion, Compt. rend., 165, 226 (1917).
- 29 SABATIER and GAUDION, Compt. rend., 165, 309 (1917).

632. By heating α -naphthylamins for 8 hours with a molecule of anilins in presence of a small amount of iodine (less than 1%), ammonia is eliminated and phenyl-naphthylamins is formed in 85% yield.

The use of very small proportions of iodine enables us to prepare the secondary amines derived from α -naphthylamine and the three toluidines, ortho and para anisidines, and meta and para chloranilines, with yields superior to those obtained by the usual methods.

By heating β -naphthylamins on the steam bath for 4 hours with less than 1% of iodine, it is almost quantitatively changed into $\beta\beta'$ -dinaphthyl-amine.

Likewise p.aminophenol heated below 200° for 5 hours with 0.0025% iodine, loses ammonia and yields about 70% pp'.dihydroxy-diphenylamine.³⁰

633. Cuprous chloride and bromide and also zinc chloride catalyze the decomposition of the phenylhydrazones derived from the lower aliphatic aldehydes and ketones, giving ammonia and substituted indols.

Thus the phenylhydrazone of *methyl-ethyl-ketone* evolves ammonia at 180° when 0.2% of cuprous chloride is added and yields 2, 3-dimethyl-indol in 2 hours.

The phenylhydrazone of propionic aldshyds gives similarly 3-methyl-indol (skatol). With copper chloride the yield is 60% and reaches 73% with zinc chloride.

With aldehydes there is some formation of nitriles resulting from splitting off aniline (635).

The formation of β -methyl-indol is thus represented:

In the same way, the phenylhydrazone of acstonyl-acstons yields dimsthyl-amino-phenyl-pyrrol.³¹

With the phenylhydrazones derived from higher aliphatic aldehydes, this reaction is of little importance as it is overshadowed by the formation of nitriles (635).

³⁰ Knoevenagel, J. prakt. Chem. (2), 89, 20 (1914).

²¹ Arbusof and Tikhvinsky, J. Russian Phys. Chem. Soc., 45, 73 (1913), C. A., 7, 2225. — Arbusof and Friauf, Ibid., 45, 694 (1913), C. A., 7, 3599. — Arbusof and Khrutzkii, Ibid., 45, 699 (1913), C. A., 7, 3599.

§ 7. — ELIMINATION OF ANILINE

634. The stability of aniline in the presence of nickel, which has been mentioned above, enables us to predict that the action of nickel on alkyl anilines will tend to split off aniline, as ammonia is eliminated from the alkyl ammonias. This is what takes place with methylaniline at 250°. Aniline is regenerated with the separation of the group CH₂ which decomposes into methane and carbon, the reaction being nearly thus:

$$2C_6H_6.NH.CH_3 = 2C_6H_6.NH_2 + C + CH_6.$$

With sthyl-anilins, we have:

$$C_0H_5.NH.C_2H_5 = C_0H_5.NH_2 + C_2H_4$$

the ethylene being entirely decomposed by the nickel (912) into carbon, methane, ethane and hydrogen, the hydrogen acting on the aniline to give a little ammonia and benzene.

Dimethyl-aniline and disthyl-aniline behave in an entirely analogous manner.22

635. The phenylhydrazones of higher aliphatic aldshydes are decomposed by copper, zinc, and platinum chlorides into nitriles and anilins:

$$R.CH: N.NH.C_6H_5 = R.CN + C_6H_5.NH_5.$$

This is true for isobutyric, isovaleric, and isoheptylic aldehydes. The simultaneous production of indols (633) is of little importance.²³

- * SABATIER and GAUDION, Compt. rend., 165, 309 (1917).
- * Arbusof, J. Russian Phys. Chem. Soc., 45, 74 (1913).

CHAPTER XIV

DEHYDROGENATION

636. We have explained direct hydrogenation by means of finely divided metals by the formation of an unstable hydride, produced rapidly by the metal and capable of readily giving up its hydrogen. If this explanation is correct, an important consequence can be readily foreseen. The catalytic metals, nickel, copper, and platinum, should be able to take up hydrogen not only from molecules of free hydrogen but also from other substances capable of furnishing hydrogen, and consequently to be dehydrogenation catalysts, a prediction which experiment has largely verified.

637. This capability has been long known in some cases. As early as 1823 it was known that *iron*, *copper*, *gold*, *silver*, and *platinum* had the power of greatly facilitating the decomposition of ammonia, without appreciable alteration of the metal. The decomposition of the ammonia can be thus effected at a much lower temperature than in the absence of these metals.¹

In 1843, Reiset and Millon noticed that alcohol vapor passed through a tube filled with fragments of porcelain and heated to 300°, is not appreciably decomposed, but that decomposition manifests itself at 220° in presence of *platinum sponge*.²

In 1866, Berthelot noticed that the presence of iron favors the decomposition of acetylens at a red heat,³ and later Schützenberger stated that platinum spongs warmed in a current of acetylene, decomposes it with incandescence, giving a voluminous mass of carbon in which the metal is diffused.⁴

This active decomposition of acetylene was rediscovered in 1896 by Moissan and Moureu, who observed it also with recently reduced iron, cobalt, and nickel.⁵ A similar decomposition of *ethylens* in contact with the same metals at 300°, was obtained in 1897 by Sabatier and Senderens,⁶ who interpreted it by assuming the temporary for-

- ¹ Dulong and Thenard, Ann. Chim. Phys. (2), 23, 440 (1823).
- ³ REISET and MILLON, Ann. Chim. Phys. (3), 8, 280 (1843).
- ³ Berthelot, Compt. rend., 62, 906 (1866).
- 4 SCHÜTZENBERGER, Traité de Chemie, I, 724.
- · Moissan and Moureu, Compt. rend., 122, 1241 (1896).
 - SABATIER and SENDERENS, Compt. rend., 124, 616 (1897).

mation of a metal hydride and were thus led to apply these metals to dehydrogenation reactions as well as to those of hydrogenation.

638. The dehydrogenation catalysts are primarily the metals, and to a less degree, certain anhydrous metal oxides and some salts derived from these oxides, carbon and, in exceptional cases, anhydrous aluminum chloride.

The effects produced by these catalysts can be divided into several groups:

- 1. Dehydrogenation of hydrocarbons.
- 2. Return of hydroaromatic compounds to aromatic with double bonds.
- 3. Conversion of primary alcohols to aldehydes and of secondary to ketones.
 - 4. Dehydrogenation of poly-alcohols.
 - 5. Dehydrogenation of amines to nitriles.
 - 6. Direct synthesis of amines from hydrocarbons.
 - 7. Formation of rings by loss of hydrogen.

§ 1. — DEHYDROGENATION OF HYDROCARBONS

639. Finely divided metals exercise an important dehydrogenating-effect on hydrocarbons, the effect being greater the higher the temperature. The separation of hydrogen is always accompanied by molecular changes, which are frequently followed by condensation into more complex hydrocarbons. We will return to the breaking down and building up of hydrocarbons by catalysts in Chapter XXI, which is devoted to that subject, and will content ourselves in the following paragraph to the regular passage of hydroaromatic hydrocarbons to the aromatic with double bonds.

§ 2. — DEHYDROGENATION OF HYDROAROMATIC COMPOUNDS

640. The various compounds formed by the hydrogenation of stable cyclic compounds tend to revert to the latter by loss of hydrogen when submitted to the action of finely divided metals at temperatures higher than those at which they are formed directly. Among the metals, reduced nickel shows itself as particularly active.

The dehydrogenation can take place in the presence of excess of hydrogen, and in some cases the excess of hydrogen, far from hinder-

⁷ This is probably a reversible reaction reaching a definite equilibrium for each temperature and pressure of hydrogen. Quantitative studies are most desirable. — E. E. R.

ing the reaction, regulates it by favoring the maintenance of the cyclic structure and diminishing the tendency to the breaking up of the molecule into many fragments (644).

641. Cyclohexane, which can not be formed by the direct hydrogenation of benzene by the aid of nickel above 300° (446), suffers a partial dehydrogenation to benzene above 300°, but a part of the benzene is transformed to methane by the liberated hydrogen:

$$3C_6H_{12} = 2C_6H_6 + 6CH_4$$

The presence of a current of hydrogen stabilizes the molecule to a certain extent so that it is only slightly broken up at 350°. At 400° about 30% of the cyclohexane passing over the nickel with the hydrogen is decomposed into benzene.

With methyl-cyclohexane alone, decomposition begins at 240° and is rapid at 275°, the gas evolved then containing:

The condensed liquid contains a large proportion of toluene.

Ethyl-cyclohexans is attacked slowly at 280 to 300° and gives a gas containing 83% methane and 17% hydrogen, a mixture of sthylbenzens and toluens being condensed.

The 1, 3-dimethyl-cyclohexane acts like cyclohexane and is stabilized by an excess of hydrogen. At 400° , the dehydrogenation to m.xylene does not exceed 25 %.10

Reduced copper exercises a similar but less intense action which does not begin till above 300°.

642. Hydroxy and amino substitution products of cyclohexane hydrocarbons undergo dehydrogenation still more readily and above 350° the reaction is not hindered by an excess of hydrogen.

In the presence of nickel above 350°, cyclohexanol and its homologs come back to the phenol condition. This effect commences at even much lower temperatures: when cyclohexanone is hydrogenated over nickel at 230°, 25% of phenol is collected along with the cyclohexanol.¹¹ In a current of hydrogen at 360° the transformation into phenol is practically complete.¹²

The same effect is even more important for the cyclic poly-alcohols and also for the amines such as cyclohexyl-amins which tends to regen-

- * SABATIER and MAILHE, Compt. rend., 137, 240 (1903).
- SABATIER and DAUDIER, Compt. rend., 168, 670 (1919).
- 19 SABATIER and GAUDION, Unpublished results.
- ¹¹ SKITA and RITTER, Berichte, 44, 668 (1911).
- ¹⁹ Padoa and Fabris, *Linesi*, 17 (1), 111 and 125 (1908), C., 1908 (1), 1395 and 1908 (2), 1103.

erate aniline and dicyclohexyl-amine which yields diphenylamine and cyclohexylaniline.

The hydrides of naphthalene act in the same way: the higher hydrides under the influence of nickel at 200° come back to the tetrahydride, and this regenerates naphthalene at 300°.

Frequently, as in the case of cyclohexane, the liberated hydrogen can break down a portion of the hydrocarbon into larger or smaller aliphatic fragments. This takes place with dodecahydrophenanthrens, which breaks down at 200° into lower hydrides and various aliphatic hydrocarbons, while the hexahydride is regularly dehydrogenated to the tetrahydride at 220°, which in turn passes to phenanthrene at 280°.

With nickel at 300-330°, the perhydrides of anthracene give the tetrahydride and decomposition products

At 250°, decahydrofluorene returns to fluorene.

643. Unsaturated cyclic hydrocarbons, cyclohexenes, cyclohexadienes, as well as the terpenes and various of their substitution products, are still more readily dehydrogenated by nickel even in a current of hydrogen.

Cyclohexene gives benzene almost quantitatively when passed over nickel at 250°. The same is true at 300° in a current of hydrogen. 18

Cyclohexadiene, C₆H₈, passed over finely divided platinum at 180°, yields benzene, but this is mixed with cyclohexane, which is stable at this temperature and which results from the utilization of the liberated hydrogen.¹⁴

644. Limonene, in a current of hydrogen over nickel at 280-300°, is changed almost entirely into *cymens* accompanied by a certain amount of *cumens* and simpler aromatic hydrocarbons.

Menthens, in hydrogen over nickel at 360°, yield 80% of cymene. Under the same conditions, pinens and camphens are dehydrogenated to aromatic hydrocarbons, C₁₀H₁₄ and lower.¹⁵

along by a current of hydrogen over nickel at 360° is simultaneously reduced and dehydrogenated to form *cymens*.

Terpinsol undergoes a similar reaction.

Pulsgons,
$$CH_3$$
. CH_2 . CH_2 . CC : CH_3 , submitted to the action

- 18 SABATIER and GAUDION, Compt. rend., 168, 670 (1919).
- 14 Böeseken, Rec. Trav. Chim. Paye-Bas, 37, 255 (1918).
- ¹⁵ SABATIER and GAUDION, Compt. rend., 168, 670 (1919).

of nickel in a current of hydrogen at 360°, is changed into a mixture of *thymol* and *cresol*, formed by the elimination of the carbon chain in the form of methane.¹⁵

646. Dodecahydrotriphenylene is completely changed to triphenylene, melting at 198°, by passing over copper at 450-500°. 16

647. Piperidine, under the action of nickel at 180 to 250°, even in the presence of hydrogen, is totally changed to pyridine:¹⁷

$$\begin{array}{c|c} CH_2.CH_2 \\ \hline CH_2.CH_2 \\ \hline \\ NH \rightarrow CH \\ \hline \\ CH:CH \\ \hline \\ N. \end{array}$$

Tetrahydroquinoline, passed over nickel at 180°, gives a certain proportion of quinoline, but the chief product is skatol:18

648. If dehydrogenation is carried out with a partially hydrogenated product, the hydrogen set free by the action of the metal on one portion may hydrogenate the other. This is what takes place when palladium sponge acts on methyl tetrahydroterephthalate which gives 1 part methyl terephthalate and 2 parts methyl hexahydroterephthalate.¹⁹

649. Palladium black is an active dehydrogenation catalyst for the hexamethylene hydrocarbons. The action begins at 170°, is vigorous at 200°, at a maximum at 300°, and yields only hydrogen and benzene or its homologs. At 100-110°, the inverse action takes place, i.e. there is hydrogenation of the benzene, but this does not take place at 200° even in excess of hydrogen. Likewise hexahydrobenzoic acid passes to benzoic.²⁰ The esters of hexahydrobenzoic acid are also dehydrogenated, but methyl cyclopentane-carbonate is not affected.²¹

- ¹⁶ Mannich, Berichte, 40, 159 (1906).
- ¹⁷ CIAMICIAN, *Lincei*, 16, 808 (1907).
- ¹⁸ Padoa and Scagliarini, Linesi, 17 (1), 728 (1908), C., 1908 (2), 614.
- 19 ZELINSKY and GLINKA, Berichte, 44, 2305 (1911).
- 26 ZELINSKY and MISS UKLONSKAJA, Berichte, 45, 2677 (1912).
- ²¹ Zelinsky and Miss Uklonskaja, J. Russian Phys. Chem. Soc. 46, 56 (1913), C. A., 7, 2224.

Below 300°, cyclopentane and methyl-cyclopentane and cycloheptane are not dehydrogenated.

Platinum black acts similarly but less energetically.22

§ 3. — DEHYDROGENATION OF ALCOHOLS

650. A long time ago Berthelot noticed that the vapors of ethyl alcohol passed through a progressively heated glass tube, begin to decompose at around 500°, that is at nearly a dull red heat, giving rise to two simultaneous reactions, namely: dshydration with separation of ethylene and dshydrogenation with the production of aldehyde, the reactions being further complicated by the decomposition of the ethylene and the aldehyde by the heat, the aldehyde being partially decomposed into carbon monoxide and methane.²⁴

Various *primary* alcohols undergo analogous decompositions at a dull red heat, being simultaneously dehydrated and dehydrogenated. We have:

$$\begin{array}{c} C_{n}H_{2n+1}.CH_{2}.CH_{2}OH \\ & \xrightarrow{\text{ethylene hydrocarbon}} \\ H_{2} + \underbrace{C_{n}H_{2n+1}.CH_{2}.CO.H}_{\text{aldehyde}} \end{array}$$

and likewise:

$$C_6H_5.CH_2OH$$

bensyl alcohol

 $H_2 + C_6H_5.CO.H$

bensaldshyde

Up to 400°, neither of these reactions takes place to any appreciable extent.

Secondary alcohols react more readily in this manner, giving hydrocarbons by dehydration and ketones by dehydrogenation, the one or the other reaction predominating as the case may be. Thus, for secondary aliphatic alcohols, ethylene hydrocarbons are formed rather than ketones, while benzhydrol yields benzophenone at as low as 290°.25

- 651. In the presence of catalysts, that is to say of substances capable of forming temporary chemical combinations with one of the products of the above reactions, the corresponding reaction will be realized at a lower temperature and rendered more or less rapid.
- ZELINSKY, J. Russian Phys. Chem. Soc., 43, 1220 (1911). Berichte, 45, 3678 (1912).
 - 2 Zelinsky and Herzenstein, J. Russian Phys. Chem. Soc., 44, 275 (1912).
- ²⁴ Berthelot and Jungfleisch, Traité élém. de Chimie Org., 2nd. Ed. Paris, 1886, I, 256.
 - * Knoevenagel and Heckel, Berichte, 36, 2816 (1903).

Dehydrogenation catalysts should specially promote the decomposition of alcohols into aldehydes or ketones, while dehydration catalysts should facilitate the formation of water and hydrocarbons.

The metals, copper, cobalt, nickel, iron, platinum, and palladium, particularly in the finely divided form, are dehydrogenation catalysts, and so are a small number of anhydrous oxides, e.g. manganous, though to a less extent.

On the contrary, certain metal oxides are exclusively dehydration catalysts for alcohols: such are thoria, alumina and the blue oxide of tungsten.

Finally a large number of substances, oxides and salts, have both functions and can to very variable extents cause the dehydration and the dehydrogenation of alcohols at the same time. *Beryllia* and *zirconia* play the two rôles almost equally well; all the intermediates are found between the two extremes of exclusive catalysts.²⁶

652. Of all the dehydrogenation catalysts, the one that serves best for the regular decomposition of primary or secondary alcohols into aldehydes or ketones, is *reduced copper*, which in practice can be replaced by the very finely divided copper which is manufactured fo imitation gilding.

Cobalt, iron, and platinum can be used, but with poorer results, while nickel is the least suitable.

Use of Copper

653. Primary Alcohols. Primary aliphatic alcohols, when passed in the vapor form over reduced copper kept between 200 and 300°, are regularly decomposed into aldehydes and hydrogen, the condensate containing, along with the aldehyde, some of the unchanged alcohol and a little of the corresponding acetal. The practical yield is usually above 50% with less than 5% of higher products and 45% of the alcohol which can be fractioned out and put through again. This is a very advantageous method for the preparation of aliphatic aldehydes, particularly for those which, on account of low volatility, are difficult to prepare by oxidation of the alcohols.

The transformation can never be complete, even when a long train of copper is used, since the hydrogen which is formed can be added to the aldehyde by copper above 200°. Hence the reaction is limited but the conditions are favorable to the decomposition because the operation is carried on in the presence of a small concentration of hydrogen.

By operating under reduced pressure, there is the double advantage of a more ready volatilization of the alcohols and a diminution

²⁶ Sabatter and Mailie. Ann. Chim. Phys. (8), 20, 289 and 341 (1310).

²⁷ SABATIER and SENDERENS, Compt. rend., 136, 738, 921 and 983 (1903).

of the reverse action of hydrogen, and consequently increasing the practical yield.

654. The apparatus used by Sabatier and Senderens is the same as that employed for hydrogenations (347) except that the tube for introducing the hydrogen is omitted.²⁸

Bouveault has used a vertical tube for the catalyst, 25–30 mm. in diameter and of varying length, up to 1 m. The lower extremity which is drawn down to 10 mm. passes through the stopper of a flask in which the alcohol is vaporized. The tube is filled with rolls of copper gauze containing copper hydroxide, resembling cigarettes; it is heated by a coil of resistance wire through which passes a current that can be suitably regulated. The reduction of the copper hydroxide is effected by hydrogen at 300° and should be carried on slowly so as to leave an adherent mass of copper.

The current is regulated so as to obtain the desired temperature and the alcohol vapors pass through the vertical catalyst tube and from it into a fractionating column which separates the more volatile aldehyde and returns the less volatile alcohol to the flask to be revaporized. A catalyst tube 1 m. long is sufficient for the preparation of 500 g. aldehyde in a day.^{29 30}

It is evident that the apparatus may be connected with a pump controlled by a regulator so as to operate in a partial vacuum, if this is desired.

655. If the temperature is above a certain point, the aldehydes formed are partially destroyed by contact with the metal with elimination of carbon monoxide:

$$R.CO.H = CO + RH.$$

But except in the case of formaldehyde and the aromatic aldehydes, this decomposition is not yet rapid at 300°.

This decomposition is more rapid with a more active catalyst. With methyl alcohol, using a light violet copper prepared by the slow reduction of the precipitated oxide, there is a rapid evolution of gas which contains about 1 volume of carbon monoxide to 2 of hydrogen: the formaldehyde produced has been completely destroyed, only traces of it being found in the condensate. We have:

$$H.CH_2.OH = CO + 2H_2.$$

On the contrary with compact reddish orange copper, prepared by reducing a dense oxide at a dull red, the evolution of gas is only about

- 28 SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 332 (1905).
- 29 BOUVEAULT, Bull. Soc. Chim. (4), 3, 50 and 119 (1908).
- ³⁰ This apparatus and its operation are more fully described by Weismann and Garrand, J. Chem. Soc., 117, 328 (1920). E. E. R.

one twelfth as rapid, but it is practically pure hydrogen and almost all of the formaldehyde survives.^{\$1}

656. Methyl alcohol is decomposed even at 200° and very rapidly at 280-300°.

By catalytic decomposition over copper, methyl alcohol can be detected in ethyl alcohol, since the formaldehyde produced can be characterized by the violet coloration which it gives with morphine and concentrated sulphuric acid.²²

The destruction of the formaldehyde is already apparent at 240–260°, hydrogen and carbon monoxide being produced along with a little methyl formate (225),²⁵ this destruction increasing rapidly with rise of temperature, till at 400° at least 75% is decomposed.

Ethyl alcohol is decomposed above 200°, the aldehyde being formed rapidly at 250 to 350°, without complications. At 420°, 16% of the acetaldehyde is destroyed and the gas collected contains 3 volumes of methane and 1 of carbon monoxide to 6 of hydrogen.³⁴

Propyl alcohol is transformed regularly at 230 to 300° and at 420° one fourth of the aldehyde is destroyed.

Butyl alcohol yields the aldehyde well at 220 to 280°, and at 370° only one sixth is destroyed.

At 240 to 300°, isobutyl alcohol is easily transformed into the aldehyde: at 400°, one half of this is decomposed.

Isoamyl alcohol yields the aldehyde at 240 to 300° without complications. At 370° only 6% of the product is decomposed and at 430°, about 25%.²⁵

An aliphatic C₁₀ alcohol is regularly changed into the aldehyde by heating in Bouveault's apparatus under reduced pressure.²⁶

The copper is never fouled by carbonaceous deposits and remains able to continue the reaction indefinitely.

657. Benzyl alcohol is transformed less readily than the aliphatic: the decomposition does not begin below 300° but is satisfactory there. At 380° the reaction is complex and some toluene and benzene are formed along with the benzaldshyds, while the gases evolved contain carbon monoxide and dioxide along with the hydrogen. From 18 parts of alcohol, only 13 go to the aldehyde, the other 5 forming benzene and toluene.

Under reduced pressure, phenylethyl alcohol, CoHs. CH2. CH2OH,

²¹ SABATTER and MAILHE, Ann. Chim. Phys. (8), 20, 344 (1910).

MANNICH and GEILMANN, Arch. Pharm., 254, 50 (1916), C. A., 11, 1114.

Mannich and Geilmann, Berichte, 49, 585 (1916).

⁸⁴ SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 463 (1905).

^{*} SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 463 (1905).

³⁶ Bouveault, Bull. Soc. Chim. (4), 3, 50 and 119 (1908).

yields phenyl-acstaldshyds readily, but there is a little decomposition of the aldehyde into toluens and carbon monoxide and there is also some dehydration of the alcohol to styrens, C₈H₈.CH: CH₃, the major part of which is hydrogenated to sthyl-benzens or condensed to the slightly volatile meta-styrens which remains on the metal and weakens its catalytic activity.

658. The unsaturated allyl alcohol, CH₂: CH.CH₂OH, is transformed over copper at 180 to 300°, with the evolution of very little hydrogen, into propionic aldehyds, with a slight amount of acroleine. The hydrogen derived from the decomposition of the alcohol serves to hydrogenate the double bond of the aldehyde formed (432).²⁵

It is the same way with undecenyl alcohol, CH₂: CH.(CH₂)₈.-CH₂OH, which yields only the saturated aldehyde, undecenal. On the contrary, under reduced pressure, geraniol (416) gives citral almost entirely.³⁶

659. Secondary Alcohols. The transformation of secondary alcohols into ketones with the separation of a molecule of hydrogen is even more readily accomplished by finely divided *copper* since, the ketones being more stable than the aldehydes, a larger temperature interval is available in which to effect the transformation. Usually even at 400° there is no appreciable complication, the gas evolved is pure hydrogen. The immediate yield of ketone may exceed 75%.

As in the case of the aldehydes, the reaction is never entirely complete, since, in contact with copper above 200°, the disengaged hydrogen is capable of hydrogenating the ketone to regenerate the alcohol. But the hydrogenating power of the copper is much less than its aptitude to decompose the alcohol and the production of ketone predominates greatly.*

Isopropyl alcohol is decomposed slowly from 150°, the production of acetons being rapid at 250 to 430°, without separation of propylene.

Secondary butyl alcohol is attacked at 160°, and furnishes butanons readily at 300° without production of butylens.

Secondary octyl alcohol produces only the octanons (2) at 250 to 300°. It is only above 400° that there is decomposition into carbon monoxide and hydrocarbons.

660. Over copper at around 300°, cyclohexanol is split cleanly into hydrogen and cyclohexanone.

At 300°, o.methyl-cyclohexanol is transformed into o.methyl-cyclohexanone, with a little water and o.methyl-cyclohexene and some o.cresol which are readily eliminated. Results almost as good are obtained with the meta but less satisfactory with p.methyl-cyclohexanol.

⁸⁷ SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 467 (1905).

The method may be used with the same facility with the various dimethyl-cyclohexanols.²⁸

- 661. By contact with copper at 300°, bornsol is changed very readily and almost totally into camphor.³⁰
- 662. Benzhydrol. C₆H₆.CH(OH).C₆H₅, when its vapors are passed over copper at 350°, yields benzophenone, which is largely changed by the liberated hydrogen into diphenyl-methane and particularly into symmetrical tetraphenyl-ethane (720).
- 663. The method is suitable for transforming a secondary alcohol group into a ketone group even in mixed compounds. The secondary alcohol-ketones of the form R.CH(OH).CO.R' readily furnish the corresponding α -diketones.⁴⁰

Under the same conditions, β -hydroxy-esters can be transformed into ketone-esters. Thus ethyl β -hydroxy-isoheptoate, $(CH_2)_2CH.CH_2.$ CH(OH). $CH_2CO_2C_4H_5$, is changed to ethyl β -keto-isoheptoate.

Use of Other Metals

664. Nickel. Reduced nickel acts more violently on the alcohols than does copper and the dehydrogenation of primary or secondary alcohols is always accompanied by a more or less considerable splitting up of the aldehyde or ketone, with the formation of carbon monoxide which may be more or less profoundly altered by the nickel; a part being hydrogenated by the hydrogen formed from the alcohol and a part being changed to carbon and carbon dioxide (614). The separation of the carbon monoxide usually begins at the same time as the decomposition of the alcohol.

Methyl alcohol is attacked as low as 180°, but two thirds of the liberated formaldehyde is destroyed. The reaction is rapid at 250° but eight ninths of the aldehyde is destroyed and the gas evolved contains only 45% of hydrogen along with methane and carbon monoxide. At 350° there is no longer any aldehyde and no carbon monoxide: the gas is a mixture of methane and carbon dioxide.

Ethyl alcohol is decomposed from 150° up, rapidly above 230°. As low as 180°, almost a third of the aldehyde formed is decomposed, and at 330° its destruction is complete.

²⁸ SABATIER and MAILHE, Ann. Chim. Phys. (8), 10, 550, 554, 557 and 568 (1907).

³⁹ Goldsmith, English patent, 17,573 of 1906; J. S. C. I., 26, 777 (1907). — Alot and Brustier, Bull. Soc. Chim. (4), 9, 733 (1911).

⁴⁴ BOUVEAULT and LOCQUIN, Bull. Soc. Chim. (3), 35, 650 (1906).

⁴¹ BOUVEAULT, Loc. cit.

^{*} SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 469 (1905).

The results are similar with propyl alcohol, with which 75% of the aldehyde is decomposed at 260°; and with n.butyl alcohol with which 92% of the aldehyde is decomposed; and for isobutyl alcohol. With ordinary isoamyl alcohol, the destruction of the aldehyde already reaches one half at 210°.

Heptyl alcohol, submitted to the action of nickel at 220°, gives only a small amount of the aldehyde, the chief product being hexane resulting from its decomposition with separation of carbon monoxide.

665. In contact with nickel, isopropyl alcohol is slowly decomposed into acetone and hydrogen from 150° up. The reaction is rapid at 210° but about 12% of the alcohol that is transformed is split into water, ethane and methane.

Secondary butyl alcohol is transformed quite regularly above 200° but 20% of the product is already decomposed, while at 310°, 80% is destroyed.

For methyl-hexyl-carbinol the decomposition is clean at 250° but at that temperature already the methyl-hexyl-ketone formed is mostly broken down into carbon monoxide, methane and hexane, only a third surviving.

666. Cobalt. The action of reduced cobalt on primary and secondary alcohols is between that of nickel and that of copper.

667. Iron. The action of iron is analogous to that of cobalt. At high temperatures, 600 to 700°, it causes a rapid destruction. An iron tube either empty or filled with iron turnings decomposes ethyl alcohol strongly at 700° giving 30% aldehyde and depositing about 7% of carbon. 45

668. Platinum. Platinum sponge acts on alcohols as does nickel but its action does not begin till above 250°. Besides the destruction of the *aldehydes* is inseparable from their formation and always predominates.

Around 250° methyl alcohol is split cleanly into hydrogen and carbon monoxide with no methane and only traces of formaldehyde.

Ethyl alcohol is attacked at 270°, and at 370° the reaction is rapid, but 75% of the aldehyde is decomposed into carbon monoxide and methane.

Propyl alcohol is split above 280°, but at 310° the aldehyde is almost completely decomposed into ethane and carbon monoxide.

The results are better with secondary alcohols since the ketones are more stable than the aldehydes.

45 IPATIEF, Berichte, 35, 1047 (1902).

BOESEKEN and VAN SENDEN, Rec. Trav. Chim. Pays-Bas, 32, 23 (1913).

⁴⁴ SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 473 (1905).

Isopropyl alcohol is transformed into acetone at 320° without notable complications and at 400° the destruction of the acetone reaches barely 3% of the product.⁴⁶

669. Palladium. The considerable affinity that this metal has for hydrogen seems to fit it for the dehydrogenation of alcohols. Benzhydrol is rapidly decomposed into benzophenone by contact with palladium sponge.

670. Zinc. Around 650° this metal decomposes alcohols strongly: sthyl alcohol yields 60% aldehyde and the gases, ethylene, carbon monoxide and methane. Isobutyl alcohol gives 75% of aldehyde and gas which is largely butylene.

Brass, an alloy of copper and zinc, acts at 600° like zinc.48

Use of Other Materials

- 671. The use of other substances to dehydrogenate alcohols is not advantageous since they act much less energetically than the metals and because they require the use of higher temperatures at which the aldehydes are decomposed into carbon monoxide and saturated hydrocarbons.
- 672. Manganous Oxide. Its action hardly begins below 320°. At 360° it decomposes *methyl alcohol* only one sixth as rapidly as compact red-orange copper; the greater part of the formaldehyde survives and the hydrogen is nearly pure.

At 360° the decomposition of *ethyl alcohol* is only one fortieth as rapid as with light copper and a part of the aldehyde is already decomposed into ethane, carbon monoxide and even carbon dioxide, the latter being formed from the carbon monoxide with a corresponding deposit of carbon, the reaction being similar to that produced by metals (614).

Propyl, isoamyl and benzyl alcohols give analogous results.49

673. Stannous Oxide. This acts above 300° as a dehydrogenation catalyst after the manner of the metals, but is slowly reduced meanwhile into metallic tin, which is easy to see in the oxide. This finely divided tin seems to possess a catalytic power similar to that of the oxide so that the mixture of metal and oxide continues to split alcohols into aldehydes and hydrogen for a long time, but as the reaction temperature is above 220°, the melting point of tin, the tiny globules

⁴⁴ SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 473 (1905).

⁴⁷ Knoevenagel and Heckel, Berichte, 36, 2816 (1903).

⁴⁴ IPATIEF, Berichte, 34, 3579 (1901) and 37, 2961 and 2986 (1904).

⁴⁹ SABATIER and MAILHE, Ann. Chim. Phys. (8), 20, 313 (1910).

of metal resulting from the reduction of the oxide gradually coalesce into larger, and consequently the activity diminishes.

Thus with *ethyl alcohol*, the brownish orange stannous oxide (resulting from the reduction of stannic oxide by the alcohol vapors) commences to act at 260°. At 350° the velocity of the reaction is almost half as great as with the same volume of very light reduced copper. The disengaged hydrogen is almost pure, the acetaldehyde being only slightly decomposed. At the end of four hours the velocity of the reaction is reduced by half.

Amyl alcohol yields the aldehyde regularly at 340°.

Methyl alcohol is attacked above 260° with the production of formaldehyde. At 350° the most of this is decomposed into carbon monoxide and hydrogen. 50

674. Cadmium Oxide. This behaves like stannous oxide and dehydrogenates while it is reduced at the same time to the metal which possesses a catalytic activity differing little from that of the oxide. Thus with *ethyl alcohol* at 300° the reaction is about one tenth as rapid as with the same volume of very active copper and maintains itself for a long time in spite of the progressive reduction of the oxide.

Benzyl alcohol acts in exactly the same way: at 350° there is a slow reduction of the oxide and at the same time a splitting of the alcohol into benzaldehyde and hydrogen. At 380° the benzaldehyde is partially decomposed into benzene and carbon monoxide. The entire absence of the resinous hydrocarbon (714) indicates that there is no dehydration.

With *methyl alcohol*, the splitting which begins at 250° is quite rapid above 300° and produces formaldehyde which is partially decomposed into carbon monoxide and hydrogen.⁵¹

675. Other Oxides. Most non-reducible metallic oxides are mixed catalysts for alcohol, causing dehydration and dehydrogenation at the same time. For some: uranous oxide, blue oxide of molybdenum, vanadous oxide, V_2O_3 , zinc oxide, dehydrogenation predominates.

In another group: beryllium oxide, zirconium oxide, chromic oxide, Cr₂O₃ (calcined above 500°), the dehydrogenating and dehydrating powers are about equal.

For a third group: chromic oxide, Cr₂O₃ (not calcined), titanium oxide, silicon dioxide, dehydration predominates.

676. With reference to methyl alcohol the classification of the oxides is quite different since in this case dehydration can not take place except by the formation of methyl ether and the conditions are

⁵⁰ SABATIER and MAILHE, Ann. Chim. Phys. (8), 20, 309 (1910).

⁸¹ SABATIER and MAILHE, Ann. Chim. Phys. (8), 20, 302 (1910).

not comparable. Except alumina, which at 390° only dehydrates, and several oxides (thoria, blue oxide of tungsten, chromic oxide and alumina above 350°) which are mixed catalysts, all metallic oxides dehydrogenate methyl alcohol with the production of formaldshyde which is more or less decomposed into carbon monoxide and methane.

The following table indicates the volume of gas obtained per minute with the same volume of various catalysts employed under the same conditions.

Oxides			Volume o	of gas in cc. per minute.				
Formaldehyde :	remaining	almost enti	irely; the gas is	nearly pure hydrogen.				
BeO .			ve	ery small				
SiO_2				. 0.3				
TiO ₂ .				. 1.2				
Formaldehyde 1	Formaldehyde partially decomposed, the hydrogen contains carbon monoxide.							
PbO a.			. .	45 (beginning)				
Formaldehyde almost completely destroyed, the gas is nearly ${ m CO} + 2{ m H_2}$.								
Fe ₂ O ₃ ⁵²				106 (beginning)				
V_2O_5 .				. 140				
SnO 52.				160 (beginning)				
Light co	opper .			152				

677. The dehydrogenating power of oxides can hardly be explained except by assuming an unstable combination of the oxide and the aldehyde. 55

678. Zinc powder, which is an intimate finely divided mixture of metallic sinc and sinc oxide, usually containing a certain proportion of cadmium and cadmium oxide, acts by virtue of these various substances as a quite active dehydrogenation catalyst, particularly toward methyl alcohol, the formaldehyde being mostly decomposed into carbon monoxide and hydrogen. Long ago Jahn noted that zinc powder splits methyl alcohol into a gas containing 30% carbon monoxide and 70% hydrogen. ⁵⁴

^{*} The gas volumes given are taken after the absorption of the carbon dioxide resulting from the slow reduction of the oxide.

^{*} SABATTER and MAILHE, Ann. Chim. Phys. (8), 20, 340 to 346 (1910).

⁴⁴ Jahn, Berichte, 13, 983 (1880).

679. Carbon. Baker's coals act towards alcohols as a mixed catalyst causing dehydrogenation and dehydration simultaneously.

Ethyl alcohol undergoes a complex reaction at 375–385°, being almost completely destroyed yielding methane and carbon monoxide. With isopropyl alcohol dehydration predominates. 56

§ 4. — DEHYDROGENATION OF POLY-ALCOHOLS

680. Glycerine is the only poly-alcohol of which the dehydrogenation has been studied. When its vapors are passed at 330° over very light reduced *copper*, prepared by the reduction of cupric carbonate at a low temperature, there is a rapid evolution of gas consisting of hydrogen mixed with methane, carbon monoxide and dioxide, the proportion of the latter rising to one third of the whole.

The initial effect of the copper is dehydrogenation to glyceric aldehyde:

$$CH_2OH.CHOH.CH_2OH = H_2 + CH_2OH.CHOH.CHO.$$

As soon as this is formed it is decomposed in the same way as it is by beer yeast into ethyl alcohol and carbon dioxide: 56

$$CH_2OH.CHOH.CHO = CO_2 + CH_2.CH_2OH.$$

A part of this alcohol is found in the distillate and a part suffers dehydrogenation by the copper to acetaldshyds, CH₂.CHO, which itself splits up, more completely when the temperature is high, into methane and carbon monoxide.

Furthermore, at the temperature of the reaction a portion of the glycerine is dehydrated to acroleine, which is mostly found in the distillate with the alcohol and water but a part of which is hydrogenated by the copper to propionic aldshyde, allyl alcohol and propyl alcohol accompanied by condensation products due to the crotonization of the aldehydes. Ethyl alcohol is the chief constituent of the liquid. 57

§ 5. — DEHYDROGENATION OF AMINES

681. Primary Amines. We have seen that nickel permits us to add hydrogen to nitriles at 200° to form primary amines (426). We may expect that it will reverse this reaction at higher temperatures and take hydrogen away from a primary amine derived from a primary alcohol, to reform the nitrils:

$$R.CH_2.NH_2 = 2H_2 + R.CN.$$

⁴⁴ LEMOINE, Bull. Soc. Chim. (4), 3, 851 and 935 (1908).

⁵⁴ GRIMAUX, Bull. Soc. Chim. (2), 49, 251 (1888).

SABATIER and GAUDION, Compt. rend., 166, 1037 (1918).

This is what takes place with benzyl-amine, with amyl-amine as well as with other primary aliphatic amines derived from primary alcohols having at least five carbon atoms.⁵⁸

When the vapors of benzyl-amine alone are passed over a layer of reduced nickel maintained at 300-50°, bensonitrile, C₆H₅.CN, is formed. But at this temperature the liberated hydrogen reacts with the amine to give toluene and ammonia (496), so that the evolution of gas is a minimum. We may write the reaction:

$$3C_6H_5.CH_2.NH_2 = C_6H_5.CN + 2C_6H_5.CH_3 + 2NH_3.$$

The yield of benzonitrile is about one third.

Likewise at 300° isoamyl-amine yields isobutyl cyanide according to the reaction:

$$3(CH_2)_2CH.CH_2.CH_2.NH_2 = (CH_2)_2CH.CH_2.CN + 2C_5H_{12} + 2NH_3.$$

The isopentans produced is partially destroyed by the nickel, depositing carbon and liberating hydrogen and lower hydrocarbons.

The reaction goes poorly with amines derived from primary alcohols having less than five carbon atoms, since with these amines nickel has a strong tendency to eliminate ammonia with the formation of sthylenic hydrocarbons (631).⁵⁹

When copper is used in place of nickel between 390 and 400°, much more complex products are obtained somewhat similar to those obtained by the hydrogenation of aliphatic nitro compounds (510).

682. Secondary and Tertiary Amines. Secondary and tertiary amines derived from primary alcohols also furnish nitriles when passed over nickel at 320-50°, by the simultaneous elimination of hydrogen and ethylenic hydrocarbons. Thus from di-isoamyl-amine and tri-isoamyl-amine, isobutyl cyanide is obtained.⁶⁰

§ 6. — SYNTHESIS OF AMINES

683. When a mixture of ammonia and benzene vapor is heated to 550° without catalyst, a slight formation of aniline is observed according to the reaction: 61

$$C_6H_6 + NH_2 = H_2 + C_6H_6.NH_2.$$

SABATIER and GAUDION, Compt. rend., 165, 224 (1917).

⁵⁹ SABATIER and GAUDION, Compt. rend., 165, 310 (1917).

MAILHE and DE GODON, Compt. rend., 165, 557 (1917). — MAILHE, Ibid., 166, 996 (1918).

MEYER and TAUZEN, Berichte, 46, 3183 (1913).

With hydrogen in presence of nickel above 350°, aniline vapors regenerate a certain amount of bensene and ammonia by the reversal of the above reaction (496).

It might be hoped that the direct production of aniline from benzene vapor and ammonia would be feasible by the use of metal catalysts at 500 to 700°. It has been found that the presence of reduced nickel, iron or copper is of no advantage, as only traces of aniline are produced. Likewise only traces of toluidine are obtained from toluens. In the most favorable case working with nickeled asbestos in an iron tube, 0.11 g. aniline was obtained from 200 g. benzene.

§ 7. — CLOSING OF RINGS BY LOSS OF HYDROGEN

684. Nickel. Methyl-o.toluidine, submitted to the action of reduced nickel at 300-30° (in presence of hydrogen), loses hydrogen to form a new cycle, yielding above 6% of indol along with methane and o.toluidine: 64

$$C_{\bullet}H$$
 CH_{\bullet}
 C

Likewise dimethyl-o.toluidine, at 300°, yields 24% of N-methyl-indol along with methane, toluidine and methyl-toluidine: 45

$$C_{\bullet}H_{\bullet}$$
 CH_{\bullet}
 CH_{\bullet}

685. Aluminum Chloride. The use of anhydrous aluminum chloride at moderate temperatures, between 80 and 140°, causes the elimination of hydrogen with the formation of new cycles.

α-Dinaphthyl yields perylens: 66

Likewise at 140°, meso-benzo-dianthrone passes quantitatively into meso-naphtho-dianthrone:

- SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 415 (1905).
- ⁶⁸ WIBAUT, Berichte, 50, 541 (1917).
- 44 CARRASCO and PADOA, Lincoi, 15 (2), 699 (1906).
- ⁶⁶ CARRASCO and PADOA, Gas. Chim. Ital., 37 (2), 49 (1907).
- 64 SCHOLL, SEER, and WELTZENBOCK, Berichte, 43, 2203 (1910).
- er Scholl and Mansfield, Berichte, 43, 1737 (1910).

At 140° phenul-a-naphthul-ketone gives a good yield of benzanthrone:



This is a typical example of many analogous reactions that can be readily carried out by this process.⁶⁸

686. Metallic Oxides. Various anhydrous metallic oxides, alumina, ferric oxide, chromium trioxide, thoria, and titania can cause the condensation of acetylene with various molecules with the elimination of hydrogen and the formation of cyclic compounds.

With ammonia pyrrol, picoline and collidines are formed, there being no hydrogen evolved in the formation of the latter:

$$\begin{aligned} &2C_{2}H_{3}+NH_{3}=H_{3}+\underbrace{C_{4}H_{4}N}_{pyrrol}\\ &3C_{2}H_{3}+NH_{3}=H_{2}+\underbrace{C_{4}H_{7}N}_{picoline}\\ &4C_{3}H_{3}+NH_{3}=\underbrace{C_{2}H_{11}N}_{collidine} \end{aligned}$$

Ferric oxide is the best catalyst for forming pyrrol.

When ethylene is used instead of acetylene, the same products are formed but at higher temperatures and with the evolution of much hydrogen.

Hydrogen sulphide gives thiophene:

$$2C_{2}H_{2} + H_{2}S = H_{2} + C_{4}H_{2}S$$

At 400-425°, water vapor forms furfural: 69

$$2C_2H_2 + H_2O = H_2 + C_4H_2O$$
.

48 SCHOLL and SEER, Sits. Akad. Wien, 120, 11, B, 925 (1911). — Annalen, 394, 111 (1912).

•• CHICHIBABINE, J. Russian Phys. Chem. Soc., 47, 703 (1915), C. A., 9, 2512 1915).

CHAPTER XV

DEHYDRATION

687. There are a large number of organic reactions which take place with the elimination of water. Many of them can be started or accelerated by the presence of so-called dehydration catalysts. As might be anticipated from the great variety of reactions of this kind, dehydration catalysts comprise many substances of very different natures, elements (phosphorus, carbon, and finely divided metals), strong mineral acids (sulphuric, hydrochloric, phosphoric, etc.), either concentrated or dilute, anhydrides of acids (phosphoric and boric), anhydrous chlorides (of aluminum, zinc and iron), various inorganic salts (ammonium salts, potassium bisulphate, calcium and aluminum sulphates, phosphates, etc.), organic acids (acetic), as well as their salts (potassium and sodium acetates).

We can distinguish two distinct modes of dehydration according to whether it takes place in the gas phase by the action of solid catalysts on the vapors which are to give up the water or in the liquid medium. We will study the two separately.

§ 1. — DEHYDRATION OF ALCOHOLS ALONE

688. Primary alcohols can undergo dehydration in two different ways: to produce an ether or a hydrocarbon, usually unsaturated. Thus with ordinary alcohol, we have:

$$2CH_3.CH_2OH = H_2O + \frac{(CH_3.CH_2)_2O}{\text{ethyl ether}}$$

and

or

$$CH_3.CH_2OH = H_2O + \underbrace{CH_2:CH_2.}_{\text{ethylene}}$$

Benzyl alcohol gives:

$$2C_{\theta}H_{5}.CH_{2}OH = H_{2}O + \underbrace{(C_{\theta}H_{5}.CH_{2})_{2}O}_{\text{bennyl ether}}$$

$$nC_{\theta}H_{5}CH_{2}OH = nH_{2}O + \underbrace{(C_{\theta}H_{5}.CH)_{n}}_{\text{resinous hydrocarbon}}$$

Methyl alcohol is an exception, as it can be dehydrated regularly in only one way, that is to form methyl ether, (CH₃)₂O.

Secondary alcohols, the dehydration of which is easier, yield ethers in exceptional cases only (e.g. benzhydrol), usually producing hydrocarbons.

The ethers can seldom be obtained from tertiary alcohols, as these are dehydrated to the hydrocarbons with still greater ease.

689. These dehydrations can be accomplished by a multitude of substances, that have affinity for water, used in excess compared with the alcohol that is to be dehydrated. But if the hydrates formed are unstable at the temperature of the operation, water is given off, regenerating the original substance which can repeat the reaction with a fresh quantity of alcohol.

This is what takes place with zinc chloride and with concentrated sulphuric acid of which a small quantity when heated can dehydrate a large amount of alcohol.

We have explained above (159) the mechanism of the action of sulphuric acid which produces either ethyl ether or ethylene from alcohol according to temperature conditions. It can continue its catalytic rôle for a long time, but is gradually diminished by being reduced to sulphur dioxide, since it slowly oxidizes the alcohol with the production of carbon dioxide and of tarry matters.

Syrupy phosphoric acid can produce an entirely analogous effect, and, as it is less readily reduced than sulphuric acid, can maintain its catalytic activity for a much longer time.^{1, 2}

Formation of Ethers

690. The formation of ethers by the direct dehydration of alcohols is possible in only a small number of cases and only with primary alcohols.

In the case of *methyl alcohol* this is the only possible manner of *dehydration* and a considerable number of catalysts can decompose its vapors into *methyl ether* and water, but they are very much less numerous than the substances which can dehydrate other alcohols to hydrocarbons.

691. Formation in Liquid Medium. Concentrated sulphuric acid is usually employed to dehydrate methyl alcohol to methyl ether.

Zinc chloride is not suitable for this reaction as it gives gaseous products by a complicated reaction and even produces a certain amount of hexamethyl-benzene.

Ethyl ether is practically prepared by the action of sulphuric acid at 140°. A mixture of 5 parts of 90% ethyl alcohol and 9 parts of

- ¹ SABATIER and MAILHE, Bull. Soc. Chim. (4), 1, 524, (1907).
- ² This is used for preparing ethylene on a commercial scale. E. E. R.
- ³ Dumas and Philigor, Ann. Chim. Phys. (2), 58, 19 (1835).
- ⁴ LE BEL and GREENE, Compt. rend., 87, 260 (1878). Jahresber. Chem., 1878, 388.

concentrated sulphuric acid is used. This mixture boils at about 140°. When it is heated to 140°, ether distils over and alcohol is added at such a rate that the boiling point does not rise. A large amount of alcohol can be transformed into ether in this way. The volume of the ether may be more than 167 times that of the sulphuric acid used. Theoretically the formation should continue indefinitely, but the yield decreases after a certain time on account of the production of a carbonaceous residue which may amount to 5% of the acid and the formation of which corresponds to the evolution of a considerable amount of sulphur dioxide.

The best yield of ether is obtained between 140 and 145°, as above that temperature more and more ethylene is formed.

Phosphoric or arsenic acid may replace the sulphuric acid in this preparation. Anhydrous zinc chloride also may be used.

Concentrated sulphuric acid at 135° produces propyl ether from the alcohol but the yield is poor because much propylene is formed.

The higher alcohols such as isobutyl do not yield ethers with concentrated sulphuric acid but only the ethylenic hydrocarbons. Nevertheless, isoamyl ether can be thus obtained (696).

Sometimes sulphuric acid at 140° enables us to obtain mixed ethers by operating on a mixture of the two alcohols. This is the case with methyl and ethyl alcohols which yield the mixed methyl-ethyl ether along with the two simple ethers. In the same way ethyl-propyl ether may be obtained, but ethyl-isobutyl can not be. Ethyl-isoamyl ether, which several chemists have failed to obtain, 1° can be prepared along with the two simple ethers by the action of 85% sulphuric acid at 135-140°.11

The mixed ethyl-tertiary-butyl ether can be obtained by heating 50 volumes of a mixture of two molecules of ethyl alcohol and one of trimethyl-carbinol with one volume of sulphuric acid in a sealed tube at 100° for 5 hours. 12

692. Although it is a secondary alcohol, benzhydrol, C₆H₅.CH(OH).-C₆H₅, is readily transformed into its ether: it is sufficient to heat it to 180° with 27% sulphuric acid.¹²

- ⁵ Evans and Sutton, Jour. Amer. Chem. Soc., **35**, 794 (1913).
- NORTON and PRESCOTT, Amer. Chem. Jour., 6, 243 (1884).
- ⁷ Boulay, Gilbert's Annalen, 44, 270 (1913).
- ⁸ Masson, Annalen, 31, 63, (1839).
- NORTON and PRESCOTT, Amer. Chem. Jour., 6, 244 (1884).
- ¹⁰ Guthrie, Annalen, 105, 37 (1858). NORTON and PRESCOTT, Amer. Chem. Jour., 6, 246 (1884).
 - ¹¹ PETER, Berichte, 32, 1419 (1899).
 - ¹² Mamontoff, J. Russian Phys. C em. Soc., 29, 234 (1897), C., 1897 (2), 408.
 - ¹² ZAGUMENNI, J. Russian Phys. Chm. Soc., 12, 431 (1880), C., 1880, 629.

The ether may be obtained also by heating benshydrol to 210-220° with finely divided copper.¹⁴

693. Formation in Gaseous Phase. Among anhydrous metallic oxides, only alumina precipitated and dried at a low temperature effects the transformation of methyl alcohol into methyl ether exclusively. The reaction commences at about 250° and is rapid at 300°, yielding methyl ether which can be completely absorbed by concentrated sulphuric acid. At about 350° the dehydration is accompanied by a slight dehydrogenation, the aldehyde produced being immediately decomposed into carbon monoxide and hydrogen.

Thoria, blue oxide of tungsten and chromium sesquioxide can dehydrate methyl alcohol to the ether above 230° but there is simultaneous dehydrogenation to the aldehydre and its decomposition products. The latter reaction is still more important with titania and takes place almost exclusively with other catalytic oxides, such as the oxides of zirconium, molybdenum, and vanadium. 15

694. Alone among the oxides, alumina at 240° enables us to obtain sthyl ether from ethyl alcohol. A little sthylens is evolved. A 90% alcohol may be used.

With propyl alcohol at 250° it gives a little propyl ether but forms propylens chiefly. It can not produce the other ethers. In the apparatus of Ipatief, under high pressures, alumina can transform ethyl alcohol into the ether, but the formation is limited by the reverse reaction. At higher temperatures only ethylene is produced. If

Ethyl ether is totally decomposed into water and ethylene by alumina at 380°. 18

Dehydration to Hydrocarbons

695. The dehydration of a single molecule to give a hydrocarbon with an ethylene double bond is the normal reaction of alcohols and also of *ethers*.

Reaction in Liquid Medium. This may readily be accomplished by concentrated non-volatile *mineral acids* and also by anhydrous zinc chlorids.

- 696. Concentrated Mineral Acids. A small proportion of concentrated sulphuric acid used at a temperature high enough to eliminate the water produced serves to prepare advantageously the lower ethylenic hydrocarbons which are gases, ethylens, propylens, and butylens.
 - 14 Knoevenagel and Heckel, Berichte, 36, 2823 (1903).
 - ¹³ SABATIER and MAILHE, Ann. Chim. Phys. (8), 20, 345 (1910).
 - 16 SENDERENS, Ann. Chim. Phys. (8), 25, 449 (1912).
 - 17 IPATIEF, Berichte, 37, 2961 (1904).
 - ¹⁸ ENGELDER, J. Phys. Chem., 21, 676 (1917).

To obtain *ethylene*, a mixture of 25 parts of alcohol and 150 parts of sulphuric acid is heated to 160-70° and a mixture of alcohol and sulphuric acid is added in drops.¹⁹

The evolution of gas is facilitated by the addition of a certain amount of fine quarts sand to the mixture. According to Senderens, this acts as a true chemical catalyst. According to the same author the results are still better when 5% of anhydrous aluminum sulphats is added to the usual mixture of alcohol and sulphuric acid. With sthyl alcohol at 157° the evolution of ethylene is thus rendered three times as rapid and propylens is formed at 130° instead of 145°; isobutyl alcohol is split at 125°.20 21

From 1500 cc. isoamyl alcohol and 100 cc. sulphuric acid in a vessel provided with a reflux condenser kept at 60-90° followed by a condenser for the amylene, 250 g. of amylene (a mixture of the 3 isomers) may be prepared in 8 hours. The alcohol remaining in the flask contains 400 g. isoamyl ether.²²

Concentrated phosphoric acid may replace sulphuric acid in these dehydrations.

697. Under analogous conditions the dehydration of molecules with mixed function may be catalyzed. Thus diacetonyl alcohol, (CH₃)₂C(OH).CH₂.CO.CH₃, warmed with traces of sulphuric acid (6 drops to 290 g. of the alcohol), furnishes mesityl oxide, (CH₃)₂C: CH.CO.CH₃, with a high yield (190 g.) on distillation.²²

698. Zinc Chloride. Anhydrous fused zinc chloride is very often employed to effect the transformation of alcohols into ethylene hydrocarbons, but it is commonly used in excess, that is in amount more than sufficient to fix as a stable hydrate all of the water that is eliminated. The same action can be exercised by the catalyst when the alcohol has a high boiling point as the alcohols of the cyclohexane series; a small quantity of the chloride serves to dehydrate these to cyclohexenes since the water that is extracted is eliminated by distillation along with the hydrocarbon so that the catalyst is continuously regenerated.

699. Iodine. In exceptional cases, iodine serves to effect the regu-

- 19 ERLENMEYER, Annalen, 192, 244 (1878).
- ²⁰ SENDERENS, Compt. rend., 151, 392 (1910).
- ** Following the observations of SENDERENS, the following method of preparing ethylene has been devised and has given excellent service. In a 500 c.c. flask 200 c.c. conc. sulphuric acid, 100 c.c. 95% alcohol and 25 g. of dehydrated alum are heated to 157 to 175°, the thermometer dipping in the mixture. One operator repeated this five times in an afternoon and obtained 667 g. ethylene bromide. E. E. R.
 - ²² ADAMS, KAMM, and MARVEL, Jour. Amer. Chem. Soc., 40, 1950 (1918).
 - ²⁸ KOHN, Monatch. Chem., 34, 779 (1913).

lar dehydration of compounds containing alcohol groups. Diacstonyl alcohol, (CH₃)₂C(OH).CH₂.CO.CH₃, which distillation alone breaks down partially into two molecules of acetone, is dehydrated by sulphuric acid to mesityl oxide (697). The same dehydration takes place quantitatively when it is distilled with a small amount of iodine.²⁴

700. Reaction in Gaseous System. This can be effected by a large number of solid catalysts among which the best are alumina, clay, thoria and the blue oxide of tungsten.

Elements. Animal charcoal, extracted with hydrochloric acid, is a rather mediocre catalyst for alcohols: above 350° it produces ethylene from sthyl alcohol, accompanied by a certain amount of methane, carbon monoxide and hydrogen resulting from the formation of acetal-dehyde which is mostly destroyed. Propyl alcohol, above 300°, gives a gas of which 87% is propylene, with ethylene and other gaseous products.²⁶

Red phosphorus acts more rapidly at much lower temperatures and probably owes this activity to small amounts of phosphorus and phosphoric acids preëxistent in the material and which are formed in considerable amounts in consequence of the oxidation of the phosphorus by the alcohol with a correlative production of phosphine.

With sthyl alcohol at 240°, a rapid evolution of ethylene is obtained containing 5% phosphine. Similar results are obtained with propyl alcohol. The proportion of phosphine is less with normal and iso-butyl alcohols and negligible with isopropyl alcohol which is already split at 150°.

The presence of phosphine, which is difficult to get rid of, takes away much of the interest in this case of catalysis.

701. Finely divided metals have an important catalytic dehydrogenating effect on primary and secondary alcohols (651); but they decompose tertiary alcohols rapidly at moderate temperatures into unsaturated hydrocarbons. Reduced nickel acts in this way without complications at 220 to 300° and reduced copper acts similarly above 280 to 300°.26

Passing an aliphatic alcohol over copper at 300° is a simple method of determining its class. A primary alcohol forms an aldehyde, a secondary one a ketone, while a tertiary breaks up into water and an unsaturated hydrocarbon.²⁷

702. Anhydrous Metal Oxides. Grigoreff in 1901 was the first to note the special aptitude of an oxide to dehydrate alcohols: he found

²⁴ Hibbert, Jour. Amer. Chem. Soc., 37, 1748 (1915).

²⁶ SENDERENS, Compt. rend., 144, 381 (1907).

²⁶ SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 467 and 472 (1905).

²⁷ SABATIER and SENDERENS, Bull. Soc. Chim. (3), 33, 263 (1905).

that alumina decomposes ethyl and propyl alcohols to the hydrocarbons with 90 % yields.²⁸

This property of alumina was studied by Ipatief and found also in the material of graphite crucibles, which is a mixture of graphite (inactive) and clay, while other oxides (of zinc, iron, tin, chromium, etc.) were revealed as dehydrogenating catalysts.²⁹

The catalytic activity of various oxides was made the object of a thorough study by Sabatier and Mailhe, ²⁰ who were able to demonstrate the great dehydrating power of thoria and of the blue oxide of tungsten. We have already noted (675) that the oxides that are not reducible, or only slowly reducible, by alcohols can be divided into dehydrogenating, dehydrating, and mixed catalysts which cause both reactions simultaneously.

The direction and the importance of the activity of the various oxides can be clearly shown by a comparison of the volume and composition of the gas evolved by them, when equal volumes of them are used at 340-50° with the same amount of ethyl alcohol; all of the oxides having been prepared below 350°: 31

		Volu	Composition			
	Oxide	gas in cc	. per min.	C ₂ H ₄ %	H ₂ %	
	(ThO ₂		31	100	trace	
Dehydrating	Al_2O_3		21	98.5	1.5	
	W_2O_5		57	98.5	1.5	
	Cr ₂ O ₃		4.2	91	9	
	SiO ₂		0.9	84	16	
	TiO ₂		7.0	63	37	
	BeO		1.0	45	55	
Mixed	ZrO_2		1.0	45	55	
IN CLOU	UO ₂		14	24	76	
	Mo ₂ O ₅ .		5	23	77	
	Fe ₂ O ₃ .		32	14	86	
	V ₂ O ₃		14	9	91	
	Z_{nO}		6	5	95	
Dalandara and a	MnO		3.5	0	100	
Dehydrogenating	∖MgO		traces	0	100	

²⁶ GRIGOREFF, J. Russian Phys. Chem. Soc., 33, 173 (1901).

²⁹ IPATIEF, Berichte, 34, 596 (1901); 35, 1047 (1902); 36, 1990 (1903).

^{**} SABATIER and MAILEE, Bull. Soc. Chim. (4), 1, 107, 341, 524 and 733 (1907). — Compt. rend., 146, 1376 (1908); 147, 16 and 106 (1908); 148, 1734 (1909). — Ann. Chim. Phys. (8), 20, 289 (1910).

²¹ SABATIER and MAILEE, Ann. Chim. Phys. (8), 20, 341 (1910).

703. We have noted (76) that the *physical condition* and the method of preparation of an oxide have a great influence on its activity and even on the direction of the catalysis.

These differences are very marked for the various varieties of chromium sesquioxide (78) the only one of which that is suitable for the dehydration of alcohols is that obtained by drying the precipitated blue hydrated hydroxide below 350°.

704. Titanium oxide, TiO₂, prepared by calcining at a red heat has very little activity. To obtain a suitable oxide, the hydroxide prepared by the action of ammonia on titanium chloride is dehydrated below 350°. **

705. Crystallized silica is almost without action on alcohols below 400°. The pure silica obtained by decomposing silicon fluoride by water, washing thoroughly and drying at 300°, is also only slightly active. The most active form is obtained by adding dilute acid to sodium silicate, washing and drying the gelatinous precipitate.

706. The most active form of alumina is prepared by precipitation from aluminum nitrate, washing the precipitate well and drying at 300°. Good results are also obtained with the oxide prepared by calcining pure ammonium alum at red heat. The basic aluminum sulphats obtained by calcining aluminum sulphate at a dull red is a very active catalyst. On the contrary, preparations of alumina which have been heated to redness for a long time are almost inactive and sometimes do not give an appreciable amount of gas from ethyl alcohol even at 420°.

Bauxite, aluminum hydroxide mingled with silica and ferric hydroxide, has low catalytic power and dehydrogenates chiefly at about 400°.34

The nature of the reaction catalyzed is closely connected with the condition of the oxide and bears a certain relation to its ease of solution in acids.²⁵

707. Thoria, on the contrary, does not present these difficulties and its catalytic activity is not sensibly diminished by calcination at

²⁵ SABATIER and MAILHE, Ann. Chim. Phys. (8), 20, 325 (1910).

⁴⁸ Sabatier and Mailer, Ann. Chim. Phys. (8), 20, 300 (1910).

Samples of bauxite from different sources differ widely in catalytic power. With isoamyl alcohol a sample of German bauxite gave gaseous products but poor yields of amylene, while a Tennessee sample gave little gas and an excellent yield of amylene. The bauxite was used in a copper tube 35 x 900 mm. at about 400°, the alcohol being admitted at about 100 drops per minute. Several pounds of amylene were thus prepared. — C. H. Milligan.

³⁵ IPATIEF, Berichte, 37, 2986 (1904).

red heat: it seems that its high molecular weight may be in the way of molecular condensations such as alumina appears to undergo when heated to redness.³⁶

708. There are great differences in the duration of the catalytic activity of various oxides; usually it goes on decreasing because the surface of the oxide is gradually covered by small amounts of tarry or carbonaceous matter which hinder gaseous exchanges and also because molecular condensations take place in the oxides, without doubt, even when the temperature of the reaction is below 400°. If we consider only the three good dehydration catalysts, alumina, thoria, and the blue oxide of tungsten, alumina, the lightest molecule (Al₂O₃, molecular weight 92) is the one which weakens most rapidly. An active specimen which disengaged 14 cc. ethylene per minute at 340°, gave only 7 cc. after three hours of use.

However certain observers have found no weakening after five hours.²⁸ ²⁹

The blue oxide of tungsten is much more permanent: the evolution of gas may continue for several hours without noticeable weakening. The same is true of thoria which has the great additional advantage of being very readily regenerated when long usage has gummed it up; calcining at a red heat for a few instants is sufficient to render it perfectly white and restore its original activity.⁴⁰

709. For a given catalyst, elevating the temperature greatly accelerates the reaction. By operating under the same conditions with ethyl alcohol and the blue oxide of tungsten, it has been found that the evolution of ethylene begins at about 250° and becomes more and more rapid as the temperature rises. The yield per minute was:

- 37 SABATIER and MAILHE, Ann. Chim. Phys. (8), 20, 299 (1910).
- ²⁸ Engelder, J. Phys. Chem., 21, 676 (1917).
- ** I have used the same alumina catalyst for many days in making ethylene without noticing any deterioration. E. E. R.
- ⁴⁰ A thoria catalyst may be cleaned by passing steam over it at 380° till all volatile material is removed and following this with nitrogen peroxide at the same temperature as long as there is any action, the oxides of nitrogen being finally displaced by steam. A catalyst so regenerated is snow white and shows its original activity. Kramer and Reid, J. Amer. Chem. Soc. 43, 884 (1921).

³⁶ In the catalytic preparation of mercaptans, Kramer and Reid (J. Amer. Chem. Soc. 43, 882 (1921)) find that the activity of a thoria catalyst depends somewhat on the temperature to which it has been subjected, being considerably diminished by heating much above 400°. Some preparations of thoria such as Welsbach gas mantles and the extremely voluminous product obtained by dropping thorium nitrate into a red hot crucible are absolutely inactive so far as this reaction is concerned.— E. E. R.

Temperatur	е										C	.c.	per minute
260°													5
300°													17.5
310°										٠.			27
. 330°													48.5
3 40°													57.5
370°													73

But it must be remembered that for any given oxide, the elevation of the temperature tends to introduce and make more and more prominent the reaction of dehydrogenation. Thus at 340° titania produces practically pure ethylene from alcohol, but at 340° with a more rapid evolution of gas there is some hydrogen, while at 360° the hydrogen amounts to one third of the gas.⁴¹

Above 400° the gas produced may contain ethans along with the hydrogen. 42

- 710. The presence of water in the alcohol is unfavorable to dehydration but does not interfere with dehydrogenation. Thus with alcohol diluted with its own volume of water, alumina gives a gas containing twice as much hydrogen as with absolute alcohol.
- 711. Increase of *pressure* retards the dehydration of alcohols, or rather raises the temperature at which this takes place; the intermediate production of the ether from primary alcohols is favored by increase of pressure which is unfavorable to the separation of the hydrocarbon.⁴³
- 712. The dehydration of alcohols higher than propyl, effected by oxides or by other catalysts, usually leads to the production of several isomeric unsaturated hydrocarbons and frequently also to the formation of a certain proportion of polymers (211).
- 713. Alumina. The best results are obtained with alumina precipitated from aluminum nitrate by ammonia, well washed and dried at 300°.

The dehydration of *methyl* alcohol begins at about 250° and is rapid below 300°, yielding exclusively methyl ether absorbable by concentrated sulphuric acid. At about 350°, the ether is accompanied by a small amount of aldehyde, a little of which is condensed, and hydrogen is collected containing carbon monoxide resulting from the partial decomposition of the formaldehyde.

With ethyl alcohol, ether is formed above 240° and at 290° pure ethylene is evolved regularly, this evolution becoming rapid at 340°.

⁴ SABATIER and MAILHE, Ann. Chim. Phys. (8), 20, 325 (1910).

⁴⁴ Engelder, J. Phys. Chem., 21, 676 (1917).

⁴⁸ IPATIEF, J. Russian Phys. Chem. Soc., 36, 786 and 813 (1904), C., 1904 (2), 1020 and 38, 63 and 92 (1906), C., 1906 (2), 86 and 87.

It is not necessary to go beyond 360° where the ethylene begins to be decomposed and where its evolution slows down rather rapidly on account of the weakening of the catalyst.⁴⁴

Propyl alcohol gives a regular current of propylene above 300° without any of the ether.

Normal and iso-butyl alcohols likewise give a regular evolution of hydrocarbons entirely absorbable by sulphuric acid. Both yield mixtures of the isomeric hydrocarbons, C₄H₃.⁴⁵ However, Ipatief obtained pure isobutylene from isobutyl alcohol.⁴⁶

With isoamyl alcohol, the dehydration goes readily, the best yield being obtained between 500 and 540°. The product contains several isomeric hydrocarbons, C₆H₁₀, but the proportion of isopropyl-sthylens is greater than in the dehydration by sulphuric acid.⁴⁷

At 450°, secondary butyl alcohol gives pure butylene and tertiary butyl alcohol, or trimethyl-carbinol, yields only isobutylene. 48

At a dull red, allyl alcohol evolves quite pure propylens with a correlative production of acrolsins. 49

714. Benzyl alcohol is readily dehydrated at above 300° to form the yellowish resinous hydrocarbon $(C_7H_6)_x$, without evolution of gas.⁵⁰

Other primary, secondary, or tertiary aromatic alcohols are readily dehydrated by alumina without complications below 350°, with the production of the corresponding unsaturated hydrocarbons. Thus phenyl-benzyl-carbinol, C₆H₅. CH (OH). CH₂. C₆H₅, yields stilbens, C₆H₅.-CH: CH. C₆H₅, quantitatively.⁵¹

Borneol gives menthens and the various secondary or tertiary cyclohexyl alcohols are readily changed to the corresponding cyclohexens hydrocarbons. Thus cyclohexanol is entirely transformed into cyclohexens and 1.2-dimethyl-cyclohexanol yields 1.2-dimethyl-cyclohexens. 52

At 350° and 30 to 40 atmospheres with alumina, decahydronaphthol yields octahydronaphthalens, C₁₀H₁₆, boiling at 197°. 55

715. Blue Oxide of Tungsten. Tungstic oxide is readily reduced by alcohol vapors above 250° and brought to the blue oxide, intermediate between WO₂ and WO₂, approaching the composition W₂O₅

- 44 SPRENT, J. Soc. Chem. Ind., 32, 171 (1913).
- 4 SENDERENS, Bull. Soc. Chim. (4), 1, 692 (1907).
- 46 IPATIEF and SDZITOWECKY, Berichte, 40, 1827 (1907).
- ⁴⁷ Adams, Kamm and Marvel, J. Amer. Chem. Soc., 40, 1950 (1918).
- 44 IPATIEF and SDZITOWECKY, Berichte, 40, 1827 (1907).
- 49 Krestinsky and Nikitine, J. Russian Phys. Chem. Soc., 44, 471 (1912).
- * SABATIER and MAILHE, Ann. Chim. Phys. (8), 20, 298 (1910).
- ⁵¹ Sabatier and Murat, Ann. Chim. (9), 4, 284 (1915).
- 12 IPATIEF and RUTALA, J. Russian Phys. Chem. Soc., 44, 1692 (1912).
- 54 IPATIEF, Berichte, 43, 3383 (1910).

more and more nearly, and which on exposure to the air, after cooling, reoxidizes spontaneously, more or less rapidly regenerating the original yellow oxide.

This blue oxide is a mediocre catalyst for *methyl* alcohol which it does not attack till 330°, dehydrating and dehydrogenating it simultaneously, but is an excellent dehydration catalyst, very active and very regular, for other alcohols.⁵⁴ By using a train of blue tungsten oxide 51 cm. long at 340° and vaporizing 17 g. of alcohol per hour a regular evolution of 101 cc. ethylene per minute containing only 1 or 2% of hydrogen, was obtained, 5.1 g. of the alcohol escaping decomposition. By doubling the rate of flow of the alcohol the evolution of gas reached 140 cc. per minute.

At 320°, propyl, isobutyl and isoamyl alcohols give good yields of the unsaturated hydrocarbons, and benzyl alcohol is rapidly transformed into crusts of the yellow polymer (714).

716. Thoria. For all the alcohols, except methyl, thoria is a very regular catalyst, the properties of which have already been mentioned (708).

With *ethyl* alcohol, the reaction begins around 280° and is readily accelerated by rise of temperature. By using a boat containing 4.7 g. thoria, at 325°, 11 cc., and at 350°, 31 cc. of practically pure ethylene were obtained per minute.

The results are equally good with propyl and isobutyl alcohols and with the other alcohols mentioned under alumina.

The secondary alcohol, isopropyl, begins to yield propylene at 260°.

717. Mineral Salts. Clay, or hydrated aluminum silicate, and particularly the white variety, kaolin, has a remarkable dehydrating power with alcohols.⁵⁵

The fragments of a graphite crucible (a mixture of graphite and clay) gave Ipatief a good yield of unsaturated hydrocarbons from alcohols.⁵⁶

In 1906, Bouveault noted the special activity of clay and designed an apparatus for using it for the dehydration of various alcohols quite similar to that which he employed for their dehydrogenation over copper (654). The catalyst consisted of clay balls about 1 cc. in volume, dried at 300° in a current of air and packed in the 1 m. vertical tube of the apparatus in which about 1 k. of alcohol per day could be dehydrated. Ethyl, propyl, isobutyl and cyclohexyl alcohols

⁵⁴ SABATIER and MAILEE, Ann. Chim. Phys. (8), 20, 328 (1910).

Kaolin was used as catalyst in preparation of ethylene at Edgwood Arsenal, U. S. A., during the war. — E. E. R.

⁵⁴ IPATIMF, Berichte, 36, 1990 (1903).

are readily dehydrated by this means. In the case of isoamyl alcohol the hydrocarbons are isomerized as with alumina or zinc chloride.⁵⁷

All aluminum salts have more or less of the catalytic power of alumina. The basic aluminum sulphates obtained by calcining neutral aluminum sulphate at a dull red ⁵⁸ and likewise the mixtures of these with alkali sulphates obtained by calcining potassium and sodium alums have this power.

- 718. Calcium sulphate is a mediocre catalyst. When obtained by calcining gypsum at a moderate temperature, it gives with alcohol at 420° an evolution of ethylene containing 6% of hydrogen, while if it is prepared at a red heat, it gives a very slow evolution of hydrogen containing 14% of ethylene at 460°.
- 719. Aluminum phosphate is recommended as a good catalyst by Senderens, who explains this aptitude as a sort of culmination of the catalytic power of alumina and that of phosphorus. Ethyl alcohol is decomposed above 330° and rapidly at 380°. With propyl, dehydration commences at 300° and is rapid at 340°; with butyl, the reaction is important at 320°. Isoamyl alcohol is attacked at above 300°, while 250° is high enough to decompose isopropyl, which goes rapidly at 300°. The decomposition of trimsthyl-carbinol begins at 140°.
- 720. The Case of Benzhydrol. We have noted above (692), that benzhydrol, C_6H_6 , $CH(OH) \cdot C_6H_6$, heated to 210° with copper powder gives the ether, $((C_6H_6)_2CH)_2O$, in 75% yield instead of benzophenons. At a higher temperature, 290°, copper powder produces benzophenone chiefly with a slow evolution of hydrogen, along with a little of the ether and of diphenylmethans.

In fact the alcohol is dehydrogenated to benzophenone but the liberated hydrogen is used for the most part immediately to form diphenylmethane and particularly symmetrical tetraphenyl-ethane:

```
\begin{array}{l} C_6H_5.\,\mathrm{CH}\,(\mathrm{OH})\,.\,C_6H_5 = H_2 + C_6H_5.\,\mathrm{CO}\,.\,C_6H_5 \\ C_6H_5.\,\mathrm{CO}\,.\,C_6H_5 + 2H_2 = H_2\mathrm{O} + C_6H_5.\,\mathrm{CH}_2\,.\,C_6H_5 \\ 2C_6H_5.\,\mathrm{CO}\,.\,C_6H_5 + 3H_2 = 2H_2\mathrm{O} + (C_6H_5)_2\mathrm{CH}\,.\,\mathrm{CH}\,(C_6H_6)_2 \end{array}
```

Dehydrating catalysts lead to the same result as copper. The vapors of benzhydrol passed over thoria at 420° give, without elimination of hydrogen, a mixture of benzophenone, diphenylmethane and sym.tetraphenyl-ethane with the separation of water simply.

- ⁵⁷ BOUVEAULT, Bull. Soc. Chim. (4), 3, 117 (1908).
- 58 SABATIER and MAILHE, Ann. Chim. Phys. (8), 20, 300 (1910).
- 50 SENDERENS, Bull. Soc. Chim. (4), 3, 633 (1908).
- ⁴⁰ SENDERENS, Bull. Soc. Chim. (4), 1, 690 (1907).
- ⁶¹ SENDERENS, Compt. rend., 144, 1109 (1907).
- ⁶² Knoevenagel and Heckel, Berichte, 36, 2816 (1903).
- ⁴⁸ Sabatier and Murat, Ann. Chim. (9), 4, 282 (1915).

721. Catalytic Passage from an Alcohol to the Corresponding Hydrocarbon. This passage is realized easily in two successive steps: 1st dehydration of the alcohol over alumina or thoria to the unsaturated hydrocarbon; 2nd hydrogenation of this hydrocarbon over a slightly active nickel at 200-50°:

$$\frac{C_{n}H_{2n+1}OH}{\frac{alcohol}{}} = C_{n}H_{2n} + H_{2}O$$

$$C_{n}H_{2n} + H_{2} = C_{n}H_{2n+2}$$

A large number of syntheses of hydrocarbons in this way have been reported by Sabatier and Murat; for example, uns.diphonyl-ethane (C₆H₅)₂CH.CH₃ was prepared from methyl-diphenyl-carbinol, (C₆H₅)₂-C (OH).CH₃.⁶⁴

722. The two reactions can be superimposed by submitting the alcohols to the simultaneous action of alumina and nickel, but a necessary condition is that the two reactions can be carried on at the same temperature which is usually impracticable at the ordinary pressure. They can be readily carried on simultaneously in the apparatus of Ipatief (585). Thus fenchyl alcohol (40 g.) with alumina (1.5 g.) and nickel oxide (2.5 g.) with hydrogen at 110 atmospheres for 40 hours at 230° gave a good yield of fenchane, boiling at 162°, and carromenthol gave menthane.

Camphor, when treated under the same conditions at 220°, is changed into isocamphene, melting at 63°. The succession is doubtless: 65

$$\begin{array}{cccc} \underline{C_{10}H_{16}O} & \longrightarrow & \underline{C_{10}H_{18}O} & \longrightarrow & \underline{C_{10}H_{16}} & \longrightarrow & \underline{C_{10}H_{18}.} \\ \underline{campbare} & \underline{borneol} & \underline{campbare} & \underline{campbare} & \underline{campbare} \end{array}$$

Catalytic Dehydration of Poly-alcohols

723. It is seldom that the dehydration of poly-alcohols leads to hydrocarbons; aldehydes and ketones are commonly formed.

However, it has been found that when the vapors of 2-methyl-butane-diol(1.3) are passed over kaolin at above 400°, isoprene is formed:66

$$HOCH_2.CH(CH_4).CH(OH).CH_4 = 2H_2O + CH_2:C(CH_4).CH:CH_2.$$

Quinite, C₆H₁₀(OH)₂, submitted to the action of alumina at 350° and 30 to 40 atmospheres pressure, is dehydrated to dihydro-benzene,

⁴⁴ SABATIER and MURAT, Ann. Chim. (9), 4, 254 (1915).

^{45, 3205 (1912).}

⁶⁶ KYRIAKIDES and EARLE, U. S. Patents, 1,094,222, 1,094,223 and 1,106,290.

C₆H₆, along with some tetrahydro-phenol, C₆H₆.OH, resulting from the incomplete dehydration.⁶⁷

724. Glycol, HOCH₂.CH₂OH, heated at 400° with alumina yields chiefly acetaldshyde which condenses partially to parallshyde.

Pinacons, (CH₂)₂C(OH).C(OH)(CH₂)₂, is changed at 300-20° into pinacolins as it is by the action of dilute sulphuric acid.⁶⁸

725. Glycerine in the liquid form to which are added small amounts of alumina, aluminum sulphate or potassium bisulphate, is dehydrated to acroleine at about 110°:

$$HOCH_2.CH(OH).CH_2OH = 2H_2O + CH_2:CH.CHO.$$

To 100 parts of glycerine, 4 parts anhydrous aluminum sulphate, 8 parts of the hydrated, or 5 of potassium bisulphate are used. The yield is 17 to 19%, or a little smaller than when 227 parts of bisulphate are used as in the ordinary method.⁶⁹

This process has the inconvenience that acetaldehyde and sulphur dioxide are evolved; the same is true when these catalysts are replaced by ferric or cupric sulphates.

Better results are obtained with anhydrous magnesium sulphats, with which more than 50% of the theoretical yield is obtained at 330–40°, with negligible amounts of by-products, while at 360° acetaldehyde appears.⁷⁰

726. Dehydration in the Gaseous Phase. When the vapors of glycerine are passed over alumina at about 360°, complete dehydration to acrolsīns takes place, but a portion of this is decomposed into sthylens and carbon monoxids while another portion is crotonized to higher aldehydes which condense along with the water and acroleIne.⁷¹

When for the alumina catalyst is substituted black uranous oxide, which dehydrates and dehydrogenates alcohols at the same time, with a predominance of the latter reaction (675), results intermediate between those with alumina and those with copper (680) are obtained.

By using kaolin at 380-400° or aluminum phosphate at 450° we can transform butane-diol(1.3) into butadiene regularly or pentane-diol-(2.4) into piperylene. The presence of a little hydrobromic acid or of aniline hydrobromide increases the yield which for piperylene reaches 50%.

⁶⁷ IPATIEF, Berichte, 43, 3383 (1901). — J. Russian Phys. Chem. Soc., 42, 1552 (1911).

⁶⁸ IPATIEF, J. Russian Phys. Chem. Soc., 38, 92 (1906).

^{**} SENDERENS, Bull. Soc. Chim. (4), 3, 828 (1908). — Compt. rend., 151, 530 (1910).

⁷⁶ Wohl and Mylo, Berichte, 45, 2046 (1912). — WITEEMANN, J. Amer. Chem. Soc., 36, 1766 (1914).

⁷¹ SABATIER and GAUDION, Compt. rend. 166, 1034 (1918).

Pinacons is likewise dehydrated to *dimsthyl-butadiens* when its vapors are passed over copper at 430-500° and the yield is raised to 70% by the presence of a little hydrobromic acid.⁷²

727. Ring Formation by the Dehydration of Poly-alcohols. Long chain molecules containing several alcohol groups can pass into the furfurans ring by catalytic dehydration in solution.

Arabinose, HOCH₂. CH (OH). CH (OH). CH (OH). CHO, when boiled with sulphuric acid diluted to one third, is converted into furfural,²

Mucic acid or saccharic acid, HOOC. (CHOH)₄. COOH, heated to 100° with hydrochloric acid, loses two molecules of water to form dehydro-mucic or furfurane-dicarbonic acid: 74

- ⁷³ Kyriakides, J. Amer. Chem. Soc., 36, 980 (1914).
- ⁷⁸ STONE and TOLLENS, Annalen, 249, 237 (1888).
- ⁷⁴ YODER and TOLLENS, Berichte, 34, 3446 (1901).

CHAPTER XVI

DEHYDRATION (Continued)

§ 2. — ELIMINATION OF WATER BETWEEN AN ALCOHOL AND A HYDROCARBON

728. The use of anhydrous aluminum chlorids enables us to condense an aromatic alcohol with an aromatic hydrocarbon in the liquid phase. Thus benzyl alcohol, C₆H₅.CH₂OH, and benzens give diphenylmethans, C₆H₅.CH₂.C₆H₅, accompanied by a certain amount of ortho and para dibenzyl-benzenes and other hydrocarbons among which is found anthracens.¹ The same reaction takes place with secondary aromatic alcohols which yield tertiary hydrocarbons. With benzene we have:

$$C_6H_6.CH(OH).R + C_6H_6 = H_2O + C_6H_6.CH$$
 C_4H_6

The yield is better when R is an aromatic residue than when it is *methyl* or specially *ethyl*. The use of an excessive quantity of aluminum chloride, particularly if the temperature is high, may lead to the elimination of a phenyl group or of an aliphatic residue, R.²

By adding aluminum chloride to a mixture of methyl-phenyl-carbinol, C₆H₅.CH(OH).CH₅, and benzene kept at 25-35°, a 20% yield of diphenyl-ethane is obtained along with ethyl-benzene, diphenyl-methane, and anthracene, due to a further action of the chloride. By operating at 10° with 5 molecules of benzene and 0.5 of aluminum chloride a 65% yield of diphenyl-ethane is obtained.

Under the same conditions, ethyl-phenyl-carbinol forms diphenyl-propane in 40 % yield.

Benzhydrol dissolved in 5 molecules of benzene to which is added 1 molecule of aluminum chloride at 35-40°, gives a 40% yield of triphenyl-methane with some diphenyl-methane. By operating below 10°, the yield of triphenyl-methane reaches 65 to 70%.

- ¹ Huston and Friedemann, J. Amer. Chem. Soc., 38, 2527 (1916).
- ² Huston and Friedemann, J. Amer. Chem. Soc., 40, 785 (1918).
- ³ Huston and Friedemann, J. Amer. Chem. Soc., 40, 785 (1918).

§ 3. — ELIMINATION OF WATER BETWEEN AN ALCOHOL AND AMMONIA OR AMINES

Reactions in Liquid Systems

729. The primary aliphatic alcohols heated for several hours at 220° in an autoclave with aniline and a very small amount of iodine as a catalyst, give good yields of the corresponding alkyl-anilines.

Thus by heating equal molecules of aniline and methyl alcohol for 9 hours at 230° with 1% of iodine, a yield of 73% of methyl-aniline is obtained. By using 2 molecules of the methyl alcohol, 86% of dimethyl-aniline is obtained in 7 hours under the same conditions.

By heating 1 molecule of aniline and 4 molecules of ethyl alcohol with 0.5 g. iodine 10 hours, 95% of disthyl-aniline is obtained.

Under the same conditions, benzyl alcohol and aniline give benzyl- or dibenzyl-aniline and isoamyl alcohol furnishes amyl- and diamyl-anilines.

With alcohols and a little iodine, α - and β -naphthyl-amines react similarly.

730. Aromatic Alcohols may condense with anilins or its homologs when they are heated gently with dilute hydrochloric acid. Thus tetra-methyl-diamino-benzhydrol, (CH₂)₂N.C₀H₄.CH(OH).C₆H₄.N(CH₃)₂, eliminates a molecule of water with aniline to give tetramethyl-leucaniline, ((CH₃)₂N.C₆H₄)₂CH.C₆H₄.NH₂.

Reactions in Gaseous Systems

731. We have seen above that the catalytic dehydration of alcohols by various anhydrous metallic oxides has been explained by Sabatier and Mailhe on the assumption of the formation of a sort of unstable ester between the alcohol and the oxide acting as an acid, e.g. an alcohol thorinate (603).

But according to the fundamental method of Hofmann, ammonia acts on the esters of mineral acids to form amines. Sabatier and Mailhe have imagined that the unstable esters formed with the oxides should behave in the same way. It was to be hoped that, at least for some oxides, the reaction of ammonia with the temporary ester should be more rapid than the decomposition of this ester into an ethylenic hydrocarbon.⁶

⁴ Knoevenagel, J. prakt. Chem. (2), 89, 30 (1914).

⁵ Badische, German Patent, 27,032 (1883).

^{*} SABATIER and MAILRE, Compt. rend., 150, 823 (1910).

Experiment has fully verified this expectation. Thus with thoria and an aliphatic alcohol we have:

$$2C_nH_{2n+1}.OH + ThO_2 = H_2O + \frac{ThO(OC_nH_{2n+1})_2}{thorinate}$$

Then:

$$ThO(OC_nH_{2n+1})_2 + 2NH_3 = H_2O + \underbrace{HC_nH_{2n+1}.NH_3}_{amine} + \underbrace{ThO_3}_{regenerated}$$

a succession of reactions which is equivalent to the single reaction:

$$C_nH_{2n+1}.OH + NH_3 = H_2O + C_nH_{2n+1}.NH_2.$$

732. This reaction does not take place in the absence of a catalyst, but does go well in the presence of thoria at 300-50°, the dehydration into an unsaturated hydrocarbon being only a side reaction. Thus with sthyl alcohol, which is largely broken down to ethylene by thoria at 350°, the presence of ammonia almost completely prevents the evolution of the hydrocarbon but causes the production of sthyl-amine. The same is true with other dehydrating catalysts, alumina, blue oxide of tungsten and equally with the mixed catalysts, such as titania, chromic oxide, blue oxide of molybdenum, zirconia, etc. The formation of the amine directs the activity of the catalysts to its profit: the decomposition of alcohols into aldehydes and hydrogen as well as into water and ethylenic hydrocarbons is almost suppressed and the formation of the amine predominates.

Furthermore the *primary amine* thus produced reacts in its turn on the alcohol in the presence of the catalytic oxide as does ammonia, and forms the *secondary amine*:

$$C_nH_{2n+1}.OH + C_nH_{2n+1}.NH_2 = H_2O + (C_nH_{2n+1})_2NH$$

and there is the possibility of the formation of some tertiary amine by the action of the secondary on the alcohol.

733. The direct action of ammonia gas on alcohols is a general method for the preparation of amines. Into a tube containing several grams of thoria heated below 350° (from 250 to 350° according to circumstances) are passed at the same time alcohol vapors and ammonia (furnished very conveniently by a cylinder of liquid ammonia). The liquid condensed at the other end of the tube is a mixture of ammoniacal water, primary and secondary amines (with traces of tertiary) and untransformed alcohol holding in solution a certain amount of the ethylenic hydrocarbon. The latter products are easily separated from the amines by fractional distillation.

From propyl alcohol, mono- and dipropyl-amines can be readily prepared and mono- and di-isoamyl-amines from isoamyl alcohol.

⁷ Sabatier and Mailee, Compt. rend., 148, 898 (1909).

- 734. Likewise benzyl alcohol and ammonia with thoria at 300-350° give only a small amount of the resinous hydrocarbon (C₇H₆)_x, but yield chiefly benzyl- and dibenzyl-amines, and a small amount of tribenzyl-amine, which solidifies in the condenser tube. By operating at 330°, benzyl-amine is the main product, while at 370-380°, dibenzyl-amine predominates, but there is at this temperature a notable decomposition of the alcohol to the aldehyde, which, in turn, is split into benzene and carbon monoxide.⁸
- 735. The secondary alcohol, isopropyl, does not suffer appreciable dehydration over thoria at 250°, but at that temperature ammonia is effective and gives about 20% of isopropyl-amine accompanied by a little di-isopropyl-amine. Around 300° a considerable evolution of propylene is observed and the condensed liquid contains about one third isopropyl-amine and about the same amount of secondary, along with water and unchanged alcohol.

Likewise diethyl-carbinol and dipropyl-carbinol give mixtures of the corresponding primary and secondary amines.¹⁰

- 736. The method is less easy to apply to benzhydrol: yet its vapors when carried by an excess of ammonia over thoria at 280° give some benzhydryl-amine, but dehydration preponderates producing tetraphenyl-ethylene.
- 737. The secondary cyclohexans alcohols (cyclohexanol and its homologs) are dehydrated rapidly in contact with thoria at 300-350° but in the presence of ammonia at 290-320° the reaction is, for the most part, directed toward the formation of amines, hardly more than 30 to 40% of the unsaturated hydrocarbons being simultaneously produced.

In this way cyclohexyl-amins and the three methyl-cyclohexyl-aminss have been prepared, some of the secondary amines being formed in all cases.¹¹

738. Mixed Amines. In this reaction the ammonia may be replaced by a primary aliphatic amine which gives us a method of preparing mixed secondary amines. It is sufficient to pass a mixture of a primary amine and an aliphatic, aromatic, or cyclohexyl alcohol in equivalent amounts over thoria at about 320°. Among the aliphatic alcohols, methyl gives the poorest results. Ethyl-isoamyl-amine, boiling at 126°, propyl-isoamyl-amine, boiling at 145°, and isobutyl-isoamyl-amine, boiling at 158°, have been prepared in this manner. 12

SABATIER and MAILHE, Compt. rend., 153, 160 (1911).

[•] SABATIER and MAILHE, Compt. rend., 153, 1204 (1911).

¹⁶ MAILHE, Bull. Soc. Chim. (4), 15, 327 (1914).

¹¹ SABATIER and MAILER, Compt. rend., 153, 1204 (1911).

¹² SABATIER and MAILHE, Compt. rend., 148, 900 (1909).

739. By associating cyclohexyl-amins with various aliphatic alcohols, with benzyl alcohols, and with cyclohexanol and its homologs, a large number of mixed secondary cyclohexyl-amines can be prepared.¹³

Thus methyl alcohol gives methyl-cyclohexyl-amins, boiling at 145°, while ethyl and other primary alcohols give the corresponding mixed amines with still better yields. Isopropyl-cyclohexyl-amins ¹⁴ and benzyl-cyclohexyl-amines have been made thus.

Cyclohexanol itself gives di-cyclohexyl-amins identical with that obtained in the hydrogenation of aniline (466). The three methylcyclohexanols give the three methylcyclohexyl-cyclohexyl-amines.¹⁵

740. At higher temperatures the aromatic amines can undergo similar reactions. By passing over alumina at 400-430° a mixture of aniline vapors and methyl alcohol in excess, the immediate formation of methyl-aniline is obtained and of dimethyl-aniline, resulting from the action of the methyl alcohol on the methyl-aniline.

Likewise o.toluidine is completely transformed by methyl alcohol over thoria into methyl-o.toluidine and then into dimethyl-o.toluidine. Similar results are obtained with meta and para toluidines. A single passage over the catalyst produces about equal proportions of the mono- and di-methyl compounds, and a second passage completes the substitution.¹⁶

By causing ammonia to act on a mixture of two alcohols, the primary and secondary amines corresponding to each alcohol are obtained and some of the mixed secondary amine. This has been found true with a mixture of propyl and isoamyl alcohols at 330°.

- 741. Alkyl-piperidines. The above method can be applied to piperidine with various alcohols over thoria at 350°. The results are satisfactory with propyl alcohol which yields only a little propylene and gives N-propyl-piperidine, boiling at 149°, and with isoamyl alcohol which furnishes N-isoamyl-piperidine, boiling at 186°, but are poor with cyclohexanol which gives much cyclohexene and only a little N-cyclohexyl-piperidine, boiling at 216°.17
- 742. Pyrrol. An analogous reaction is carried out by the aid of zinc dust with a mixture of *ethyl alcohol* and *pyrrol* which give α -*ethyl-pyrrol*.¹⁸
 - ¹² Sabatier and Mailhe, Compt. rend., 153, 1207 (1911).
 - ¹⁴ MAILHE and AMOROUX, Bull. Soc. Chim. (4), 15, 777 (1914).
 - 15 SABATIER and MAILHE, Compt. rend., 153, 1207 (1911).
 - ¹⁶ Mailhe and de Godon, Compt. rend., 166, 467 and 564 (1918).
 - 17 GAUDION, Bull. Soc. Chim. (4), 9, 417 (1911).
- ¹⁸ Dennstedt, Berichte, 23, 2563 (1890). Zanetti, Gas. Chim. Ital., 21 (2), 167 (1891).

§ 4. — ELIMINATION OF WATER BETWEEN AN ALCOHOL AND HYDROGEN SULPHIDE

Synthesis of Mercaptans

743. If the direct action of alcohols on the dehydrating oxides, such as thoria, gives rise to the formation of a sort of unstable ester (thorinate), it can be predicted that when this is brought into contact with an acid more energetic than the hydrate of the oxide, such acid will displace the oxide at least in part to give a new ester. We will have:

$$\frac{\text{ThO}(\text{OC}_n\text{H}_{2n+1})_2}{\text{thorinate}} + 2\text{AH} = \underbrace{2\text{A.C}_n\text{H}_{2n+1}}_{\text{ester}} + \text{ThO}_2 + \text{H}_2\text{O}$$

and if the acid is incapable of forming a stable salt with thoria as a base, the thoria will be regenerated and will react with a new portion of alcohol to repeat the cycle.

Sabatier and Mailhe believed that hydrogen sulphide, which does not react with thoria (nor with alumina), would act in this manner, since it appears to be a stronger acid than thoria. We would have in succession:

$$\frac{\text{ThO}(OC_nH_{2n+1})_2}{\text{thorinate}} + 2H_2S = \underbrace{2C_nH_{2n+1}.SH}_{\text{mercentan}} + \text{ThO}_2 + H_2O$$

and then, with greater difficulty, on account of the acid function still remaining in the mercaptan:

$$\frac{\text{ThO}(OC_nH_{2n+1})_2 + 2C_nH_{2n+1}.SH = 2(C_nH_{2n+1})_2S + \text{ThO}_2 + H_2O.}{\text{thorinate}}$$

The thoria being regenerated can react with a fresh portion of alcohol and if the hydrogen sulphide continues to act, the thoria can function indefinitely as a catalyst to produce mercaptans and alkyl sulphides, provided that the reaction of the hydrogen sulphide on the unstable thorinate is more rapid than the decomposition of the thorinate into the unsaturated hydrocarbon, water and thoria.

744. Experiment has shown that this is usually the case. This is a direct method for the preparation of mercaptans from the alcohols. It is sufficient to pass a mixture of the alcohol vapors and hydrogen sulphide over a train of thoria maintained between 300 and 380°. The mercaptan along with a small amount of the neutral sulphide is condensed with the water and unchanged alcohol.

A portion of the alcohol is dehydrated to the unsaturated hydro-

carbon, but with the primary aliphatic alcohols this is not important, provided the reaction temperature is not too high, but it is considerable with the secondary alcohols which decompose into hydrocarbons more readily.

Methyl, ethyl, propyl, isobutyl, and isoamyl mercaptans have been thus prepared with yields above 75%, so long as the condensation of the products is efficient. The yield is equally good for allyl mercaptan from allyl alcohol. Benzyl alcohol gives a rather large proportion of benzyl mercaptan and some sulphide. 19 20

745. The yields are less satisfactory, hardly above one third, when secondary alcohols are used. The following mercaptans have been obtained in this way: propane-thiol(2), pentane-thiol(3), heptane-thiol(5), 2.4-dimethyl-pentane-thiol(3), cyclohexyl mercaptan and the three o.m. and p.methyl-cyclohexyl mercaptans, 21 and also the mercaptan from benzhydrol, C₆H₅.CH(SH).C₆H₅, boiling at 278°.22

746. Various other catalytic oxides have been found to be inferior to thoria. With isoamyl alcohol and thoria maintained at 370-80°, the approximate yields of mercaptans for 100 parts of alcohol destroyed were:

Thoria						70
Zirconia						44
Uranous oxide						30
Blue oxide of tungsten						22
Chromic oxide						18
Blue oxide of molybdenum						
Alumina						10

Alumina gives amylene chiefly.22

They find that the amounts of unsaturated hydrocarbons formed are surprisingly low, usually only 2 to 3%, while considerable amounts of the aldehydes, 7 to 15% (estimated by the hydrogen produced), are formed. — E. E. R.

¹⁹ SABATIER and MAILHE, Compt. rend., 150, 1217 (1910).

²⁶ Working at 360-380° Kramer and Reid [J. Amer. Chem. Soc. 43, 887 (1921)], obtain the following yields from the alcohols named: methyl 42%, ethyl 35%, propyl 45%, n.butyl 52%, isobutyl 36%, isoamyl 42%. A part, at least, of the discrepancy between these figures and those given by Sabatier and Mailhe is due to a different method of estimating the mercaptan produced.

²¹ MAILHE, Bull. Soc. Chim. (4), 15, 327 (1914).

²² SABATIER and MAILHE, Bull. Soc. Chim. (4), 11, 99 (1912).

²² SABATIER and MAILHE, Compt. rend., 150, 1569 (1914).

§ 5. — ELIMINATION OF WATER BETWEEN ALCOHOLS AND ACIDS

Esterification

747. It is known that the formation of esters by the direct action of organic acids on alcohols takes place very slowly at ordinary temperatures and that the transformation is never complete as it is limited by the inverse action of water on the ester. Several years of contact are required for this limit to be reached. Elevation of the temperature hastens the reaction greatly but it still requires considerable time, several days at 110°, several hours at 156°.

The production of ester is very slow in the gaseous state also, even at temperatures above 250°: when a mixture of equivalent amounts of the vapors of ethyl alcohol and acetic acid is passed through a tube heated above 250°, the esterification effected is entirely negligible.

But either in the liquid or in the vapor condition, the presence of small amounts of catalysts accelerates the production of ester enormously so that the limit is soon reached.

Esterification by Catalysis in the Liquid State

- 748. The catalysts for esterification in liquid system are chiefly the strong mineral acids, hydrochloric and sulphuric, and several salts, ammonium salts, alkaline bisulphates, zinc chloride, sodium acetate mixed with water.
- 749. Catalysis by Mineral Acids. When equal molecules of ethyl alcohol and acetic acid are mixed and the mixture is distilled, the amount of ester produced is less than 1%.

But a long time ago, Berthelot found that it is sufficient to add to a mixture of an organic acid and an alcohol a few per cent of hydrochloric or sulphuric acid to cause an abundant formation of ethyl acetate, benzoate, etc.²⁴ He showed that traces of sulphuric acid are sufficient for the preparation of ethyl acetate.²⁵

750. To a mixture of equal molecules of ethyl alcohol and acetic acid (106 g.) small quantities of hydrochloric acid were added, namely:

To the first 0.67 g. or 0.017 molecule second 4.77 g. or 0.125 molecule third 11.84 g. or 0.33 molecule

²⁴ Berthelot, Bull. Soc. Chim. (2), 31, 342 (1879).

BERTHELOT and JUNGFLEISCH, Traité de Chim. Organ., 3rd Ed. 1886, I, 208.

The amounts of ester formed were as follows:

	At ordinary temperature								
	First	Second	Third						
Immediately after mixing	9.6%	58.7 %	82.3 %						
After six hours	9.6	73.6	75.8						

The limit without the mineral acid would be 66.6%: this limit is raised by the presence of the hydrochloric acid, and is practically attained in six hours with the above mixtures. In the cold, without this acid, several years would have been required. Besides, no ethyl chloride was formed.

751. Analogous results were obtained with sulphuric acid. To a mixture of 1 mol. ethyl alcohol, 1 mol. acetic acid and 0.5 mol. water was added 0.02 mol. (about 2 g.) sulphuric acid and in 24 hours in the cold, the esterification had reached 59.6%. In 2 hours at 100°, 60.6% was reached, which is the limit for this system.

By boiling under reflux a mixture of 25 cc. propionic acid, 25 cc. propyl alcohol, and 50 cc. 5% sulphuric acid, the proportion of ester was: 28

After	0.5	hour									45.1 %
	1	hour									51.8
	2	hours									56.9
	3	hours									58.3

752. The action of the sulphuric acid can be explained by the formation of acid ethyl sulphate, the immediate product of the action of the sulphuric acid, and the action of which on the acetic acid would produce ethyl acetate and regenerate sulphuric acid, which would renew the action. In the case of hydrochloric acid, Berthelot explains the accelerating action by assuming the formation of an addition product of the hydrochloric acid and the alcohol.^{29 20}

Bodroux has proposed a different explanation based on the temporary formation of an addition compound of the mineral acid cata-

- ²⁶ Berthelot explained the elevation of the limit by the taking part of the hydrochloric acid in the equilibrium, in which it increases the total amount of acid relative to the alcohol.
 - ²⁷ BERTHELOT, Bull. Soc. Chim. (2), 31, 342 (1879).
 - 28 Bodroux, Compt. rend., 157, 939 (1913).
 - 29 BERTHELOT, Bull. Soc. Chim. (2), 31, 342 (1879).
- ³⁰ It is curious how many chemists have given entirely different explanations for the action of hydrochloric and sulphuric acids. All the facts go to show that all acids act alike and that whatever explanation is given in any one case must fit all others. E. E. R.

lyst with the organic acid considered as the anhydride of an ortho acid:

$$AH + R.C$$
 $OH = R.C$
 OH
 AH

then:
$$R.C \stackrel{OH}{\underset{A}{\smile}} + R'OH = AH + R.C \stackrel{OH}{\underset{OR'}{\smile}}$$

and finally by the immediate spontaneous loss of water: 31

$$R.C\sqrt{\frac{(OH)_2}{OR'}} = H_2O + R.CO.OR'.$$

753. Many chemists still continue to think that the presence of a large amount of the mineral acid is favorable to esterification and it has become common usage to saturate the mixture of alcohol and acid with hydrogen chloride when preparing esters. Many seem to have forgotten that the same end can be attained by employing very small proportions of acids as catalysts. In 1895 Emil Fischer and Speier made exact measurements on this matter and showed that the use of small quantities of the mineral acids makes the operation more convenient and leads to satisfactory yields.**

754. Thus for the preparation of ethyl benzoate, the classical method was to saturate with hydrogen chloride a mixture of 1 part of benzoic acid with 4 parts ethyl alcohol, which gave only 73% yield. Erdmann recommended heating on the water bath for 10 to 12 hours 1 part of the acid, 0.8 of alcohol and 0.4 of concentrated sulphuric acid, the yield being 75%.

By dissolving 3% hydrogen chloride in a mixture of 2 parts of alcohol to 1 part of acid, Emil Fischer obtained 76% of the ester, while for 1% hydrogen chloride, the yield was 64.5% for the same time of heating.

755. The use of sulphuric acid is very advantageous. A mixture of 1 part of benzoic acid, 2 parts of alcohol and 0.2 part concentrated sulphuric acid is heated 3 hours under reflux and a practical yield of 90% is obtained. If account is taken of inevitable losses during the washings, the yield is practically quantitative and as the excess of alcohol can be recovered almost entirely, the operation is very advantageous economically.

²¹ Bodroux, Compt. rend., 157, 1428 (1913).

²⁵ E. FISCHER and SPEIER, Berichte, 28, 3252 (1895).

756. Emil Fischer has shown that this process can be applied not only to the aliphatic acids as Berthelot had found, and to benzoic acid, but also to a large number of types of acids whether aliphatic or aromatic:

```
Monobasic acids (naphthoic, phenyl-acetic);
Unsaturated monobasic acids (crotonic, cinnamic);
Saturated dibasic acids (succinic, phthalic), or unsaturated (fumaric);
Hydroxy-acids (glycolic, phenyl-glycolic);
Phenol-acids (salicylic);
Ketone-acids (lasvulinic);
Polybasic hydroxy-acids (malic, tartaric, citric, mucic).
```

The yields obtained are usually very satisfactory; we quote some of the results obtained by heating for 4 hours a mixture of 1 part of the acid with 3 to 4 parts of ethyl alcohol.

	% Catalyst	% Ester
α -Naphthoic acid	2.2 HCl	74.8
Phenyl-acetic		87.0
Cinnamic	0.7 HCl	78.8
Cinnamic	7.5 H ₂ SO ₄	89.7
Crotonic	7.5 H ₂ SO ₄	54.3
Phenyl-glycolic	2.2 HCl	67.5
Laevulinic		76.5
Succinic	0.8 HCl	73.9
Succinic	8.0 H ₂ SO ₄	73.9
Fumaric	0.8 HCl	68.2
Tartaric	0.8 HCl	72.8
Malic	0.8 HCl	70.5

757. To obtain slightly soluble esters, Bodroux adds to a mixture of an organic acid and alcohol its weight of pure commercial hydrochloric acid diluted with its own volume of water: in the cold the mixture becomes turbid in a few hours and finally gives 60 to 90% of ester. This process works well for phenyl-acetic acid with various saturated alcohols but not so well for benzoic, salicylic and cinnamic acids.¹²

The yields are less satisfactory, hardly more than 50%, with allyl alcohol or with the secondary alcohols, isopropyl and cyclohexyl. They are worse still with dimethyl-ethyl-carbinol as well as with glycerine and mannite.³⁴

²³ Bodroux, Compt. rend., 157, 939 (1913).

²⁴ Bodroux, Compt. rend., 157, 1428 (1913).

758. Senderens and Aboulenc, who do not seem to have known of the work of Berthelot and of Emil Fischer, have described as new the method of direct esterification of alcohols in presence of small amounts of sulphuric acid. The results which they give are a verification and extension to other alcohols of a part of the results of Emil Fischer. But they have thought that they were able to make an essential distinction in the mechanism of the reaction between the aromatic acids that can be regarded as substitution products of acetic acid, e.g. phenyl-acetic, on the one side and straight aromatic acids in which the carboxyl is attached to the nucleus, e.g. benzoic and toluic, on the other.

For the first class, they consider the speed of the esterification and the amount of ester formed as independent of the amount of the sulphuric acid, while for benzoic acid, for example, these increase with the amount of the acid, "which consequently does not act simply as a catalyst."

This distinction can not be admitted. A solid catalyst, up to a certain limit, acts in proportion to its active surface. Soluble catalysts, such as diastases or acids in hydration reactions, or sulphuric acid in this case, act proportionally to their mass, at least if this is not too large for the total volume of liquid, and this is as true for acetic as for benzoic. The results quoted above from Berthelot for the formation of ethyl acetate in the presence of hydrochloric acid, show that, in the cold, the rapidity of the esterification is approximately proportional to the amount of the catalyst.

The difference between the aliphatic acids and their analogs and benzoic acid is that the velocity of the esterification of the former by the catalytic acid is much greater than for benzoic. To obtain the same yield of ester from benzoic a larger amount of the catalyst would have been required.

Oxalic acid is esterified regularly like succinic.

Furthermore the practical yields are much better with the higher alcohols since with the less soluble esters the losses in the necessary washings with water and alkaline carbonate solutions are less serious.

759. According to the same authors the sulphuric acid can be replaced by double its weight of anhydrous aluminum sulphats or potassium bisulphats.²⁶

²⁵ The slowness of esterification of benzoic acid as compared with acetic acid is shown by the work of Freas and Reid, [(J. Amer. Chem. Soc., 40, 569 (1918)], who found it necessary to heat benzoic acid with methyl and ethyl alcohols to 200° for 96 hours to insure reaching the limit of esterification while Berthelot and St. Gilles found 24 hours sufficient even at 170°. — E. E. R.

²⁶ SENDERENS and ABOULENC, Compt. rend., 152, 1671 and 1855 (1911) and 153, 881 (1911).

760. Glycerine mixed with acetic acid (1 molecule of glycerine to 3 of the acid) gives on boiling under reflux for 1 hour, esterification amounting to 0.4 molecule of acid: by the addition of 5% potassium bisulphats, the amount esterified reaches 1.2 molecules.

With	2%	anhydrous	aluminum	sulphate					1.5 mol.
	1 %	sulphuric (acid		_	_	_		1.5 mol.

By starting with 1 molecule of glycerine and 12 acetic acid, the amount esterified by boiling 1 hour is:

Triacetine is thus reached and there would be no advantage in increasing the amount of the catalyst.**

761. Esterification by Acetanhydride. A common method of preparing the acetates of alcohols or of poly-alcohols is to heat them with acetanhydride.

$$R.OH + (CH_3.CO)_2O = 2CH_3.CO_2R + H_2O.38$$

By this means all the hydroxyl groups of a complex molecule are esterified. The presence of a certain amount of sodium acetate favors the action of the anhydride.

Still better results are obtained by adding to an alcohol four times its weight of acetanhydride and a small fragment of fused zinc chloride. The reaction becomes very rapid immediately. In the case of glycerine a veritable explosion is caused. With mannite, however, it is regular and yields in a few minutes mannite hexa-acetate, melting at 120°. 20°

Catalytic Esterification in Gaseous System

762. The assumption of an unstable combination between the dehydrating oxides and the alcohols has been a basis for the prediction of various reactions which have been realized by experiment, such as the formation of mercaptans and aliphatic amines. Sabatier and Mailhe thought that it might be expected that these combinations

$$R.OH + (CH_4.CO)_3O = CH_4CO_2R + CH_4CO_3H.$$

An excess of the anhydride is always used and the reaction goes to completion since no water is formed to reverse it. — E. E. R.

³⁷ SENDERENS and ABOULENC, Compt. rend., 158, 581 (1914).

³⁸ I think it better to write the reaction thus:

⁸⁹ Franchimont, Berichte, 12, 2059 (1879).

would play a part analogous to that of the acid sulphuric esters, that is, that the dehydrating oxides would act as esterification catalusts.40

763. As has already been indicated, if a mixture of the vapors of an alcohol and an organic acid be passed through a 60 cm. tube heated between 300 and 360°, the proportion of ester formed during the passage is absolutely negligible, but the presence of a catalytic oxids changes the case entirely. Let us suppose that the tube contains a catalytic oxide, MO, derived from the metallic hydroxide, M(OH)₂, an amphoteric hydroxide.

The reaction can take three different courses:

1st. The acid may combine to form a salt, unstable for those oxides which catalyze acids, and breaking down to regenerate the oxide and forming a symmetrical ketone (837):

(1)
$$MO + 2R.COOH = H_2O + (R.COO)_2M = H_2O + MO + CO_3 + \frac{R.CO.R.}{betons}$$

2nd. The oxide may combine with the alcohol to form an unstable salt:

$$MO + 2C_nH_{2n+1}.OH = H_2O + M(OC_nH_{2n+1})_2$$

This unstable complex can decompose in two ways, either by itself to give the unsaturated hydrocarbon: 41

(2)
$$M(OC_nH_{2n+1})_2 = MO + H_2O + 2C_nH_{2n}$$

or with the aid of the acid to form an ester:

(3)
$$M(OC_nH_{2n+1})_2+2R.CO.OH=MO+H_2O+2R.CO.OC_nH_{2n+1}.$$

In any case the catalytic oxide is regenerated and can continue the same effects. Furthermore in reaction (3), the water produced tends to destroy the combination, $M(OC_nH_{2n+1})_2$, and consequently *limits* the formation of the ester which results from it. Since these reactions are very rapid, the esterification limit will be reached quickly, the catalytic oxide acting like the platinum sponge in the combination of iodine and hydrogen (19).

764. We may have simultaneously formation of a *kstons*, production of unsaturated hydrocarbon (or ether), and the rapid reversible formation of the ester; this is what is observed when a mixture of the vapors of ethyl alcohol and acetic acid is passed over *thoria* or alumina heated to about 400°.

If the conditions are such that reactions (1) and (2) do not take place, (3) will be the only one and we will have an advantageous catalytic formation of ester.

⁴⁰ SABATIER and MAILHE, Compt. rend., 150, 823 (1910).

⁴¹ In the case of methyl alcohol, this decomposition gives methyl ether.

To obtain this result it is necessary to operate at such low temperatures that the acids are not decomposed and that the decomposition into unsaturated hydrocarbon is not too rapid.

765. Thoria, which is the most active catalyst for the destruction of acids and which, likewise, has a powerful dehydrating effect on alcohols, would doubtless be less advantageous than titania, which produces these effects less vigorously.

766. With aromatic acids, such as benzoic and its homologs which have the carboxyl attached to the nucleus, thoria does not produce any appreciable decomposition even up to 450°: it can be predicted that reaction (1) will not take place. Experiment has shown that this is the case and that at 350° reaction (2) is negligible as compared with reaction (3), which goes very rapidly. By vaporizing a saturated solution of benzoic acid in an alcohol (there are at least 12 molecules of the alcohol to 1 of the acid) and passing the vapors over a train of thoria at 350°, there is no appreciable formation of the unsaturated hydrocarbon, but the benzoic acid is almost totally esterified. Methyl, sthyl, propyl, butyl, isobutyl, isoamyl and allyl benzoates have been obtained advantageously in this way.

In spite of their greater tendency to form unsaturated hydrocarbons, the secondary alcohols can form benzoic esters with fairly good yields: this is the case with *isopropyl alcohol* with which the formation of propylene is of minor importance. Cyclohexyl alcohol is more delicate, but nevertheless gives a fairly good yield of the benzoate.

Analogous results are obtained with the three toluic acids which are readily esterified by thoria at 350–380°, but the practical preparation of these esters is less advantageous on account of the smaller solubility of these acids, particularly the para, in the alcohols: the meta is the most soluble.⁴²

767. Titania enables us to esterify various acids in the same manner. If a mixture of equivalent amounts of the vapors of a primary alcohol and an aliphatic acid, other than formic, is passed over a train of this oxide maintained at 280–300°, rapid esterification takes place, reaching a limit slightly above that observed by Berthelot and Menschutkin in their experiments on direct esterification. The production of gas on account of the destruction of the acid or the alcohol is absolutely negligible.

768. It is known that the presence of a catalyst does not change the location of the limit in reversible reactions, but diminishes greatly the time required to reach that limit. In this particular case, Berthe-

SABATIER and MAILHE, Compt. rend., 152, 358 (1911).

lot found that the limit is moved somewhat by elevation of the temperature. For equivalent amounts of ethyl alcohol and acetic acid, he found the following values of the limit:

In the cold (10 years)						65.2 %
At 100° (200 hours) .						65.6
170° (42 hours)						66.5
200° (24 hours)						

The figures show that the limit is not fixed but progresses slowly with the temperature and suggest a still higher value for the limit at 280-300°.

769. At 155°, Menschutkin found for various alcohols mixed with equivalent amounts of different acids, the following limits: 4

Acetic acid	+ methyl alcohol .			69.6 %
Acetic acid	+ ethyl alcohol			66.6
Acetic acid	+ propyl alcohol .			66.9
Acetic acid	+ butyl alcohol			67.3
Acetic acid	+ isobutyl alcohol.			67.4
Propionic acid	+ isobutyl alcohol			68.7
Butyric acid	+ isobutyl alcohol			69.5
Isobutyric acid	+ isobutyl alcohol			69.5

Sabatier and Mailhe obtained the following limits with titania at 280-300°:

```
Acetic acid
                   + isobutyl alcohol . . . .
                                                69.5%
                   + methyl alcohol . . . . .
Propionic acid
                                                72.9
                   + isoamyl alcohol . . . .
Propionic acid
Butyric acid
                   + ethyl alcohol . . . . .
                                                71
Butyric acid
                   + isoamyl alcohol . . . . .
Isobutyric acid
                   + ethyl alcohol . . . . . .
                                                71
```

These values are slightly higher than the corresponding figures obtained at lower temperatures.

770. Furthermore, in this rapid catalytic esterification, the same laws are found to hold as Berthelot formulated for direct esterification. An excess of one constituent increases the amount of the other combined. Thus for 1 molecule of isobutyric acid with 1, 2, and 4 molecules of ethyl alcohol, the following percentages of the acid were esterified:

With 1 molecule							•		71.0%
2 molecules									83.5
4 molecules		•	•		•	•			91.0

MENSCHUTEIN, Ann. Chim. Phys. (5), 20, 289, and 23, 64 (1880).

In the presence of more than 10 molecules of alcohol the esterification of the acid is nearly complete and, conversely, almost all of the alcohol is esterified by a large excess of the acid. The relative cost of the alcohol and acid in such cases decides which conditions are most economical.

771. Sabatier and Mailhe have prepared easily the methyl, ethyl, propyl, butyl, isobutyl and isoamyl esters of acetic, propionic, butyric, isobutyric, isobutyric, caproic, pelargonic and crotonic, etc., acids.

Bensyl alcohol gives equally good results with various acids. The dehydration to the resinous hydrocarbon, $(C_7H_6)_x$, which is effected so rapidly by catalytic oxides, hardly takes place at all in the presence of acid vapors.⁴⁴

772. Sabatier and Mailhe have found further that it is not indispensable to use as high a temperature as 280°, which is usually the most advantageous.

The catalytic activity continues, though it falls off gradually, to temperatures much lower where the acids and alcohols are stable. In this *titania* is superior to thoria.⁴⁵ By operating with equal molecules of ethyl alcohol and acetic acid and passing the vapors over a 50 cm. train of the oxide at the rate of 0.2 molecule, or 21 g., per hour, Sabatier and Mailhe obtained the following percentages of esterification:

						W	ith thoria		N	ith titania
At	150°.						11 %			20 %
	170°.						26			
	230°.						45			60

Besides, the catalytic power of titania persists almost indefinitely; it was not diminished by experiments on varied mixtures of alcohols and acids extending over 20 days.

773. Formic acid can be esterified at these temperatures at which it is fairly stable. By operating with equal molecules of formic acid and ethyl alcohol, distributed by the same capillary tube through which the molecular volume passed very rapidly, in spite of this unfavorable circumstance, the following amounts were esterified over titania:

At	120°										47 %
	150°										65 %

The esterification limit is nearly reached even at 150° at which the decomposition of formic acid into gaseous products is still inconsiderable.

- 44 SABATIER and MAILHE, Compt. rend., 152, 494 (1911).
- * MAILHE and DE GODON [(Bull. Soc. Chim., 29, 101 (1921)] conclude that ZrO₁ is as good as or better than TiO₂. MILLIGAN and REID (unpublished work) find silica gel to be a better esterification catalyst than either. E. E. R.

In practice formic acid, mixed with an excess of the desired alcohol, is passed over thoria at 150°. Methyl, ethyl, propyl, butyl, isoamyl and benzyl for mates have been readily prepared in this way.

774. The comparison of these results has led Sabatier and Mailhe to conclude that the rapidity of the esterification of the primary alcohols by the aliphatic acids, in presence of catalysts, is directly proportional to the kinstic velocities of the reacting molecules; it is as much greater as the molecules are lighter and it can be inferred that the reason is to be found in the greater rapidity of gaseous interchanges on the catalyst.

775. The secondary alcohol, isopropyl, mixed with isobutyric acid does not give any evolution of propylene with titania below 300°. The proportion esterified was:

\mathbf{At}	235°										16.5 %
	256°										21
	292°										37

For primary propyl alcohol, the amount is 50% at 235° and 72% at 292°.

776. Trimethyl-carbinol (tertiary butyl alcohol) likewise mixed with isobutyric acid, gives 6% ester at 235° with no formation of the hydrocarbon. With the isomeric primary alcohol, isobutyl, it is 22%. It is only at 255° that the decomposition into butylene begins to manifest itself. At 265° it is quite rapid and the acidity of the mixture increases on account of the destruction of the alcohol in place of diminishing by esterification.

777. These results agree well with the weakening of the alcoholic function in secondary and particularly in tertiary alcohols. The velocity of the catalytic esterification should be at the same time a function of the speed of the gaseous interchanges, in consequence of the smallness of the molecules and also of the facility with which the alcohol forms the temporary unstable complexes with the catalytic oxide.⁴⁶

778. Beryllium oxide also can be employed as an esterification catalyst. With this oxide heated to 310°, yields of above 70% of ester can be obtained. The catalyst can be regenerated by calcining at a red heat. With this catalyst esters of tertiary alcohols and of high molecular weight acids can be prepared.⁴⁷ 48

⁴⁶ SABATIER and MAILEE, Compt. rend., 152, 1044 (1911).

⁴⁷ HAUSER and KLOTS, Chem. Zeit., 37, 146 (1913).

⁴⁸ I have tried to prepare esters by the use of beryllia and so has Dr. Milligan but neither of us has been able to verify the statements of Hauser and Klotz.

— E. E. R.

§ 6. — ELIMINATION OF WATER BETWEEN ALCOHOLS AND ALDEHYDES OR KETONES

779. The elimination of water between alcohols and aldehydes or ketones can take place in several ways. The one way is to a certain extent comparable to *esterification* and leads to *acetals*; it can hardly be realized except in liquid systems. The other, more exceptional, gives rise to *hydrocarbons* and is effected in gaseous systems.

I. — Formation of Acetals

780. Aldehydes can combine directly with alcohols to give acetals:

$$\frac{R.CHO}{\text{alcohol}} + \frac{2R'OH}{\text{alcohol}} = H_2O + \frac{R.CH(OR')_2}{\text{acetal}}.$$

But the direct formation is very imperfect, unless suitable catalysts are used.

Good yields are obtained by passing for a long time a current of pure *phosphins* through a well cooled mixture of the aldehyde and alcohol: by this means *acstaldshyds* has been made to combine with *ethyl*, *propyl* and *isobutyl* alcohols.⁴⁹

The combination of alcohols and aldehydes is greatly aided by the presence of a certain amount of glacial acetic acid. 50

- 781. Trioxymethylene, the condensation product of formaldehyde, readily forms methylal, HCH(OCH₂)₂, when it is mixed with *methyl alcohol* and heated on the water bath for 10 hours with 3% of ferric chloride.
- 782. A good method for preparing acetals is to mix the aldehyde with the proper amount of alcohol containing 1% of hydrogen chlorids (the gas dissolved) and digest the mixture for 18 to 20 hours: the yields are usually satisfactory.⁵¹

To obtain acetals from acetaldshyds with various aliphatic alcohols, 40 g. of acetaldshyde is mixed with 60 g. of the alcohol and 1 cc. concentrated hydrochloric acid is added and this mixture digested 24 hours with a saturated solution of sodium chloride and 10 g. of the solid salt.²²

783. The action of ethyl ortho-formate on aldehydes or ketones readily produces their combinations with ethyl alcohol; but this re-

- Engel and Girard, Jahreeb., 1880, 694.
- 56 GEUTHER, Annalen, 126, 62 (1863).
- E. FISCHER and GIEBE, Berichte, 30, 3053 (1897).
- ** King and Mason, English patent, 101,428 of 1916, J. Soc. Chem. Ind., 35, 1131 (1916).

action does not take place without the aid of suitable catalysts. These may be quite varied, e.g. strong mineral acids, ferric chloride, ammonium chloride, ethyl-, disthyl-, or tristhyl-amine hydrochlorides, potassium bisulphate, ammonium sulphate or nitrate. Boiling for a few minutes is sufficient to assure the formation.

Thus to prepare the acetal from *ethyl alcohol* and *bensaldehyde*, 1 molecule of the aldehyde is mixed with 0.1 molecule ethyl ortho-formate and poured into 3 molecules of the alcohol and a little dry hydrogen chloride is passed in. After ten minutes boiling, the acetal, C_0H_5 . CH $(OC_2H_5)_2$ is obtained in 99 % yield. By using 2 g. ammonium chloride, the yield is 97 %.

The diethyl acetal of acetons is obtained thus with 66% yield. If the boiling is prolonged too greatly, the yield is more and more diminished, which shows that the catalyst tends to destroy by hydrolysis the acetal which it has formed. **

II. - Formation of Hydrocarbons in Gaseous System

784. The dehydrating action of oxides such as alumina on a mixture of an alcohol and an aldehyde can eliminate all of the oxygen as water producing a doubly unsaturated hydrocarbon.

This takes place when a mixture of sthyl alcohol and acstaldshyds is passed over the impure alumina formed by calcining ammonium alum. Butadiens, boiling at 2°, is obtained:

$$CH_2OH.CH_3 + OCH.CH_3 = 2H_2O + CH_2:CH.CH:CH_3$$

With pure alumina, methyl-allens, CH₁.CH: C: CH₃, is also formed. This reaction can be applied to the synthesis of rubber by the polymerization of the hydrocarbon obtained (213). From 100 g. of the mixture of aldehyde and alcohol, 25 g. of the crude hydrocarbon may be obtained or 16 to 18 g. of pure butadiene which may be totally transformed into rubber.⁵⁴

Similarly acetaldehyde, with isopropyl or propyl alcohols, leads to piperylens, CH₃.CH: CH.CH: CH₃, boiling at 42°.54

⁵⁸ Claisen, *Berichte*, 40, 3903 (1907).

⁵⁴ OSTROMUISSLENSKII and Kelbasinski, J. Russian Phys. Chem. Soc., 47, 1509 (1915); C. A., 10, 3179 (1916).

CHAPTER XVII

DEHYDRATION (Continued)

§ 7. — DEHYDRATION OF PHENOLS ALONE

785. One method of preparing sthers from phenols is to distil dry aluminum phenylates: this works well for phenyl ether and for the ethers of ortho and para cresols. This method of preparation leads us to foresee that phenyl ethers can be prepared catalytically by the action of a catalytic oxide such as thoria on the vapors of the phenol at a suitable temperature, the mechanism of dehydration depending, as with the alcohols, on the formation of an unstable thorinate which decomposes regenerating thoria.

We have: $2C_6H_5.OH + ThO_2 = H_2O + Th(OC_6H_5)_2$ thorinate

and then: $Th(OC_6H_5)_2 = ThO_2 + \underbrace{(C_6H_5)_2O.}_{\text{ether}}$

This prediction having been verified, Sabatier and Mailhe have based on it an advantageous method for the preparation of phenol ethers by the use of thoria.²

786. Simple Phenol Ethers. The vapors of the phenol are passed over a train of thoria kept at 400-500°. If the phenol is a liquid, it is introduced directly by means of the capillary tube (181); if it is a solid, its benzene solution is used. The reaction products are shaken with caustic soda, which extracts the unchanged phenol leaving the ether which is obtained entirely pure by one distillation. Phenyl ether can be prepared in this way very economically and in great purity with a yield of 50% or better; meta and para cresyl ethers can be readily obtained, while ethers are more difficult to obtain from ortho cresol and from xylenol(1,3,4), and poor results are gotten with carvacrol.

787. Diphenylene Oxides. This method leads to the simultaneous formation of diphenylene oxides, fluorescent compounds, less volatile than the ethers, and formed by the loss of H₂.

With ordinary phenol at 475°, there is formed along with phenyl

- ¹ GLADSTONE and TRIBE, J. Chem. Soc., 41, 9 (1882), and 49, 25 (1886).
- ² Sabatier and Mailhe, Compt. rend., 151, 492 (1910).
- ³ SABATIER and MAILHE, Bull. Soc. Chim. (4), 11, 843 (1912).
- 4 SABATIER and MAILHE, Compt. rend., 158, 608 (1914).

ether, boiling at 253° and melting at 28°, a considerable amount of C₄H₄ O, boiling at 287° and melting at 85°, which

had previously been obtained by distilling calcium phenylate.⁵ The cresols, xylenols, and naphthols give rise to the formation of similar products.⁶

788. Mixed Phenol Ethers. By dehydrating over thoria, not a single phenol, but a mixture of two phenols, the product contains along with the simple ethers of the two phenols and the diphenylene oxides, an amount, usually considerable, of the mixed ether derived from the two phenols which can be separated by careful fractionation. Sabatier and Mailhe have prepared the following mixed ethers, phenylocaresyl, phenyl-m.cresyl, phenyl-p.cresyl, phenyl- α -naphthyl, phenyl-carvacryl, p.cresyl-carvacryl, as well as the phenylene-naphthylene oxides.

§ 8. — ELIMINATION OF WATER BETWEEN PHENOLS AND ALCOHOLS

Synthesis of Alkyl Phenol Ethers

789. Sabatier and Mailhe have shown that catalytic oxides such as thoria readily eliminate water from a phenol and an alcohol with the formation of a mixed ether. This is a very advantageous method of preparing mixed ethers. All that is necessary is to pass a mixture of the phenol with an excess of the alcohol over thoria at 390-420. With methyl alcohol, which is dehydrated by thoria very slowly, the results are particularly good. The excess of the alcohol and most of the unchanged phenol are separated from the ether by fractionation. The remainder of the phenol is extracted by caustic soda from the mixed ether, which is purified by a single distillation.

In this way, Sabatier and Mailhe have prepared the methyl ethers of phenol, the three cresols, xylenol(1,3,4), thymol, carvacrol, and α -and β -naphthols.

At the same time small quantities, more or less important according to the phenol, of the phenol ether and diphenylene oxide are obtained. A mixture of methyl alcohol and carvacrol gives methylcarvacryl ether, along with di-carvacryl ether and carvacrylene.

- ⁵ NIEDERHÄUSERN, Berichte, 15, 1120 (1882).
- SABATIER and MAILHE, Compt. rend., 151, 494 (1910).
- ⁷ Sabatier and Mailhe, Compt. rend., 155, 260 (1912), and 158, 608 (1914).
- * SABATIER and MAILHE, Compt. rend., 151, 359 (1910).
- * SABATIER and MAILHE, Compt. rend., 158, 608 (1914).

The other alcohols, in spite of their own rapid decomposition by thoria, can readily give the mixed ethers: one operates around 420° on phenol dissolved in excess of the alcohol, a part of which is decomposed forming the unsaturated hydrocarbon. *Ethyl-, propyl-,* and *isoamyl-phenyl* ethers have been prepared in this way.

§ 9. — ELIMINATION OF WATER BETWEEN A PHENOL AND AN AMINE

790. Nothing worth while is accomplished by passing a phenol and ammonia over a catalytic oxide at 400°. The production of amines is quite negligible.

We may mention as a catalytic reaction of this sort, the action of α - and β -naphthols, on anilins, the toluidines and other aromatic amines, when they are heated 7 hours to about 200° in the presence of 1% of iodins. The corresponding secondary amines are obtained with satisfactory yields.¹⁰

§ 10. — ELIMINATION OF WATER BETWEEN PHENOLS AND HYDROGEN SULPHIDE

791. Sabatier and Mailhe have found that by passing the vapors of a phenol and hydrogen sulphide over thoria between 430 and 480°, the corresponding thiophenol is obtained:

$$C_aR_a.OH + H_aS = H_aO + C_aR_a.SH.$$

But the yield is not so good as with alcohols (744), and is not above 10% in the most favorable case. A temperature of 500° decreases the yield on account of the serious decomposition of the hydrogen sulphide. Hence the reaction is of no practical use but is of only scientific interest.¹¹

The yields are still less when other oxides are used. With phenol, the following yields were obtained at 450°:12

Alumina	0.4 %
Zirconia	1.5
Blue oxide of molybdenum	1.8
Blue oxide of tungsten	1.5
Chromic oxide	2.5
Uranous oxide	3.8
Thoria	8.0

¹⁰ Knoevenagel, J. prakt. Chem. (2), 89, 16 (1914).

¹¹ SABATIER and MAILHE, Compt. rend., 150, 1220 (1910).

¹² SABATIER and MAILHE, Compt. rend., 150, 1570 (1910).

§ 11. — ELIMINATION OF WATER BETWEEN PHENOLS AND ALDEHYDES

792. For some years there has been prepared under the name of bakelite, a resinous material very resistant to shock and to pressure and endowed with insulating properties of the first order. It results from the condensation of phenol or cresols with formaldehyds in the presence of various catalysts, chiefly substances with alkaline reaction.¹² According to Backeland,¹⁴ who has given his name to the product, one of the materials, called bakelits C, results from the reaction of 7 molecules of formaldehyde with 6 of phenol:

$$6C_6H_5OH + 7CH_2O = C_{45}H_{25}O_7 + 6H_2O.$$

The formaldehyde can be replaced by methylal, trioxymethylene, or hexamethylene-tetramine.

The products obtained are very variable according to the operating conditions and either liquid intermediate substances or solids corresponding to an advanced stage of molecular condensation may be obtained. The condensation to the solid products can be effected by acid catalysts such as *hydrochloric* acid.

§ 12. — FORMATION OF PHENOLIC GLUCOSIDES

793. Quinoline employed in small proportion causes *phenols* to react with acetyl-brom-glucose forming the acetylate of the corresponding phenyl-glucosids. By warming for 1 hour 50 g. acetyl-brom-glucose with 160 g. phenol in the presence of 19 g. quinoline, the tetra-acetyl-phenyl-glucoside is obtained, the hydrolysis of which by baryta water separates the phenyl-glucoside. 15

§ 13. — DEHYDRATION OF ALDEHYDES OR OF KETONES

794. Frequently the presence of certain substances causes the condensation of two or more molecules of aldehydes or of ketones with the elimination of water and the formation of a single molecule retaining only one aldehyde or ketone group and containing double bonds.¹⁶

¹⁵ LEBACH, J. Soc. Chem. Ind., 32, 559 (1913). — Caoutchouc and Gutta-percha, 14, 9339 (1917). HUTIN, Ibid., 16, 9987 (1919).

¹⁴ BARKELAND, J. Ind. Eng. Chem., 1, 149 (1909).

¹⁴ E. FISCHER and VON MECHEL, Berichte, 49, 2813 (1916).

¹⁶ SABATIER and MAILHE, Compt. rend., 150, 1220 (1910).

This process is called *crotonization* from *croton aldehyde* which is formed from *acstaldehyde*:17

$$CH_{\bullet}.CHO + CH_{\bullet}.CHO = H_{\bullet}O + CH_{\bullet}.CH : CH.CHO.$$

Reactions of this kind can take place between molecules of different aldehydes or ketones.

Crotonization in Liquid Medium

795. The catalysts that are able to cause the crotonization of aldehydes and ketones in liquid medium are quite varied and their action is generally quite slow: soda, potash, hydrochloric acid, zinc chloride, lime, aluminum chloride and sodium acetats may be mentioned.

In order to transform acetaldehyde into croton aldehyde, it is heated to 97° for 36 hours with 20% of its weight of a water solution of sodium acetate, 18 or better to 100° for 48 hours with a solution of zinc chloride. 19

Paraldshyds in contact with sulphuric acid also gives croton aldehyde.20

The same process applies to the crotonization of propionic aldehyde which can be crotonized by heating with a solution of caustic soda.²¹ The same agent is employed for butyric aldehyde.²² Dry hydrogen chloride ²³ or a solution of sodium acetate may be used to crotonize isobutyric aldehyde.²⁴

Zinc chloride, or alcoholic potash, causes two or four molecules of heptaldehyde to condense.²⁶ Contact with zinc turnings is sufficient to crotonize isovaleric aldehyde: sodium, caustic potash and hydrochloric acid produce the same effect.²⁶

796. Croton aldehyde itself mixed with acetaldehyde and zinc chloride at 100°, undergoes a second like reaction and forms hexadienal (boiling at about 172°).²⁷

```
CH_{2}.CH:CHO+CH_{2}.CHO=
H_{2}O+CH_{2}.CH:CH.CH:CH.CHO.
```

- 17 SABATIER and MAILHE, Compt. rend., 150, 1570 (1910).
- 18 LIEBEN, Monatsh., 13, 519 (1892).
- 19 MULLER, Bull. Soc. Chim. (3), 6, 796 (1891).
- ²⁰ Delépine, Ann. Chim. Phys. (8), 16, 136 (1909), and 20, 389 (1910).
- ²¹ HOPPE, Monatsh., 9, 637 (1888).
- 22 RAUPENSTRAUCH, Monatsh., 8, 112 (1887).
- 28 ŒCONOMIDES, Bull. Soc. Chim. (2), 36, 209 (1881).
- 24 FOSSEK, Monatsh., 2, 616 (1881).
- ²⁶ Perkin, Berichte, 15, 2804 (1882).
- RIBAN, Bull. Soc. Chim. (2), 18, 64 (1872). Kekulé, Berichte, 3, 135 (1870). Borodin, Berichte, 6, 983 (1873).
 - 27 KEKULÉ, Annalen, 162, 105 (1872).

797. Ordinary acetone 28 kept for a long time in contact with lime, 29 or aluminum chloride 30 is transformed into mesityl oxide:

and then into phorons:

$$(CH_3)_2C: CH.CO.CH: C(CH_3)_2.$$

Cyclohexanons, in contact with sodium ethylate or hydrochloric acid, condenses to an oily compound similar to mesityl oxide, *1

$$CH_{2} \cdot CH_{2} \cdot CO \\ CH_{2} \cdot CH_{2$$

798. Crotonization can take place in a similar manner between different molecules, principally between a molecule of acetone and one or two molecules of aldehyde. The presence of aqueous or alcoholic soda is most frequently efficient in causing these condensations with the elimination of one or two molecules of water giving compounds containing a ketone group and one or two double bonds.

Benzaldshyds gives such products readily. Thus with acetone in prolonged contact with aqueous soda, it forms successively benzalacetone and dibenzal-acetone:

$$C_6H_6$$
.CHO + H_2 CH.CO.CH₈ = H_2 O + C_6H_6 .CH:CH.CO.CH₈ and

$$2C_6H_6.CHO + CH_2.CO.CH_3 = 2H_2O + (C_6H_6.CH:CH)_2CO.$$

In the presence of a little soda solution, o.nitrobenzaldehyde condenses with acetone to give o.nitrobenzal-acetone: 25

$$O_2N.C_6H_4.CHO + CH_3.CO.CH_3 = H_2O + O_2N.C_6H_4.CH : CH.CO.CH_3.$$

799. Benzaldehyde condenses with acetophenons in the presence of hydrogen chloride,³⁴ or of a few cubic centimeters of sodium methylate,³⁵ to give diphenyl-propenons, C₆H₅. CH: CH.CO.C₆H₅.

- ²⁸ Pure acetone passed over heated freshly prepared alumina forms condensation products, only about 60% of the acetone passing through unchanged. No gaseous products are formed. HOMER ADKINS.
 - 29 FITTIG, Annalen, 110, 32 (1859).
 - 30 LOUISE, Compt. rend., 95, 602 (1882).
 - ²¹ Wallach, Berichte, 29, 2955 (1896), C., 1897 (1), 322.
 - 22 CLAISEN and PONDER, Annalen, 223, 139 (1884).
 - BAEYER and DREWSEN, Berichte, 15, 2856 (1882).
 - ²⁴ Claisen and Claparede, Berichte, 14, 2463 (1881).
 - ³⁶ Claiben, Berichte, 20, 657 (1887).

The same aldehyde gives benzylidene-hydrindone with hydrindone and a little alcoholic potash:

$$C_0H_4 \xrightarrow{CH_3} CH_3 + OCH \cdot C_0H_4 \rightarrow C_0H_4 \xrightarrow{CH_3} C : CH \cdot C_0H_5.$$

Cinnamic aldehyds, digested several hours with acetophenone in contact with soda, passes into diphenyl-pentadieneone:

Cyclopentanons condenses with two molecules of benzaldehyds in contact with soda.²⁸

800. Citral (50 cc.) and acetone (65 cc.) shaken several hours with 1 l. of 4% baryta water condense to pseudo-ionone.²⁰

Condensations in Gaseous Phase

801. Catalytic dehydrating oxides can cause regular condensations of aldehydes or ketones in vapor phase.

The vapors of acetaldehyde, or of paraldehyde, passed over thoria at about 260° yield, along with a mixture of methane and carbon monoxide resulting from the decomposition of the aldehyde, a liquid containing water, crotonic aldehyde, hexadienal, without doubt associated with a certain amount of octatriensal. Careful hydrogenation of the liquid over nickel at 180°, gives essentially a mixture of normal butyl and hexyl alcohols.⁴⁰

The vapors of acetone passed over thoria at 410-20° give considerable mesityl oxide.41

Elimination of Water from a Single Molecule

- 802. We have seen (308 and 310) that the presence of certain catalysts permits the addition of a molecule of water to certain doubly unsaturated hydrocarbons, the products usually being aliphatic ketones. The inverse reaction, the formation of a doubly unsaturated hydrocarbon by the abstraction of a molecule of water from a ketone, can be realized also. It has been found that the vapor of methylisopropyl-ketone, passed, under reduced pressure, over kaolin between
 - ⁸⁶ Kipping, J. Chem. Soc., 65, 498 (1894).
 - ³⁷ SCHOLTZ, Berichte, 28, 1726 (1895).
 - ³⁸ Vorländer and Hobohm, Berichte, 29, 1840 (1896).
- ³⁰ Tiemann and Krüger, Berichte, 26, 2691 (1893). Bull. Soc. Chim. (3), 9, 798 (1893).
 - 44 SABATIER and GAUDION, Compt. rend., 166, 632 (1918).
 - 4 MAILHE and DE GODON, Bull. Soc. Chim. (4), 21, 63 (1917).

400 and 600°, gives isoprene (which doubtless results from the isomerization of 3-methyl-butadiene (1,2)). We would have:

$$\mathrm{CH_4.CO.CH} \overset{\mathrm{CH_4}}{\underset{\mathrm{CH_4}}{\longrightarrow}} + \mathrm{H_4O} + \mathrm{CH_2} : \mathrm{C} : \mathrm{C} \overset{\mathrm{CH_4}}{\underset{\mathrm{CH_4}}{\longrightarrow}} + \mathrm{CH_2} : \mathrm{CH.C} \overset{\mathrm{CH_2}}{\underset{\mathrm{CH_4}}{\longleftarrow}}.$$

Condensations of Aldehydes or Ketones with Various Organic Molecules

803. Condensations with elimination of water comparable to crotonizations can frequently take place between aldehydes or ketones and molecules of various kinds, nitro compounds, phenols, esters, indols, pyrrols, etc. These reactions are most frequently brought about by the usual condensing agents, sulphuric or hydrochloric acids, zinc chloride, etc., or ammonia and amines, or anhydrous aluminum chloride. The products are generally unsaturated at the point where the aldehyde or ketone groups have disappeared.

Thus benzaldehyde condenses with nitromethane in the presence of zinc chloride to give a nitro derivative of phenyl-ethylene: 42

$$C_6H_5$$
, $CHO + CH_2NO_2 = H_2O + C_6H_5$, $CH : CH . NO_2$.

804. The same aldehyde condenses with malonic acid or its esters to form benzylidene-malonic acid when heated in presence of alcoholic ammonia or of hydrochloric acid: 44

$$C_6H_6.CHO + H_2C(CO_2R)_2 = H_2O + C_6H_6.CH : C(CO_2R)_2$$

From equal molecules of benzaldehyde and malonic acid warmed 1.5 hours to 55° with an 8% alcoholic ammonia solution, a 60% yield of the condensed acid is obtained. The ammonia can be replaced by sthyl-amins or piperidins. 45

805. Acetone condenses with pyrrol on the addition of a few drops of concentrated hydrochloric acid to give a crystalline product, the molecule of which is doubtless quadruple the formula given:

$$C_4H_4N + CH_2 \cdot CO \cdot CH_2 = H_2O + C_7H_0N$$

- ⁴² Earle and Kyriakides, U. S. Patent, 1,106,290. J. Soc. Chem. Ind., 33, 942 (1914).
 - 4 PRIEBS, Annalen, 225, 321 (1884).
 - 44 CLAISEN, Berichte, 14, 348 (1881).
 - 4 KNOEVENAGEL German patents, 94,132, 97,735 and 164,296 (1904).

With 1 cc. hydrochloric acid, 14 g. pyrrol and 14 g. acetone dissolved in 80 cc. alcohol and heated, the yield is about 95%.46

806. Trioxymethylene can condense with benzene or its homologs in presence of anhydrous aluminum chloride to give at the same time diphenyl-methane (or a homolog) and anthracene:

$$4C_6H_6 + (CH_2O)_8 = C_6H_5 \cdot CH_2 \cdot C_6H_6 + C_{14}H_{10} + 3H_2O + H_2$$

Chloral and bromal can react in the same way in the presence of anhydrous aluminum chloride on various aromatic compounds with the elimination of water and the loss of the aldehyde function. This takes place with benzene and its homologs:

$$2C_6H_6 + CCl_3.CHO = H_2O + CCl_3.CH(C_6H_6)_2.$$

Resorcine (in carbon disulphide solution) gives a similar reaction but with the simultaneous loss of hydrochloric acid: 48

$$2C_6H_4(OH)_2 + CCl_2CHO = H_2O + HCl + CCl_2 : C[C_6H_4(OH)_2]_2.$$

Anisol reacts with chloral to give the compound, CCl₃. CH-(C₆H₄OCH₃)₂.⁴⁹

Naphthalene, anthracene, and phenanthrene react in an analogous manner with chloral and bromal in the presence of aluminum chloride but with the simultaneous loss of water and halogen hydride. Thus naphthalene gives the compound, $CCl_2: C(C_{10}H_7)_2.50$

§ 14. — ELIMINATION OF WATER BETWEEN ALDEHYDES OR KETONES AND AMMONIA

807. Catalytic oxides can bring about the condensation of aldehydes and ammonia in various ways.

Acetaldehyde and ammonia passed over alumina below 300° give a certain amount of pyrrol by simultaneous dehydration and dehydrogenation: 51

$$CH_3.CHO + NH_3 = 2H_2O + H_2 + CH:CH$$
 $CH_3.CHO$

- ⁴⁶ Chelintzev and Trunov, J. Russian Phys. Chem. Soc., 48, 105 (1916); C. A. II. 452 (1917).
 - 47 Frankforter and Kokatnur, J. Amer. Chem. Soc., 36, 1529 (1914).
 - 48 Frankforter and Daniels, J. Amer. Chem. Soc. 36, 1511 (1914).
 - 49 Frankforter and Kritchevsky, J. Amer. Chem. Soc., 37, 2560 (1915).
 - ⁵⁰ Frankforter, J. Amer. Chem. Soc., 37, 385 (1915).

Acetaldehyde and bensaldehyde carried over alumina by ammonia at above 300°, vield α- and γ-phenyl-puridines: 51

$$C_6H_5$$
. CHO + 2CH₂. CHO + NH₂ = 2H₂ + 3H₂O + C_5H_4N . C_6H_2 .

808. Aldehydes and ammonia passed over thoria at 420-50° give, by simultaneous dehydration and dehydrogenation, a considerable proportion of nitriles:

$$R.CHO + NH_s = R.CN + H_sO + H_s$$

With isovaleric aldehyde, the yield of nitrile reaches 40% and equally good results are obtained with isobutyric, propionic, and even benzoic and anisic aldehydes. 52

809. In contact with thoria at 300-400°, ketones and ammonia give ketimines. With benzophenone the yield is almost theoretical. We have

$$R.CO.R' + NH_4 = H_4O + \frac{R'}{R}C: NH.$$

§ 15. — ELIMINATION OF WATER BETWEEN ALDEHYDES AND HYDROGEN SULPHIDE

810. In contact with alumina below 300°, acetaldehyde condenses with hydrogen sulphide with simultaneous dehydration and dehydrogenation to give thiophene: ⁵¹

$$2CH_{2}CHO + H_{2}S = H_{2} + 2H_{2}O + C_{4}H_{4}S$$

§ 16. — DEHYDRATION OF AMIDES

811. The dehydration of *amides* to *nitriles* can be effected by appropriate catalysts. The amide mixed with the catalyst is heated to 250-60° for 4 hours in a flask fitted with a reflux condenser. Four parts by weight of catalyst are used to one of amide.⁵⁴

Acetamide gave the following yields of acetonitrile:

With alumina									68 %
lamp blac									
pumice									
powdered									
sand .	-								

⁸¹ CHICHIBABIN, J. Russian Phys. Chem. Soc., 47, 703 (1915); C. A., 9, 2512 (1915).

MAILHE and DE GODON, Compt. rend., 166, 215 (1918).

¹⁰ Mignonnac, Compt. rend., 169, 237 (1919).

⁵⁴ Boehner and Andrews, J. Amer. Chem. Soc., 38, 2503 (1916).

But better yields are obtained by carrying the amide in a current of air over the catalyst heated to 420°, the yields being: 55

With pumics									91.5
alumina									82
sand .									86.5
graphite									75.5

812. The same process can be applied to nascent amides furnished by the vapors of the acid with ammonia in excess in the presence of alumina or thoria at around 500°. Alumina gives the best results. Starting with acetic acid an 85% yield of the nitrile is obtained.⁵⁶

813. We may put along side of the catalytic dehydration of amides to nitriles, the action of ammonia gas on the chlorides of acids in the presence of catalytic oxides. The amide formed is immediately dehydrated to the nitrile.

The mixture of ammonia and the acid chloride is passed over alumina at 490-500° and water and hydrogen chloride are eliminated:

$$R.COCl + NH_2 = R.CN + H_2O + HCl.$$

High yields of the nitriles are obtained in this way from *propionyl*, *isobutyryl*, *isobutyryl* and *benzoyl chlorides*. As the ammonia gas is used in excess, ammonium chloride is formed and deposits in crystals in a receiver placed at the end of the reaction tube.⁵⁷

§ 17. — DEHYDRATION OF OXIMES

814. The aldoximss which are isomeric with the amides can be transformed into nitriles in the same way. The vapors of the aldoximes are passed over alumina or thoria maintained at 350-60° and give the nitriles. Isovalerald-oxime gave isovalero-nitrile and oenanthald-oxime gave hexyl cyanide. The ketoximes, when submitted to the action of these dehydration agents, undergo a complex reaction in which nitriles with one less carbon are formed.⁵⁸

§ 18. — DIRECT SULPHONATION OF AROMATIC COMPOUNDS

815. The direct sulphonation of aromatic compounds by means of concentrated sulphuric acid corresponds to the elimination of water and can be facilitated or modified by the presence of certain catalysts.

- ⁵⁵ BOEHNER and WARD, J. Amer. Chem. Soc., 38, 2505 (1916).
- ⁵⁶ VAN EPPS and REID, J. Amer. Chem. Soc., 38, 2128 (1916).
- ⁵⁷ MAILHE, Bull. Soc. Chim. (4), 23, 380 (1918).
- ⁵⁸ Mailhe and de Godon, Bull. Soc. Chim. (4), 23, 18 (1918).

The addition of 1 part of *iodins* to 240 parts of benzene warmed with 584 parts sulphuric acid brings about complete sulphonation in 5 hours; the iodine is readily recovered.⁵⁹ ⁶⁰

816. The catalyst most commonly employed is mercuric sulphate.

Benzoic acid heated with sulphuric acid alone gives only the meta
and para derivatives, but in the presence of mercuric sulphate, the
ortho is obtained.⁶¹

Anthraquinons gives only the β -monosulphonic acid with sulphuric acid alone, or the 2,6 and 2,7 disulphonic acids, when fuming sulphuric is used.

By heating to 130° with 0.5 part mercury to 110 parts sulphuric acid and 29 parts sulphur trioxide, the α -monosulphonic acid is obtained. At 160° with 1 part mercury to 200 parts sulphuric acid and 40 parts of the trioxide the *disulphonic* acids (1, 5), (1, 6), (1, 7), and (1, 8) are obtained. **at is a constant.

Vanadium sulphate has been proposed for aiding the sulphonation of pyridine.44

§ 19. — CONDENSATIONS BY ELIMINATING MOLECULES OF ALCOHOLS

817. It is convenient to consider reactions in which molecules of aliphatic alcohols are eliminated along with those in which water is abstracted. Anhydrous aluminum chloride is specially suitable as a catalyst for such condensations.

Ethyl ether reacts with benzene in the presence of aluminum chloride to form ethyl-benzene with the elimination of alcohol:66

$$C_6H_6 + (C_2H_5)_2O = C_2H_5.OH + C_6H_5.C_2H_6.$$

- ** Heinemann, English patent, 12,260 of 1915. J. Soc. Chem. Ind., 35, 1008 (1916).
- ⁶⁰ According to RAY and DEY (J. Chem. Soc. 117, 1405 (1920)) iodine influences the sulphonation of many compounds notably that of benzoic acid, the sulphonic acid group taking the ortho position under the influence of iodine instead of meta and para: toluene, chlor- and brombenzenes are sulphonated in para position only instead of ortho and para. E. E. R.
 - ⁶¹ DIMMROTH and VON SCHMAEDEL, Berichte, 40, 2411 (1907).
 - 62 ILJINSKY, Berichte, 36, 4194 (1914).
- es In the absence of mercury salts, a very small proportion, about 3% of α sulphonic acid is formed along with the beta. The presence of the mercury salt does not seem to affect the rate of sulphonation in the beta position but increases the rate of sulphonation in the alpha position so enormously that the operation can be carried on at much lower temperatures and with weaker oleum under which conditions the formation of the β -sulphonic acid is slow. E. E. R.
 - H FARBW. V. F. BAYER & Co., German patent, 160,104.
 - 'Jannasch and Bartels, Berichte, 31, 1716 (1898).

818. Under the same conditions benzene reacts with the chlormethyl ethers by the elimination of alcohol to form benzyl chloride along with some of the ether C₆H₆.CH₃.O.R(889). We have:⁶⁶

$$C_6H_6 + ClCH_2.O.R = ROH + C_6H_5.CH_2Cl.$$

819. Ethyl nitrate with benzene and aluminum chloride, gives a vigorous reaction which leads to nitrobenzene and the separation of alcohol:

$$C_6H_6 + O_2N.O.C_2H_5 = C_6H_5.NO_2 + C_2H_5.OH.$$

- 66 SOMMELET, Compt. rend., 157, 1443 (1913).
- 47 BOEDTKER, Bull. Soc. Chim. (4), 3, 726 (1908).

CHAPTER XVIII

DECOMPOSITION OF ACIDS

820. The aliphatic acids are very stable under the action of heat, except formic, which is decomposed by heat under many conditions. We will take up separately the catalytic decomposition of formic acid and then the decomposition of other aliphatic and aromatic acids under the influence of metals and of oxides. The action of the oxides leads to important applications which will be studied in succession, the preparation of symmetrical ketones, of mixed ketones and of aldehydes.

DECOMPOSITION OF FORMIC ACID

821. The decomposition of formic acid by heat may go in several distinct directions, either by the separation of carbon dioxide:

$$H.CO_2H = CO_2 + H_2 \tag{1}$$

or by the elimination of water:

$$H.CO_2H = CO + H_3O \tag{2}$$

or by the simultaneous elimination of water and carbon dioxide from two molecules:

$$2H.CO_2H = H.CO.H + CO_2 + H_2O.$$
(3)

If reactions (1) and (3) coexist, the nascent hydrogen from (1) may sometimes act on the *formaldehyds* produced in (3) to transform it into *methyl alcohol*. We will then have:

$$3H.CO_{2}H = CH_{2}.OH + 2CO_{2} + H_{2}O.$$
 (4)

The presence of a given catalyst will have the effect of turning the decomposition either into one of these directions or into several at the same time, by lowering more or less the temperature of the decomposition.

¹ Berthelot, Ann. Chim. Phys. (4), 18, 42 (1869). — Sainte-Claire-Deville and Debray, Compt. rend., 78, 1782 (1874). — Blackadder, Zeit. phys. Chem., 81, 385 (1912).

822. Reaction (1) which is a dehydrogenation, is produced at the ordinary temperature by rhodium black, or by palladium black.

Reaction (2) which is a dehydration is brought about by substances that take up water, sulphuric acid which acts below 100°, dry oxalic acid above 105° or anhydrous sodium and potassium formates above 150°.

823. Sabatier and Mailhe have studied the decomposition of formic acid under the influence of various catalysts, including finely divided metals, anhydrous oxides and some other substances.⁴ Comparisons have been made under analogous experimental conditions, the addition of the formic acid being at about 0.27 g. per minute and the pulverulent solid catalyst forming a layer 50 cm. long in a horizontal Jena glass tube heated to a known temperature.

The tube without catalyst gave a negligible decomposition below 300°, while at 340° 2.6 cc. of gas was collected per minute, chiefly a mixture of hydrogen and carbon dioxide (reaction 1) with a few percent of carbon monoxide (reaction 2).

824. The catalysts studied can be divided into three groups:

1st. Dehydrogenating Catalysts. These are the ones that cause reaction (1) almost exclusively, doubtless because they give rise to a temporary compound with one of the products, either hydrogen or carbon dioxide. The metals doubtless combine with the hydrogen:

Palladium (sponge) acts at 110° and produces total decomposition at 245°.

Platinum (sponge) begins to decompose it at 120°, the reaction being complete at 215°.

Reduced copper (light violet) evolves at 190° 278 cc. of gas containing equal amounts of hydrogen and carbon dioxide.

Reduced nickel at 280° disengages 290 cc. gas containing only traces of carbon monoxide.

Finely divided *cadmium*, prepared by reducing the oxide, gives 325 cc. gas per minute at 280°.

Stannous oxide begins to act above 150°, while at 285° it evolves 172 cc. of gas, being slowly reduced to small globules of tin which continue the catalysis. The gas contains a small excess of carbon dioxide due to reaction (3) which takes place to a slight extent.

An analogous result is produced by zinc oxids where the temporary production of zinc carbonate is doubtless the cause of the reaction: it begins to act at about 190° and at 230° disengages 172 cc. of gas containing 51% of carbon dioxide and 49% of hydrogen by volume.

- ² Zelinsky and Glinka, Berichte, 44, 2305 (1911).
- ² LORIN, Compt. rend., 82, 750 and Bull Soc. Chim. (2), 25, 517 (1876).
- 4 SABATIER and MAILHE, Compt. rend., 152, 1212 (1911).

The production of *formaldehyde* according to equation (3) amounts to 2 %. At 245° this may be raised to 12 % of formaldehyde.

825. 2nd. Dehydrating Catalysts. Reaction (2) takes place exclusively with *titania* above 170°, and at 320° 180 cc. of practically pure carbon monoxide is collected per minute.

The blue oxide of tungsten (715) acts the same way: at 270° it gives 195 cc. carbon monoxide practically pure.

The reaction goes in the same direction, but with reaction (3) as a side reaction to a slight extent, that is formaldehyde is produced equivalent to the carbon dioxide without hydrogen, with alumina, silica, zirconia, and uranous oxide, UO₂.

With alumina; the decomposition, which begins at about 234°, yields carbon monoxide containing 6% of the dioxide. Reaction (2) dominates but about 10% is decomposed according to (3) giving formaldehyde.

At 340° the disengagement of gas reaches 192 cc. per minute, but the gas then contains a little hydrogen resulting from the partial decomposition of the formaldehyde.

Silica, which is less active than alumina, gives about 3% of reaction (3).

At 340°, zirconia gives 144 cc. gas containing 5% of carbon dioxide: reaction (3) takes place to an extent of 10%.

With uranous oxide, reaction (3) is almost as important as (2).

826. 3rd. Mixed Catalysts. These are the most numerous of all. They produce reactions (1) and (2) simultaneously, usually with (3) as a minor side reaction.

This is what takes place with thoria. The decomposition shown by a slight evolution of gas, begins around 230. It is still quite slow at 250°, and gives a gas which contains 75% carbon monoxide, 15% carbon dioxide, and 10% hydrogen; the condensed liquid contains formal-dehyds. These figures show that of 100 molecules of formic acid, 79 undergo reaction (2), the other 21 being equally divided between (1) and (3).

Elevation of temperature modifies the conditions of the decomposition which is more and more rapid. At 320° the gas amounts to 120 cc. per minute and the carbon dioxide reaches 45% and the liquid contains considerable *methyl alcohol*, resulting from the intervention of reaction (4) which may be regarded as a reduction of formic acid by formaldehyde:

$$H.CO_2H + H.CO.H = CO_2 + CH_3.OH.$$

⁵ SABATIER and MAILHE, Compt. rend., 152, 1212 (1911).

HOFMANN and Schibsted, Berichte, 51, 1398 (1918).

The amount of *methyl alcohol* increases above 350° and as formaldehyde is then partially decomposed into carbon monoxide and hydrogen, the proportion of hydrogen increases while that of carbon dioxide decreases. At 375°, the gas is 144 cc. per minute containing only 33 % carbon dioxide. The condensate contains methyl alcohol.

827. For certain mixed catalysts, reaction (1) predominates; this is the case with the blue oxide of molybdenum, Mo₂O₂, resulting from the reduction of molybdic oxide by the formic acid at 340°. The decomposition, already clean at 105°, gives at 340°, 325 cc. of gas containing only 5% carbon monoxide. Of 12 molecules of the acid, 9 decompose according to reaction (1), 2 according to (3) and 1 according to (2).

Ferrous oxide, an active catalyst, and lime and broken Jena glass, mediocre catalysts, favor reaction (1) decidedly.

828. The two reactions (1) and (2) are of about equal importance with powdered white glass which acts at 240°.

The dehydration reaction (2) predominates as is indicated by the proportion of carbon dioxide being less than 33%, with a large number of substances whose absolute activities differ greatly, thus:

Powdered pumice liberates at 340° 4 cc. per minute
Magnesia 10
Charcoal from light wood 95
Light chromic oxide 150
Black vanadium oxide 215
Manganous oxide 225
Beryllium oxide 250
Reaction (3) takes place more or less with all of these.

DECOMPOSITION OF MONOBASIC ORGANIC ACIDS

829. In the action of heat on monobasic organic acids we find the three types of reactions given above for formic acid (821), namely: 1st. Elimination of carbon dioxide:

$$R.CO.OH = CO_2 + RH$$
hydrogerbon (1)

2nd. Separation of water alone, which can take place with primary or secondary acids only:

$$RR'.CH.CO.OH = H_2O + \frac{RR'C:CO}{ketene}$$
 (2)

3rd. Simultaneous elimination of water and carbon dioxide from two molecules of acid, giving a symmetrical ketone:

$$2R.CO.OH = CO_2 + H_2O + \underbrace{R.CO.R}_{\text{botons}}$$
 (3)

Reaction (2) is realized only exceptionally, as in the case of the action of an incandescent platinum spiral on the vapors, not of acstic acid but of acstanhydrids, giving the kstene, CH:CO, because the ketenes that are formed are very unstable and tend to polymerize ending up with carbonaceous substances.

Reactions (1) and (3) are of great importance.

830. Without the aid of a catalyst these two reactions take place simultaneously at a dull red heat: but the hydrocarbon and even the ketone are more or less destroyed and a complex pyrogenetic mixture results. The presence of a catalyst, either a finely divided metal or an oxide, orients the reaction and lowers the reaction temperature.

With aliphatic acids, reaction (1) is the most difficult to effect and is obtained only with difficulty by the use of finely divided metals. On the contrary, reaction (3) is easily brought about by the aid of oxide catalysts and leads to a practical method for the preparation of symmetrical ketones.

Aromatic and cyclic acids frequently give reaction (1) under the action of heat alone so that the aid of catalysts is often superfluous. However, the presence of suitably chosen catalysts can either accelerate reaction (1) or substitute for it, partially or entirely, reaction (3) which would not take place in their absence. But among the aromatic acids it is necessary to distinguish between those in which the carboxyl is joined directly to the nucleus and those in which the carboxyl is in a side chain. For the latter, e.g. phenyl-acetic acid, CeH₅. = CH₂.COOH, reaction (3) is easily realized by the aid of catalytic oxides as is the case with aliphatic acids.

For the former, s.g. benzoic, C₆H₃.COOH, and the toluic acids, reaction (1) is the one that always tends to take place, reaction (3) being very difficult to obtain, at least with the aromatic acids alone.

SIMPLE ELIMINATION OF CARBON DIOXIDE

831. In the case of aliphatic acids this is accomplished more or less by finely divided metals.

Finely divided copper commences to decompose the vapors of acetic acid at 260°, and an evolution of gas is obtained, slow at first but quite regular at 390-410°, containing 7 volumes of carbon dioxide to 1 of methane. The formation of some acetone is observed. Reactions (1) and (3) are catalyzed and the composition of the gas shows that of 13

¹ Wilsmore, J. Chem. Soc., 91, 1938 (1901).

molecules of the acid, 1 has decomposed according to reaction (1) and 12 have given acetone.

832. Reduced nickel causes, slowly below 240°, rapidly above 320°, an analogous decomposition. The gas contains 50% of methane and reaction (1) seems to have taken place exclusively, but a portion of the acids is decomposed into carbonaceous substances which are deposited on the metal.

833. Other aliphatic acids give analogous results. The action of copper is slow. That of nickel is much more rapid: at 230°, propionic acid is broken down into carbon dioxide and ethane which is largely decomposed into methane, carbon and hydrogen. No ketone is formed, but a part of the acid is reduced to the aldshyds. At 250°, butyric acid gives analogous results, so do isobutyric and caproic.

834. With aromatic acids the decomposition into carbon dioxide and hydrocarbon is usually quite easy.

The vapors of benzoic acid carried along by carbon dioxide over reduced copper at 550°, are totally decomposed into benzene and carbon dioxide.

Over nickel, or over the oxide which is rapidly reduced at that temperature, the benzene produced is almost entirely broken up with the deposition of carbon and the liberation of hydrogen and *methane*. Under the same conditions, reduced *iron* gives *benzene*, which is partially destroyed, and some *diphenyl*.¹⁰

835. In contact with copper powder, coumanic acid is regularly transformed into γ -pyrone:¹¹

836. The presence of alkaloids favors the decomposition of the carboxy-camphor acids at 70° into camphor and carbon dioxide. With an inactive alkaloid, in polarized light, the dextro and laevo acids are decomposed at the same rate; with an active alkaloid, the velocities are different. Thus with quinine a difference of 46% is found.¹²

The conditions of this decomposition in the presence of quinoline.

- SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 467 (1905).
- MAILHE, Bull. Soc. Chim. (4), 5, 616 (1909).
- 10 SABATIER and MAILHE, Compt. rend., 159, 217 (1914).
- 11 WILLSTÄTTER and PUMMERER, Berichte, 38, 1461 (1908).
- ¹² FAJANB, Zeit. physik. Chem., 73, 25 (1910).

pyridine, piperidine, and of other amines, benzyl-amine, allyl-, iso-amyl-amine, etc., have been studied in detail, in various solvents at 80° with the result that the formation of a complex by the acid and the amine appears to be the cause of the catalysis in every case. 18

SIMULTANEOUS ELIMINATION OF WATER AND CARBON DIOXIDE

I. Preparation of Symmetrical Ketones

837. This is the reaction that is specially catalyzed by metallic oxides.

It is derived in fact from the old method of preparing symmetrical ketones by calcining at a red heat the calcium or barium salts of monobasic organic acids:

$$(R.CO_2)_2Ba = R.CO.R + BaCO_2.$$

Squibb conceived the idea of transforming this reaction into a catalytic one. By passing the vapors of acstic acid over barium carbonate heated to about 500°, a regular and continuous decomposition of the acetic acids into acetone, water and carbon dioxide is obtained:

$$2CH_{2} \cdot CO_{2}H = CH_{2} \cdot CO \cdot CH_{2} + CO_{2} + H_{2}O.$$

The process which gives a yield of better than 90% has been used industrially. The carbonates of all the metals whose acetates give acetone on calcination may be used.¹⁴ We have studied above (161) the mechanism of this reaction.

- 838. Ipatief described an analogous formation when he used zinc oxide or carbonate or the carbonates of calcium, barium and strontium. Acetic acid gives acetone, and propionic acid, disthyl-ketone.¹⁵
- 839. Calcium Carbonate. This is an excellent catalyst for acetic acid, a short column at 450° is sufficient to transform the acid completely into practically pure acetone with the evolution of nothing but carbon dioxide and water.

With propionic acid, the yield of diethyl-ketone is very satisfactory, some propionic aldshyds is obtained and the gas contains a little ethylene. This formation of the aldehyde increases with the complexity of the molecule and appears to be correlative to the production of the unsaturated hydrocarbon. We have:

$$C_nH_{3n+1}CO.OH = C_nH_{3n} + H.CO.OH.$$
hydrocarbon formic acid

- Bredig and Joyner, Zeit. f. Elektrochem., 24, 285 (1918).
- ¹⁴ SQUIBB, J. Amer. Chem. Soc., 17, 187 (1895), and 18, 231 (1896). CONBOY, Rev. gén. Sci., 13, 563 (1902).
 - ¹⁸ IPATIEF and SCHULMANN, J. Russian Phys. Chem. Soc., 36, 764 (1904).

The formic acid thus produced can break up at once into $CO_2 + H_2$ or into $CO + H_2O$ (821), but it can act also on the acid that is being used reducing it to the *aldshyds* (851).

The secondary reactions, still more important for butyric acid, are exaggerated with isobutyric and isoraleric acids.

The calcium carbonate used is blackened by the decomposition of a small portion of the acid, but nevertheless conserves its catalytic activity almost indefinitely and remains as carbonate for the most part.¹⁶

Benzoic acid is scarcely attacked till about 550°, and gives chiefly benzene and carbon dioxide with only a little benzophenone and traces of anthraquinone.¹⁷

The same difficulty is encountered with the typical aromatic acids in which the carboxyl is united directly to the nucleus, such as ortho, meta, and paratoluic acids and the naphthoic acids.

On the contrary, aromatic acids in which the carboxyl is in a side chain, such as phenyl-acetic, and phenyl-propionic acids can be advantageously transformed into the corresponding symmetrical ketones at 430-70°.

840. Among the metallic oxides the most suitable for the production of ketones are *thoria* and *manganous oxide*. It is sufficient to pass the vapors of the acid over a layer of the oxide, usually below 450°.

Alumina. Alumina gives very good results with acetic, not quite so good with propionic and quite poor with isobutyric. With benzoic only a slow decomposition into benzene is effected.

Chromic Oxide. The results are analogous to those obtained with alumina.

Uranium and Zirconium Oxides. These give nearly the same results but their activity diminishes quite rapidly.

Lims. Lime acts as the carbonate. When it is used at 400°, it is possible to observe the formation of the intermediate salt, the decomposition of which furnishes the ketone and which is decomposed only above 420° for the acstats and 460° for the butyrats. The temperatures reached can account for some decomposition of the ketones formed.

841. Zinc Oxide. With zinc oxide, the acstate is decomposed above 280° and very rapidly at 340°; the production of acetone is therefore very easy. The difficulty of forming the ketone increases with the molecular weight of the acid and is partly due to the volatility of the zinc salt. Benzoic acid is not attacked below 500° and then gives only benzene.

¹⁶ Sabatier and Mailes, Bull. Soc. Chim. (4), 13, 319 (1913) and Compt. rend., 156, 1730 (1913).

¹⁷ SABATIER and MAILHE, Compt. rend., 159, 217 (1914).

¹⁸ SENDERENS, Bull. Soc. Chim. (4), 3, 824 (1908).

842. Cadmium Oxide. This is slowly reduced by the acid vapors but without the activity being diminished by the formation of the metal which can be seen sublimed in the tube. At 400–450° it can readily transform acetic, propionic, butyric and valeric acids into their symmetrical ketones; the results are not so good with branched chain acids as isobutyric and isovaleric, with which the gas evolved is no longer pure carbon dioxide but contains considerable amounts of the unsaturated hydrocarbons, carbon monoxide and hydrogen.¹⁹

It acts at 450°, and violently at 500°, on the vapors of *benzoic* acid to give benzene, the reduction of the oxide hardly modifying the catalysis.²⁰

843. Oxides of Iron. Ferrous oxide resulting from the calcination of the oxalate as well as ferric oxide which is rapidly reduced to the ferrous, can give good results with aliphatic acids at 450-90°. The yield of ketone is excellent with acetic or propionic, good with caprylic and poor with isobutyric or isovaleric acid.²¹

The immediate formation of a ferrous compound is the basis of a process for the preparation of ketones by heating an acid with 10% of its weight of iron scale: this works well for the higher fatty acids from lauric up to melissic. Thus stearic acid gives 80% of the ketone. The results are not so good with oleic, elaidic, and brassidic and are poor with the lower acids, acetic, butyric, etc., as well as with phenylacetic, benzoic, suberic, and sebacic.²²

With benzoic acid at 550°, ferric oxide acts like iron (834), but its simultaneous reduction causes the formation of a certain amount of phenol which results from the oxidation of the benzene formed.²²

844. Thoria. Thoria of which the valuable qualities of constant activity and ready revivification have been mentioned in connection with the dehydration of alcohols (708), gives excellent results with various monobasic aliphatic acids and enables us to prepare with good yields, acstone, disthyl-ketone, dipropyl-ketone, disophotyl-ketone, distophotyl-ketone, dibutyl-ketone, etc., as well as ketones derived from aromatic acids in which the carboxyl is not joined immediately to the nucleus, such as phenyl-acetic, β -phenyl-propionic, etc.²⁴

Benzoic acid is only slowly attacked by thoria even at 550° and then is only decomposed into benzene and carbon dioxide.²⁵

- 19 MAILHE, Bull. Soc. Chim. (4), 13, 666 (1913).
- ²⁰ Sabatier and Mailhe, Compt. rend., 159, 217 (1914).
- ²¹ Mailhe, Compt. rend., 157, 219 (1913).
- * Easterfield and Taylor, J. Chem. Soc., 99, 2298 (1911).
- ²² Sabatier and Mailhe, Compt. rend., 159, 217 (1914).
- M SENDERENS, Ann. Chim. Phys. (8), 18, 243 (1913).
- * SABATIER and MAILEE, Compt. rend., 159, 217 (1914).

845. Manganous Oxide. This oxide prepared by calcining the precipitated carbonate in the vapor of methyl alcohol, is on account of its low price and its great activity, very useful for the preparation of ketones at 400–450°. The carbonaceous deposits do little harm and the same lot of oxide has been used in 22 different preparations. In the case of slightly volatile acids, carbon dioxide is used to carry their vapors along.

The yields of symmetrical ketones are very high, not only for acetic, propionic and valeric acids, but also for isobutyric, with which an experiment carried out at 400-410° gave a 70% yield, with caproic, heptoic, nonylic as well as with phenyl-acetic. With benzoic acid at 550° a little benzophenone is formed, but chiefly benzone.²⁶

- 846. Lithium Carbonate. At 550° this is the most advantageous catalyst for transforming *benzoic* acid into *benzophenone*, always accompanied by a little anthraquinone; but even in this, the most favorable case, much benzene is formed.²⁷
- 847. Formation of Ketones in Liquid Medium. In the case of monobasic acids which boil above 300°, the ketones can be formed by heating the acids somewhat above 300° with various catalysts, including the oxides mentioned above, silica, silicates and also finely divided metals. Stearic acid yields stearone in 3 hours.²⁷

II. Preparation of Mixed Ketones

848. A long time ago Williamson showed that the calcination of a mixture of the calcium salts of two fatty acids gave the mixed ketone: 28

$$(R.CO_2)_2Ca + (R'.CO_2)_2Ca = 2CaCO_2 + 2R.CO.R'.$$

It might be expected that the catalytic decomposition by means of oxides when applied, not to a single acid but to a mixture of two acids, would give the mixed ketone derived from the two acids instead of the symmetrical ketone. Senderens found this to be the case. We have:

$$R.CO_2H + R'.CO_2H = CO_2 + H_2O + R.CO.R'.$$

A simple method of preparing mixed ketones is to pass a mixture of the vapors of the two acids over thoria at about 400°.

For the success of this method it is sufficient that one of the acids is catalyzed by thoria: we may use two aliphatic acids or one aliphatic with benzoic or a toluic acid, but not benzoic with a toluic.

- ³⁶ Sabatier and Mailhe, Compt. rend., 158, 830 (1914).
- ²⁷ Schicht Act. Ges. and Grün, German patents, 295,657 and 296,677. J. Soc. Chem. Ind., 36, 569 and 615 (1917).
 - 28 WILLIAMSON, Annalen, 81, 86 (1852).

The chief reaction is usually that which furnishes the mixed ketone, but it is always accompanied by the reactions that the two acids would undergo separately. We obtain three ketones if we start with two aliphatic acids or an aliphatic and phenyl acetic, but only two when an aliphatic acid is used with benzoic, a toluic, or a naphthoic.

The separation of the ketones is easily accomplished by fractionation. Numerous mixed ketones have been prepared in this way.

849. The green oxide of uranium, though less active, can replace thoria for this preparation: zirconia acts in the same way, but does not give as good results with the homologs of benzoic acid. Lims, zinc oxide, alumina, and chromic oxide produce acstophenone easily but give poorer and poorer results as the aliphatic acid increases in molecular weight.

Titania, stannic oxide and ceria give decomposition products chiefly.²⁹

Cadmium oxide, ferrous and ferric oxides, 30 and calcium carbonate are excellent catalysts for mixed ketones. 31

850. Manganous oxide at 400-450° is as good as thoria and by its use mixed ketones have been prepared from benzoic acid with lauric, muristic, C₁₄H₂₀O₂, and stearic, C₁₅H₂₀O₂, as well as phenyl-acetic.²²

CATALYTIC PREPARATION OF ALDEHYDES

851. If in Williamson's method for preparing mixed ketones, one of the calcium salts is a formate, an aldehyde a is produced accompanied by the decomposition products of the individual salts, the symmetrical ketone, R.CO.R, formaldshyde, and methyl alcohol as well as gaseous products from the formate:

$$(R.CO_2)_2Ca + (H.CO_2)_2Ca = 2CaCO_3 + \underbrace{2R.CO.H}_{aldehyde}$$

Analogies would lead us to expect that a mixture of the vapors of formic acid and another monobasic organic acid passed over an oxide catalyst would give the aldehyde corresponding to that acid according to the reaction:

$$R.CO_2H + H.CO_2H = R.CO.H + CO_2 + H_2O.$$

- 29 SENDERENS, Loc. cit.
- ²⁰ MAILHB, Compt. rend., 157, 219 (1913).
- ²¹ Sabatier and Mailhe, Compt. rend., 156, 1732 (1913).
- SABATIER and MAILHE, Compt. rend., 158, 830 (1914).
- ²⁵ Limpricht, Annalon, 97, 368 (1856). Piria, Ann. Chim. Phys. (3), 48, 113 (1856).

852. Sabatier and Mailhe were able to realize this reaction with titania as a catalyst. This constitutes a general method for the preparation of aldehydes from acids. It is sufficient to pass the vapors of the acid mixed with an excess for formic acid over titania heated to 300°. There is evolved a mixture of carbon monoxide resulting from the decomposition of the formic acid by the titania (825) and carbon dioxide from the desired reaction. The condensate is a mixture of water, aldehyde, and unchanged acids from which the aldehyde is easily separated.

Aldehydes derived from various aliphatic acids up to C₂ have been thus prepared with yields above 40% and reaching 90%.

Thus nonylic, or pelargonic acid, gives 85% of nonylic aldehyde. Usually no ketone is formed; only very small amounts of the ketones are formed with acids containing more than 5 carbon atoms.

The unsaturated acid, *crotonic*, is likewise transformed into the aldehyde. The reaction gives poor results with *benzoic acid* but works well with phenyl-acetic, the constitution of which is more like the aliphatic acids. It gives a 70% yield.⁸⁴

853. Manganous oxide can be substituted for titania and has the advantages of being readily prepared and of retaining its activity because it gives rise to less tarry deposits. The operation should be conducted at a little higher temperature, 300–350°. The yields are very satisfactory, reaching 50% with isovaleric acid. Caproic, heptoic, octoic, and nonylic aldehydes have been prepared in this way.²⁵

854. The use of thoria is less advantageous because it requires a higher temperature and because it favors the formation of ketones which are found with the aldehydes; however, by operating at 270–300°, 25 to 30%, and sometimes more, of the aldehydes are obtained.³⁴

DECOMPOSITION OF DIBASIC ACIDS

855. Solid oxalic acid, HOOC.COOH, mixed with alumina is decomposed below 100° into water, carbon monoxide and dioxide. 36

Glycerine mixed with crystallized oxalic acid produces a different result: at 100-110° carbon dioxide and formic acid are produced:

$$HOOC.COOH = CO_2 + H.CO_2H.$$

When the reaction dies down all that is necessary to start it again is to add some more oxalic acid and so on, the glycerine being able to

²⁴ Sabatier and Mailhe, Compt. rend., 154, 561 (1912).

^{*} Sabatier and Mailhe, Compt. rend., 158, 985 (1914).

³⁶ SENDERENS, Bull. Soc. Chim. (4), 3, 828 (1908).

serve almost indefinitely, and hence playing the part of a catalyst. In reality there is first the production of a glycerine mono-oxalate:

$$HOH_2C.CH(OH).CH_2OH + HOOC.COOH = H_2O + HOH_2C.CH(OH).CH_2.CO_2.COOH.$$

At 100-110° this ester loses water and gives the monoformine, HOH₂C.CH(OH).CH₂.CO₂H, which is saponified by the water set free in the first reaction liberating formic acid and glycerine which is thus free to recommence the process.

856. The use of manganous oxide permits the preparation of cyclopentanones from ϵ -dibasic fatty acids.

The vapors of adipic acid carried along by a current of carbon dioxide over manganous oxide at 350°, give an 85% yield of cyclopentanone:

$$CH_{2}.CH_{2}.CO_{2}H$$
 $CH_{3}.CH_{2}.CO_{2}H$
 $= CO_{2} + H_{2}O + CH_{3}.CH_{3}$
 $CO.$

Likewise β -methyl-cyclopentanone is prepared from β -methyl-adipic acid. 27

But with suberic acid, in which the carboxyl groups are separated by 6 carbon atoms, the process gives only a poor yield of suberone and tarry substances are formed which gum up the catalyst.²⁸

CATALYTIC DECOMPOSITION OF ACID ANHYDRIDES

857. The anhydrides like the acids can be decomposed catalytically to form the corresponding symmetrical ketones, carbon dioxide only being eliminated:

$$R.CO.O.CO.R = CO_2 + R.CO.R$$

Precipitated calcium carbonate gives good results at 450-500° with the anhydrides of acetic, propionic, isovaleric, etc., acids. Thoria is also suitable for this reaction.

This process gives a mixed ketone along with the two symmetrical ketones when an acid and the anhydride of another acid are used together.³⁹

- 87 SABATIER and MAILHE, Compt. rend., 158, 985 (1914).
- ⁸⁸ GODCHOT and TABOURY, Bull. Soc. Chim. (4), 25, 352 (1919).
- ⁸⁹ Sabatter and Mailee, Bull. Soc. Chim. (4), 13, 320 (1913), and Compt. rend., 156, 1733 (1913).

CHAPTER XIX

DECOMPOSITION OF ESTERS OF ORGANIC ACIDS

& I. - ESTERS OF MONOBASIC ACIDS

858. In the absence of catalysts the esters of monobasic acids are difficult to decompose by simply heating; the decomposition is slow and such high temperatures are required that the molecules are broken up. We may mention that ethyl benzoate heated in a sealed tube above 300° is slowly decomposed into bensoic acid and ethylene. Colson, who noted this reaction and a similar decomposition of ethyl stearate, considered this tendency to decompose into the acid and an unsaturated hydrocarbon a general property of esters.

The presence of a catalyst acting at the same time on the alcohols and on the acids should greatly facilitate the decomposition of esters which should yield, in conformity with what has been said above, the unsaturated hydrocarbon and the decomposition products of the acid, water, carbon dioxide, and the symmetrical ketone. Some observations relative to the action of alumina on ethyl acetate, propionate and butyrate confirmed this prediction, but on the contrary, these same esters gave with thoria a complicated decomposition which has not been cleared up.²

Sabatier and Mailhe have studied a great number of cases of the action of various catalytic oxides on esters of various sorts and have indicated the general conditions that govern the decomposition.³

Formic esters require separate treatment and will be taken up after the other esters.

859. If an ester derived from a primary aliphatic alcohol and from a monobasic organic acid, other than formic, be brought in contact with a catalytic oxide, MO, derived from an amphoteric hydroxide, M(OH)₃, the following reaction will take place:

$$\frac{2R.CO.OC_{n}H_{2n+1}}{} + 2MO = (R.COO)_{2}M + (C_{n}H_{n2+1}O)_{2}M.$$

The salt, $(RCO.O)_2M$, and the alcohol derivative, $(C_nH_{2n+1}O)_2M$ are both unstable, if the oxide chosen is at the same time a catalyst

- ¹ Colson, Compt. rend., 147, 1054 (1908).
- ² SENDERENS, Bull. Soc. Chim. (4), 5, 482 (1909).
- ² Sabaties and Mailes, Compt. rend., 152, 669 (1912), and 154, 49 and 175 (1912).

for the decomposition of alcohols and of acids at the operating temperature.

860. First Case. If the instability of the two temporary compounds is of the same order they will decompose simultaneously and the decomposition becomes:

(1) $2R.CO.OC_nH_{2n+1} + 2MO = \left\{\frac{R.CO.R + CO_2}{\text{ketone}}\right\} + 2MO.$

A symmetrical ketone is produced and an unsaturated hydrocarbon which, if it is a gas (ethylene, propylene and butylene) has twice the volume of the carbon dioxide produced. This was the case in the experiments with alumina mentioned above (858).

If the ester is a *methyl* ester there is no separation of water, and methyl ether, (CH₂)₂O. is formed.

861. Second Case. If the catalyst is more active toward acids than with alcohols, the decomposition of the complex, (R.CO.O)₂M, is more rapid than that of the alcohol compound. The water formed in reaction (1) has time to react with an equivalent amount of the latter and decomposes it to regenerate the alcohol:

$$(C_nH_{2n+1}O)_2 + H_2O = MO + \underline{2C_nH_{2n+1}.OH}$$

This combined with the former reaction gives:

(2)
$$4R.CO.OC_nH_{2n+1} = 2R.CO.R + 2CO_2 + 2C_nH_{2n} + 2C_nH_{2n+1}.OH.$$

There is the simultaneous formation of ketone and alcohol and of equal volumes of carbon dioxide and of unsaturated hydrocarbon (if it is a gas). This is usually the case with decompositions caused by thoria, e.g. at 310° with ethyl acetate, propyl acetate, propyl propionate, isobutul acetate, and ethyl caproate.

- ⁴ Titania prepared by the precipitation of the hydroxide from the sulphate catalyses the decomposition of ethyl acetate two thirds according to the equation:
- (1) CH₂CO₂C₂H₄ = C₂H₄ + CH₄COOH and one third according to:
 - (2) $2CH_1CO_2C_2H_3 = CH_1COCH_4 + CO_2 + C_2H_4 + H_2O$

Titania prepared by precipitating blue titanous hydroxide from a solution of titanous chloride, and then allowing this to oxidise to the white titanic hydroxide while suspended in the solution, catalysed the reaction one third according to (1) and two thirds according to (2).

Thoria prepared by ignition of the nitrate gives very little ethylene, as was found by Sabatier, but thoria prepared by precipitation of the hydroxide gives almost as much ethylene as would be called for by (2).

Alumina does not only give reaction (2) but a fifth to two thirds of the ethyl acetate is decomposed according to (1). The method of preparation of the catalyst and the length of time it has been used determine the proportions.—Homer Adkins.

862. Elevation of temperature accelerates the decomposition of the unstable intermediates and tends to bring the reaction nearer to (1). This is the case with *isobutyl acetate* over thoria at above 350° and for *ethyl caproate* at about 360°. Besides when the temperature becomes high, the alcohols suffer more or less decomposition into hydrogen and *aldehydes*, easy to recognize, and these may be partially split up into hydrocarbons and carbon monoxide.

863. Third Case. If the catalyst is less active with acids than with alcohols, the temporary complex, (R.CO.O)₂M, will be decomposed only slowly. The water set free by the rapid decomposition of the alcohol complex will act on the above to set the acid free:

$$(R.CO.O)_2M + H_2O = MO + \underbrace{2R.CO.OH}_{\text{acid}}$$

In this case the formation of ketone and liberation of carbon dioxide are less important: the production of unsaturated hydrocarbon and setting free of acid predominate.

This is what takes place over *titania* with esters of acetic, propionic, butyric and valeric acids, which acids it decomposes more slowly than it does the alcohols.

864. Fourth Case. The exaggeration of the preceding case is found with those catalysts which are active with alcohols but are incapable of decomposing acids. This is the case with various catalytic oxides, e.g. thoria and titania with esters of benzoic and toluic acids, and with boric anhydride with esters of aliphatic acids, since boric acid can form the temporary complexes with the alcohols only. In such cases we may write the reaction as follows:

$$\begin{split} 2R.CO.OC_nH_{2n+1} + MO &= M(OC_nH_{2n+1})_2 + \underbrace{(R.CO)_3O}_{anhydride} \\ &= MO + 2C_nH_{2n} + H_2O + (R.CO)_3O \\ &= MO + 2C_nH_{2n} + \underbrace{2R.CO_3H}_{aoid}. \end{split}$$

There will be a total regeneration of the acid with the formation of the unsaturated hydrocarbon exclusively. It has been found that *ethyl benzoate* is decomposed into benzoic acid and ethylene by thoria above 400°, as in Colson's sealed tube.

Likewise ethyl valerate catalyzed by boric anhydride above 400°, gives ethylene and valeric acid exclusively.

865. Methyl esters which can give only methyl ether are difficult to decompose: the reaction, which requires a higher temperature, yields exclusively carbon dioxide, methyl ether and the ketone, frequently partially decomposed, and resulting water which may saponify a part of the ester to form free acid and methyl alcohol.

Catalytic Decomposition of Formic Esters

866. In the absence of catalysts, formic esters are quite stable: when the vapors of *ethyl formate* are passed through a glass tube at 400°, no appreciable decomposition is observed, but the decomposition is very rapid in contact with catalysts that decompose formic acid (821), and takes place at temperatures lower than those required for the esters of other aliphatic acids, but higher than those required by formic acid.

Sabatier and Mailhe have shown that this decomposition takes place according to two different reactions at the same time, the one similar to the usual decomposition of esters of other aliphatic acids:

(1)
$$2H.CO_2C_nH_{2n+1} = H.CO.H + CO_2 + \underbrace{(C_nH_{2n+1})_2O}_{\text{other}}$$

the ether surviving only in the case of methyl ether, splitting in other cases into water and unsaturated hydrocarbon $(H_2O + 2C_nH_{2n})$; the other always predominating, is peculiar to formic esters:

(2)
$$\mathbf{H}.CO_2C_n\mathbf{H}_{2n+1} = CO + \frac{C_n\mathbf{H}_{2n+1}.O\mathbf{H}.}{alcohol}$$

A portion of this alcohol is decomposed at the reaction temperature, either into aldehyde and hydrogen (with metals or manganous oxide), or into unsaturated hydrocarbon and water (with thoria and alumina), or in both ways (with mixed catalysts). The water resulting from reaction (1) or from the dehydration of the alcohol formed according to equation (2) can saponify a part of the ester to alcohol and free formic acid which is then decomposed in the way already described (821).

867. Metals. Finely divided metals can easily cause the decomposition of formic esters, nickel above 220°, platinum above 270°, and copper above 350°. Reaction (2) greatly predominates and gives the alcohol which the metal breaks down to aldehyde, along with the carbon monoxide. With copper or with nickel at a low temperature the aldehyde survives, but with platinum or with nickel at a high temperature (618) it is largely destroyed.

868. Titania. Reaction (2) takes place almost exclusively. From methyl formate, methyl alcohol and methyl ether resulting from its partial dehydration are obtained. The gas collected over water is practically pure carbon monoxide without the dioxide because the formic acid, under these conditions, gives only carbon monoxide and water (825).

SABATIER and MAILHE, Compt. rend., 154, 49 (1912).

869. Zinc Oxide. It is again reaction (2) that predominates, but the formic acid that is set free by the water resulting from the dehydration of the alcohol, is decomposed by the catalyst into water and carbon dioxide which is found mixed with the monoxide.

870. Thoria. Reaction (2) predominates but is accompanied by (1) which furnishes a certain amount of formaldehyde which is diminished as the reaction temperature is raised.

§ 2 — DECOMPOSITION OF ESTERS IN THE PRESENCE OF AMMONIA

871. When the vapors of an ester of an organic monobasic acid mixed with ammonia are passed over *thoria* or *alumina* at about 480-90°, *nitriles* are obtained by the elimination of water and alcohol or of decomposition products of the alcohol.

Methyl esters give the alcohol, partially split into methyl ether and water or into formaldehyde and hydrogen.

Esters of other aliphatic alcohols give the unsaturated hydrocarbons, while phenol esters yield phenol the major portion of which remains:

$$R.CO.OR' + NH_4 = H_4O + R'OH + RCN.$$

Ethyl acetate gives ethylene and acotonitrile and isoamyl acetate yields amylene and acetonitrile, while phenyl acetate liberates the same nitrile and phenol.

Analogous results have been obtained with esters of propionic, butyric, isovaleric, nonylic and caproic acids.

Methyl benzoate gives more than 80% of benzonitrile with methyl alcohol and formaldehyde. Ethyl and isopropyl benzoates yield the same nitrile. The esters of the three toluic acids behave in the same way.

Ethyl α - and β -naphthoates are almost quantitatively transformed into the α - and β -naphthonitriles.

Ethyl phenyl-acetate gives an excellent yield of benzyl cyanide.

§ 3.—ESTERS OF DIBASIC ACIDS

872. The catalytic decomposition of esters of dibasic acids has been as yet very incompletely studied.

Catalytic oxides such as alumina and thoria cause decompositions readily. If we extend to the esters of dibasic acids the interpretation above set forth for the action of these oxides, we can predict that an oxide, MO, will effect the reaction:

³ MAILHE, Bull. Soc. Chim. (4), 23, 232 (1918).

$$(CH_2)_x \underbrace{CO.OR}_{CO.OR} + 2MO = \underbrace{(CH_2)_x}_{CO.O} \underbrace{M + M}_{OR}$$

$$\underbrace{CO.OR}_{ester}$$
metal salt

If the oxide is at the same time a catalyst for acids and for alcohols, the compounds thus formed will be unstable and will decompose as follows:

$$(CH_3)_x \xrightarrow{CO.O} M = MO + (CH_3)_x \xrightarrow{CO} O$$
and:
$$M \xrightarrow{OR} = MO + R \xrightarrow{ether} O.$$

The oxide, MO, is entirely regenerated and can carry on the reaction indefinitely. We will obtain as results of the catalysis, the acid anhydride, or its debris, if it is unstable, the ether, or in most cases, the catalytic decomposition products of that ether, i.e. water and an unsaturated hydrocarbon.

873. These predictions have been verified by Sabatier and Mailhe in the case of the neutral esters of oxalic, malonic and succinic acids over thoria.

equally unstable and decomposes into carbon suboxide, CO:C:CO, which polymerizes into reddish products or decomposes into carbon monoxide, dioxide and carbon.

Succinic anhydride is stable if the temperature is not too high and can be collected as crystals melting at 177°. If the temperature is above 350°, it is decomposed into carbon monoxide and dioxide, ethylene and condensation products.

These results have been verified for the ethyl, propyl, isobutyl and isoamyl esters of the three acids: except in the case of the ethyl esters, where the stable ethyl ether can be collected, the debris of the esters, water and the unsaturated hydrocarbon, are found.

⁷ SABATIER and MAILHE, Bull. Soc. Chim. (4), 11, 369 (1912).

^{*} SABATIER and MAILHE, Loc. cit. and unpublished results.

With esters of oxalic acid, the catalytic decomposition begins at very moderate temperatures and is already rapid at 220°: higher temperatures are required for malonates and still higher for succinates.

A decomposition of this nature has been found in the particular case of *ethyl oxalate* over *alumina*: at 200°, ethyl ether, carbon monoxide and dioxide are obtained, while at 360°, the ethyl ether is replaced by ethylene.

874. The catalytic decomposition of ethyl succinate over alumina at 400°, according to Senderens, liberated ethylene and carbon dioxide and produced p. cyclohexadione.¹⁰ But Sabatier and Mailhe were not able to verify this and obtained only succinic anhydride along with ethylene and carbon dioxide. The same results were obtained with alumina at 260°.

Ethyl glutarate over alumina at 270°, gave only ethylene and glutaric acid. Ethyl adipate furnished ethylene and adipic acid at 300°. 11 At higher temperatures this should have given cyclopentanone (856).

[•] SENDERENS, Bull. Soc. Chim. (4), 3, 826 (1908).

¹⁰ SENDERENS, Bull. Soc. Chim. (4), 5, 485 (1909).

¹¹ Michiels, Bull. Soc. Chim. Belge, 27, 227 (1913); C. A., 8, 1106 (1914).

CHAPTER XX

ELIMINATION OF HYDROGEN HALIDES OR SIMILAR MOLECULES

875. The elimination of hydrogen halides can take place from a single molecule or by condensation from two molecules. Anhydrous chlorides are the chief catalysts in both cases.

§ 1. — ELIMINATION OF HYDROGEN HALIDE FROM A SINGLE MOLECULE

876. When an alkyl mono-chloride is passed over a layer of various anhydrous metal chlorides in a tube maintained at above 260° there is rapid decomposition into unsaturated hydrocarbon and hydrochloric acid:

$C_nH_{2n+1}Cl = HCl + C_nH_{2n}$

Methyl chloride alone does not decompose in this way.

Barium, nickel, cobalt, lead, cadmium and ferrous chlorides are suitable for effecting this reaction. Primary chlorides are decomposed above 260° and rapidly at 300° while secondary and tertiary are still more easily acted on.

The same metal chlorides decompose alkyl mono-bromides or monoiodides in the same way to form hydrobromic or hydroiodic acids, but higher temperatures are required.¹

The recombination of the unsaturated hydrocarbon with the liberated hydrogen halide takes place to a certain extent in the tube beyond the catalytic chloride and may yield a certain amount of secondary or tertiary isomers of the original alkyl halide.

Dry barium chloride gives very good results and can effect this decomposition indefinitely; if it is dissolved in water after long use, there is a small residue of viscous very condensed hydrocarbons with a petroleum odor.

The chlorides of monovalent metals, silver, sodium, and potassium are inactive.

The process applies to monochlor derivatives of cyclohexane and cyclopentane. It can succeed with unsaturated mono-chlorides and even

¹ SABATTER and MAILHE, Compt. rend., 141, 238 (1905).

with halogenated alcohols 2 as well as with dichlorcyclohexane which is converted to dihydrobenzene.3

877. Anhydrous aluminum chloride acts actively in the same manner, but it has the disadvantage of producing liquid products that hinder the continuation of the catalysis. It has been used to transform propyl chloride into propylene.

878. The catalysis can be explained by the assumption of an unstable organo-metallic combination derived from the alkyl chloride:

$$C_nH_{2+n+1}.Cl + BaCl_2 = HCl + Cl.Ba.C_nH_{2n}.Cl.$$

The mixed complex thus formed would decompose rapidly to give the unsaturated hydrocarbon:

$$Cl.Ba.C_nH_{2n}.Cl = BaCl_2 + C_2H_{2n}.$$

The regenerated chloride can repeat the cycle of reactions indefinitely. The formation of such a mixed complex can be observed in the case of anhydrous aluminum chloride: mixed with isobutyl chloride at -10° , no reaction takes place but if the mixture is warmed to 0° , hydrogen chloride and isobutylene are evolved and an intensely colored liquid is formed.

At 300°, ferric chloride causes the elimination of hydrogen chloride, but no isobutylene is formed; a solid of high molecular weight is produced.

Chromic chloride, CrCl, does not act.5

879. As has been mentioned above (876), this kind of catalysis can be applied to poly-halogen derivatives.

Heptachlorpropane, CCl₂.CCl₂.CHCl₂, is decomposed above 250° by cuprous chloride with the elimination of hydrogen chloride to give pentachlor-propylene, CCl₂.CCl: CCl₂ at the same time that a splitting of the molecule yields chloroform and tetrachlorethylene, C₂Cl₄. Zinc and barium chlorides have little action. Aluminum chloride gives a reaction which is limited by the reverse combination of the chloroform with the tetrachlorethylene (902).

880. Benzyl chloride is easily decomposed by various anhydrous chlorides particularly those of barium and nickel, into hydrochloric acid and a very high molecular weight compound of the empyrical formula C_7H_{\bullet} , previously discovered by Cannizzaro and identical with

- ² Badische, German patent, 255,519 (1913).
- BADISCHE, French patent, 441,203.
- ⁴ Kerez, Annalen, 231, 306 (1885).
- 5 SABATIER and MAILHE, Compt. rend., 141, 238 (1905).
- ⁶ Bosseen, van der Scheer and de Vogt, Rec. Trav. Chim. Pays-Bas, 34, 78 (1915).

that formed by the dehydration of benzyl alcohol (714) and which is perhaps hexaphenyl-cyclohexane, (C₆H₅.CH)₆. The reaction is:

$$x(C_6H_5.CH_2Cl) = xHCl + (C_6H_3.CH)_{x}.$$

- 881. Anhydrous metaflic oxides can likewise effect the decomposition of alkyl halides in consequence of the formation of a certain amount of the corresponding chloride. When isobutyl chloride is passed over alumina at above 250°, the slight dissociation of the alkyl chloride at that temperature can account for the formation of a little chloride or oxy-chloride of aluminum which starts the catalytic action and the amount of which increases rapidly in consequence of the hydrochloric acid evolved. This is the explanation of the decompositions of alkyl chlorides with alumina that have been described. Thoria above 390° has been proposed for the decomposition of tetrachlorethans into trichlorethylens: there is the simultaneous formation of carbon hexachloride, C₆Cl₄.
- 882. It is probably the formation of nickel chloride also, to which may be attributed the identical catalytic effect of *reduced nickel* on alkyl chlorides in the presence of hydrogen; the decomposition takes place easily above 250°. 10

§ 2. — CONDENSATIONS EFFECTED BETWEEN MOLECULES WITH ELIMINATION OF HYDROGEN HALIDE

883. Anhydrous aluminum chloride causes condensation with elimination of hydrochloric acid between aromatic hydrocarbons and various alkyl or cyclo-alkyl chlorides and bromides effecting the synthesis of a large number of aromatic compounds. This is the basis of the *Friedel* and *Crafts synthesis*. 11

I. Alkylation of Aromatic Hydrocarbons

- 884. Method of Operating. It is common to use a large well dried flask with a stopper through which passes a very large tube the upper end of which is closed by a stopper and which permits the introduction of the solid aluminum chloride and on the side of which is fused a tube inclined upward and connected with a reflux condenser. The hydrogen chloride which is evolved escapes at the top of this con-
 - ⁷ CANNIZZARO, Annalen, 92, 114 (1854).
 - ⁸ SENDERENS, Bull. Soc. Chim. (4), 3, 823 (1908).
- CHEM. FABR. BUCKAU, German patent, 274,782, J. Soc. Chem. Ind., 33, 807 (1914).
 - 10 SABATIER and MAILHE, Compt. rend., 138, 407 (1904).
 - ¹¹ Friedel and Crafts, Ann. Chim. Phys. (6), 1, 489 (1884).

denser and may be led into a tarred flask of water, the gain in weight of which serves as a means of following the reaction so that it may be stopped when the theoretical amount of this acid has been liberated.

The aromatic hydrocarbon in large excess (usually 10 times the calculated amount) is mixed with the halogen compound with which it is to react and put into the flask which is warmed on the water bath. The well pulverized anhydrous aluminum chloride is added in small portions, 2 to 20 g. at a time. Whenever the evolution of hydrogen chloride dies down a fresh portion of the chloride is added.

If the alkyl halide is a gas (methyl or ethyl chloride), it may be passed into the flask after the addition of a certain amount of aluminum chloride.

When the reaction is considered finished, the flask is cooled and the mixture is poured into a large excess of cold water acidulated with hydrochloric acid; the oily layer is separated, washed and dried and fractionated.

The simplest case is methyl chloride with bensene:

$$C_0H_0 + CH_3Cl = HCl + C_0H_5 \cdot CH_3 \cdot \frac{C_0H_5 \cdot CH_3}{toluone}$$

885. Often excellent yields are obtained but the chief product is always accompanied by others, particularly the di- and tri- substituted, resulting from the reaction of the first product with a second molecule of the halide. Thus in the simplest case, that of methyl chloride on benzene, the latter reacts with the toluene that is formed to give a mixture of the xylenes. These can react in their turn to yield trimethyl-benzenes (1, 2, 4 and 1, 3, 5) and if the reaction is prolonged, tetramethyl-benzene (1, 2, 4, 5), then pentamethyl- and finally hexamethyl-benzene are formed.

By stopping the reaction when the calculated amount of acid has been evolved these complications are avoided for the most part.

The yield of monosubstituted hydrocarbon is considerably increased when an amount of aluminum chloride equal to 15 or 20 % of the weight of the alkyl chloride is used.

The use of carbon disulphide as a solvent sometimes facilitates the reaction.¹²

Alkyl chlorides, bromides or iodides may be used interchangeably, the latter evolving hydrogen bromide and iodide.

886. In place of using aluminum chloride as above described, the flask may be filled with aluminum turnings (previously cleaned by boiling with alcohol and washing with ether) and a current of dry hydro-

¹⁹ Anschütz, Annalen, 235, 207 (1886).

gen chloride passed.¹³ Aluminum turnings may be used with mercuric chloride which attacks the metal rapidly forming the chloride ¹⁴

In some cases the aluminum chloride is put in the flask first and covered with carbon disulphide and then the mixture of the two substances that are to react is run in.

887. Reversal of the Reaction. The addition of alkyl groups may be limited by the reverse reaction of removing them, and this is also catalyzed by aluminum chlorids.

When the poly-alkyl benzenes are treated with aluminum chloride and a current of hydrogen chloride, the alkyl side chains are eliminated as alkyl chlorides. ¹⁵ From hexamethyl-benzene we may pass to pentamethyl-, to tetramethyl- (1, 3, 4, 6) and (1, 3, 4, 5) to trimethyl- (1, 3, 4) and (1, 3, 5) to meta and para xylenes, then toluene and finally benzene. ¹⁶

888. It may happen that side chains are taken off of one molecule and put on another in consequence of splitting off an alkyl halide which then reacts with the other molecule.

Thus poly-ethyl-benzenes in presence of benzene and aluminum chloride, retrograde towards ethyl-benzene, particularly in a current of hydrogen chloride which carries off the ethyl chloride.¹⁷

Ethyl-benzene kept in contact with aluminum chloride furnishes simultaneously benzene and disthyl-benzene. Isomerizations may result from an alkyl group being taken off and put on again. From p.xylene we may get m.xylene and inversely; pseudocumene (1, 3, 4 - tri-methyl-benzene) may give mesitylene (1, 3, 5).18

889. Results Obtained. The reaction goes well with various aromatic hydrocarbons, benzene and its homologs as well as with naphthalene 19 and diphenyl. The homologs of benzene frequently give better results than benzene itself.

It was developed first for alkyl mono-chlorides but may go equally well with cyclohexyl monochlorides: cyclohexyl chloride and benzene give phenyl-cyclohexane.²⁰

- 13 STOCKHAUSEN and GATTERMANN, Berichte, 25, 3521 (1891).
- 14 RADZIEWANOWSKI, Berichte, 28, 1135 (1895).
- ¹⁵ JACOBSEN, Berichte, 18, 339 (1885).
- ¹⁶ This reaction has been extensively used for the manufacture of toluene from the xylenes. E. E. R.
- ¹⁷ RADZIEWANOWSKI, Berichte, 27, 3235 (1894). BOEDTKER and HALSE, Bull. Soc. Chim. (4), 19, 444 (1916).
 - ¹⁸ Anschütz and Immendorf, Berichte, 17, 2816 (1884), and 18, 657 (1885).
- ¹⁹ It is remarkable that when a solution of naphthalene in bensene is treated with phthalic anhydride in the presence of aluminum chloride, the naphthalene reacts to the exclusion of the bensene. Heller and Schülke, Berichte, 41, 3627 (1908). E. E. R.
- ²⁶ KOURSANOF, J. Russian Phys. Chem. Soc., 33, 527 (1901); Bull. Soc. Chim. (3), 28, 271 (1902).

It is also applicable to the *chlormethyl ethers*, R.O.CH₂Cl, which form the ether R.O.CH₂R' with an aromatic hydrocarbon, R'H. With *benzens* the reaction goes regularly in the cold but the yield is only 30%, because *benzyl chlorids* is also formed by a side reaction which liberates the alcohol, R.OH (818).²¹

The reaction applies to derivatives of aromatic hydrocarbons which are chlorinated in a side chain, e.g. benzyl chloride, C₆H₆.CH₂Cl.²²

Unsaturated monochlorides or monobromides may be used. Thus vinyl bromide, CH₂: CHBr, condenses with benzene to form styrens.²²

890. Dihalogen derivatives may also be used. Ethylene chloride reacts with benzene to form symmetrical diphenyl-ethane ²⁴ and 1, 1-dibrom-ethylene forms 1, 1-diphenyl-ethylene, CH₂: C(C₆H₅)₂. ²⁵

Ethylidene chloride, CH₂.CHCl₂, gives similarly 1, 1-diphenylethane, CH₂.CH(C₆H₅)₂, but the reaction may be complicated by the formation of ethyl-benzene and dihydro-dimethyl-anthracene.²⁶

Benzal chloride, C₄H₅.CHCl₂, with 5 parts of benzene and a little aluminum chloride, yields triphenyl-methane,²⁷ which may also be formed from benzene and chloroform, CHCl₃.²⁸

II. Synthesis of Ketones

891. The Friedel and Crafts reaction is still more easily applied to the production of ketones, by the reaction of aromatic hydrocarbons with carbonyl chloride, or with the chlorides of aliphatic or aromatic acids.

Thus carbonyl chloride and benzene form benzophenone:

$$COCl_2 + 2C_6H_6 = 2HCl + C_6H_5 \cdot CO \cdot C_6H_5$$

Acetyl chloride produces acetophenone from benzene:

$$CH_3.COCl + C_6H_6 = HCl + C_6H_5.CO.CH_3.$$

- 892. For these preparations equal molecules of the hydrocarbon and the acid chloride are mixed and carbon disulphide, ligroine or nitrobenzene is added till a limpid liquid is obtained. Care must be taken to protect from all moisture. This solution is added a little at a time to another flask which contains an equal volume of solvent and
 - ²¹ SOMMELET, Compt. rend., 157, 1443 (1913).
 - ²² Friedel and Crafts, Ann. Chim. Phys. (6), 1, 478 (1884).
 - ²⁸ Anschütz, Annalen, 235, 231 (1886).
 - ²⁴ SILVA, Compt. rend., 89, 606 (1879).
 - 25 Demole, Berichte, 12, 2245 (1879).
 - ²⁶ GENVERESSE, Bull. Soc. Chim. (2), 49, 579 (1888).
 - ²⁷ LINEBURGER, Amer. Chem. Jour., 13, 270 (1891).
- ²⁶ FRIEDEL and CRAFTS, Bull. Soc. Chim. (2), 37, 6 (1882).— E. and O. FISCHER, Annalen, 194, 252 (1878).— ALLEN and KÖLLIKER, Annalen, 227, 107 (1885).

aluminum chloride equal in weight to the acid chloride.²⁰ The mixture is warmed slowly on the water bath till no more hydrogen chloride is evolved.

Nitrobenzene as a solvent has the advantage of dissolving aluminum chloride.²⁰

The aluminum chloride may be added a little at a time to the mixture of the hydrocarbon and the acid chloride.

893. Results. Acetyl chloride, CH₂.COCl, condenses with benzene to form acetophenone, C₆H₅.CO.CH₂,²¹ while benzoyl chloride, C₆H₅.-COCl, gives benzophenone, C₆H₅.CO.C₆H₃,²² which may also be obtained by condensing benzene with carbonyl chloride. Benzoyl bromide may be used with benzene and aluminum bromide.²²

Chlor- brom-, or nitro- ring substitution products of the aromatic acid chlorides may be used with the same facility. Thus m.nitroben-zoyl chloride, O₂N.C₆H₄.COCl, reacts with benzene to form m.nitrobenzophenone, O₂N.C₆H₄.CO.C₆H₅,³⁴ and similar compounds can be obtained from the chlor ³⁵ and brom ³⁶ derivatives.

The chlorides of dibasic acids can give a double reaction to form diketones. Thus succinyl chloride and benzene furnish 1, 4-diphenyl-butadione (1,4), C₆H₅.CO.CH₂.CH₂.CO.C₆H₅. The reaction is carried out in carbon disulphide.⁵⁷

Malonyl and glutaryl chlorides react similarly.88

On account of its tautomeric nature, phthalyl chloride can give different products according to the way the reaction is carried out.

- ²⁰ It is better to calculate the amount from its molecular weight and that of the acid chloride; to 1 mol. RCOCl, 1 mol. AlCl₂ = 133.5, is required but 10% excess is of advantage. E. E. R.
 - ³⁰ Behn, German patent, 95,901 (1897).
 - ²¹ Friedel and Crafts, Ann. Chim. Phys. (6), 14, 455 (1888).
 - 22 FRIEDEL and CRAFTS, Ann. Chim. Phys. (6), 1, 510, and 518 (1884).
 - ²⁵ OLIVIER, Rec. Trav. Chim. Pays-Bas, 37, 205 (1918).
 - ²⁴ Grigy and Königs, Berichte, 18, 2401 (1885).
- ²⁶ Overton, Berichte, 26, 29 (1893). Hantesch, Berichte, 24, 57 (1891). Demuth and Dittrich, Berichte, 23, 3609 (1890).
 - ²⁶ CATHCART and MEYER, Berichte, 25, 1498 (1892).
 - ²⁷ CLAUS, Berichte, 20, 1375 (1887).
 - ³⁶ Auger, Ann. Chim. Phys. (6), 22, 349 (1891).
- ** FRIEDEL and CRAFTS, Ann. Chim. Phys. (6), 1, 523 (1884). BARYER, Annalen, 202, 51 (1880).
 - 44 HALLER and GUYOT, Bull. Soc. Chim. (3), 17, 873 (1897).

o. benzoyl-benzoic acid, 41 C₆H₅.CO.C₆H₄.COOH, and other products are obtained.

Acid chlorides may react with pyridine or quinoline in the presence of aluminum chloride to give ketones when traces of thionyl chloride are present. From benzoyl chloride and pyridine, pyridyl-phenyl-ketone is obtained: 48

$$C_0H_5.CO.Cl + C_5H_5N = HCl + C_0H_5.CO.C_5H_4N.$$

894. Thiophosgene, CSCl₂, reacts with aromatic hydrocarbons to form thioketones: thus with benzene, thiobenzophenone, C₄H₄.CS.-C₄H₄.44

III. Formation of Amides

895. By the action of carbamic chloride, Cl.CO.NH₂, aromatic amides are formed: thus from benzene, benzamide, C₆H₅.CO.NH₂ is obtained.⁴⁵

IV. Formation of Cyclic Compounds

896. Methylene chloride condenses with diphenyl, C₆H₅. C₆H₅, to form fluorene,

897. Tetrabromethane (1,1,2,2,), or acetylene tetrabromide, reacts with benzene to form anthracene:

Condensation may take place between two or more molecules of a chlor-compound. Thus 2-phenyl-ethyl chloride, C₆H₃.CH₂.CH₂Cl, reacts vigorously with aluminum chloride in carbon disulphide or ligroine to form an insoluble resin (C₆H₄.CH₂CH₃)x.

Dissolved in 6 parts of ligroine with 1 part of aluminum chloride, 4-phenyl-butyl chloride gives an excellent yield of tetrahydro-naphthalene:

- 41 SCHEIBER, Annalen, 389, 121 (1912).
- ⁴² COPISAROW, J. Chem. Soc., 111, 10 (1917).
- WOLFFENSTEIN and HARTWICH, Berichte, 48, 2043 (1915).
- 44 BERGREEN, Berichte, 21, 341 (1888).
- 46 GATTERMANN, Annalen, 244, 29 (1888).
- 44 ADAM, Ann. Chim. Phys. (6), 15, 253 (1888).
- 47 ANTSCHÜTE, Annalen, 235, 165 (1886).

Similarly 5-phenyl-pentyl chloride gives phenyl-cyclopentane boiling at 213°.48

898. Mechanism of the Reaction. We have shown above how the rôle of the aluminum chloride in the Friedel and Crafts reaction may be explained (173). The catalytic nature of the action is not doubted though sometimes it is necessary to employ large amounts of the salt, sometimes larger than the amount of the aromatic hydrocarbon. This is the case when the aluminum chloride combines with one of the products of the reaction and is thus withdrawn from its catalytic function.⁴⁹

899. Other Catalytic Chlorides. Several anhydrous metallic chlorides can be employed in the same way as aluminum chloride in the Friedel and Crafts synthesis: zinc, ferrous, ferric and stannic chlorides and antimony pentachloride.

The use of ferric chloride is quite advantageous in preparing ketones. Thus benzoyl chloride and benzene give benzophenone. Its action, like that of the other chlorides mentioned above, is milder than that of aluminum chloride. For that reason these chlorides sometimes give rise to less formation of byproducts.

For the preparation of bensophenone, the following comparative yields have been obtained: 51

With	aluminum chlorid	e						70–71 %
	ferric chloride .							60-62
	zine chlorida							26_32

Aluminum chloride serves poorly for condensing toluene with chlormethyl ethers (889), while good results are obtained with antimony pentachloride and particularly with stannic chloride.⁵²

The use of *zinc chloride*, or better metallic zinc which immediately forms some of the chloride, has been recommended for reactions with naphthalene. Thus the *di-naphthyl* ketones are prepared by the action of zinc on a mixture of naphthalene with α - or β -naphthoyl chlorides. α -

900. A different isomer may be obtained when other chlorides are substituted for the aluminum chloride. Isobutyl chloride condensed

- 48 VON BRAUN and DEUTSCH, Berichte, 45, 1267 (1912).
- 49 HELLER and SCHULKE, Berichte, 41, 3627 (1908).
- ⁵⁶ NENCKI, Berichte, 30, 1766 (1897), and 32, 2414 (1899). MEISSEL, Berichte, 32, 2419 (1899).
 - 61 GANGLOFF and HENDERSON, J. Amer. Chem. Soc., 30, 1420 (1917).
 - ³⁰ SOMMELET, Compt. rend., 157, 1443 (1913).
 - ALEXYEF, Meth. de transform. des comb. organ., Paris, 1891, 186.
 - GRUCAREVIC and MERS, Berichte, 6, 1242 (1877).

with toluene in the presence of ferric chloride gives p.methyl-isobutylbenzene, while in the presence of aluminum chloride the meta compound is obtained.⁵⁶

Formation of Aromatic Amines by Hofmann's Reaction

901. Traces of cuprous iodide can readily effect the condensation of primary aromatic amines with phenyl bromide, with elimination of hydrobromic acid. The acetyl derivative of the amine may be used. Thus by boiling 90 g. brombenzene, 10 g. acetanilide, 6 g. sodium carbonate and a little cuprous iodide for 15 hours, acetyl-diphenyl-amine is obtained and this can readily be transformed into diphenyl-amine. The cuprous iodide can be replaced by copper and iodine or even by copper and potassium iodide. 56

The presence of copper powder greatly facilitates the action of ammonia under pressure at 170° on chlor-nitro-benzene to form amino-nitro-benzene.

It is also useful in the similar reaction of aniline or its homologs on o.chlor-benzoic and 2,4-chlor-nitro-benzoic acids in the preparation of the corresponding amino compounds.⁵⁷

Likewise pyridine heated 7 hours to 250° with benzyl chloride and a little copper powder gives a good yield of 2-benzyl- and 4-benzyl-pyridine. Ethyl iodide and pyridine give the ethyl-pyridines under the same circumstances. The copper can be replaced by cuprous chloride. Aluminum and magnesium powders give poorer results.⁵⁸

Condensations in the Aliphatic Series by Anhydrous Chlorides

902. The use of ferric chloride enables us to effect important condensations in the aliphatic series. Thus with propionyl chloride in the presence of alcohol, two molecules of the acid chloride condense to form the ester of a keto-acid: 50

$$CH_1.CH_2.COCl + CH_1.CH_2.COCl + C_0H_0OH = CH_1.CH_2.CO.CH(CH_0).CO_2C_2H_0 + 2HCl.$$

903. Chloroform condenses with pentachlorethane on contact with aluminum chloride with evolution of hydrogen chloride to form heptachlor-propane: 60

$$CHCl_2 + CCl_2 \cdot CHCl_2 = HCl + CHCl_2 \cdot CCl_2 \cdot CCl_3$$

- BIALOBRZESKI, Berichte, 30, 1773 (1897).
- ⁵⁶ GOLDBERG, Berichte, 40, 4541 (1907).
- ⁶⁷ ULLMANN, Annalen, 355, 312 (1907).
- 56 CHICHIBABINE and RYUMSHIN, J. Russian Phys. Chem. Soc., 47, 1297 (1915).
- 50 HAMONET, Bull. Soc. Chim. (3), 2, 334 (1899).
- 60 PRINS, J. prakt. Chem. (2), 89, 414 (1914).

§ 3. — ELIMINATION OF A MOLECULE OF AN ALKALINE CHLORIDE, BROMIDE, OR IODIDE

904. The action of the aromatic halogen derivatives, phenyl chloride, bromide, and iodide on the alkali salts of phenol should form phenyl ether, but practically the yield is trifling. It becomes very high when the reaction is carried on under pressure at 150° to 200° in the presence of finely divided copper as catalyst. The yield reaches 25% with the chloride, 82% with the iodide and 78% with the iodide.

This process may be applied to the formation of ethers of diphenols.⁶¹

el Ullmann and Sponagel, Annalen, 360, 83 (1907).

CHAPTER XXI

DECOMPOSITIONS AND CONDENSATIONS OF HYDROCARBONS

905. The action of high temperature on hydrocarbons is to dissociate the molecules, from which hydrogen tends to separate, at the same time that it produces a greater or less breaking up of the molecules into groups, CH₂, CH₂, and CH which are capable of uniting to form new complex molecules. There result complicated mixtures of varied constitution to which Berthelot has given the name of pyrogenetic equilibria which as the temperature rises tend to produce larger and larger proportions of hydrogen and methane along with substances very rich in carbon and very condensed hydrocarbons.

906. The petroleum industry has taken advantage of reactions of this sort in the process known as "cracking." This process, which was accidentally discovered at Newark, N. J., in 1861, consists in carrying petroleum vapors to high temperatures, above a dull red. Along with usable gases, new hydrocarbons are produced which increase the proportion either of gasoline or of heavy oils as compared with the original oil.

The effect of temperature begins to be felt at about 325° but is not important below a red heat. The presence of catalysts lowers the temperature of these reactions and makes them easier to carry out. The finely divided metals, copper, iron, cobalt, nickel, platinum, magnesium, and aluminum can be employed and so may the anhydrous oxides, titania, zinc oxide and alumina, etc.¹

It is important to know the results of the pyrogenetic decomposition of the hydrocarbons in the absence of catalysts.

From this point of view, benzene, petroleums and the coal tar hydrocarbons known as solvent naphtha have been the most studied.

907. Benzene is hardly affected below 500°, at which it begins to decompose into diphenyl, the formation of which increases till it reaches a maximum at 750°. It is accompanied by diphenyl-benzene: carbon is deposited and hydrogen set free without any production of acetylene or of naphthalene below 800°.

¹ ZELINSKI, J. Russian Phys. Chem. Soc., 47, 1808 (1915).

² ZANETTI and EGLOFF, J. Ind. Eng. Chem., 9, 356 (1917).

908. American petroleum, under the action of heat alone, gives increasing amounts of gas from 450° to 875°, while the density of the liquids produced increases also with the temperature. Between 450° and 600°, the products formed contain more toluene than xylene, more xylene than benzene, and neither naphthalene nor anthracene. At 650° the proportion of benzene is still lower than that of toluene but above that of xylene. From 700° to 850°, benzene is more abundant than toluene and especially than xylene. The formation of naphthalene begins at 750° and that of anthracene at 800° and both increase rapidly with the temperature.

For 100 parts of petroleum thus treated, the benzene in the product reaches its maximum of 4.7% at 750°, toluene its maximum of 3.1% at 650°, and xylene its maximum of 1.9% at 700°. At 800° we have 2% naphthalene and 0.3% anthracene. These aromatic hydrocarbons are associated with various aliphatic.

909. Solvent naphtha contains considerable amounts of higher hydrocarbons. When it is heated in steel tubes under 11 atmospheres pressure to 500-800°, it yields considerable amounts of lower hydrocarbons. In the product, benzene reaches its maximum of 42.5% at 800° and toluene its maximum of 39.9% at 750°. But as the temperature is raised higher and higher the yield of liquid decreases rapidly in consequence of the more abundant production of gaseous products and of materials poor in hydrogen, the real maximum yield based on 100 parts of solvent naphtha is:4

Under the action of a red heat, pinene gives a large number of hydrocarbons, both gaseous and liquid, among which have been found benzene, toluene, m.xylene, naphthalene, anthracene, methyl-anthracene and phenanthrene.

By operating at a barely visible red, along with a terpene isomeric with pinene but boiling higher, *isoprene*, C₅H₅, benzene and its homologs, and poly-terpenes are formed.⁷

Action of Catalysts.

- 910. The presence of catalysts usually enables us to carry out the same reactions at lower temperatures which is more favorable to the preservation of sensitive products that may be formed. Usually
 - ² EGLOFF and Twomey, J. Phys. Chem., 20, 121 (1916).
 - ⁴ EGLOFF and MOORE, J. Ind. Eng. Chem., 9, 40 (1917).
 - ⁵ Berthelot, Ann. Chim. Phys. (3), 39, 5 (1853), and (4), 16, 165 (1869).
 - SCHULTZ, Berichte, 10, 114 (1877).
 - ⁷ TILDEN, Ann. Chim. Phys. (6), 5, 120 (1885).

nickel and iron act violently tending to produce very advanced dehydrogenation with charring more and more intense as the temperature is raised.

911. Aliphatic Hydrocarbons. Methane is only slightly attacked by nickel up to 360° but towards 390° the deposition of carbon is appreciable.

The decomposition is not yet rapid at 910° at which methans heated 10 minutes in a porcelain tube, without catalyst, gives only 10% of hydrogen. The presence of silica in the tube does not increase the decomposition, but with lims the proportion of hydrogen reaches 35%, with wood charcoal, 69% while with metallic iron it is 73%.

Ethane decomposes slowly above 325° giving carbon, methane and free hydrogen.

Pentane decomposes in an analogous way: at 350-400° methane is produced with intermediate hydrocarbons and carbon is deposited on the nickel.

Lengthening the carbon chain makes these decompositions more easy;¹⁰ but only above 550° and towards 600° are the liquid hydrocarbons such as are found in Pennsylvania petroleum attacked.

912. Unsaturated Hydrocarbons. If a current of ethylene is passed over reduced nickel heated above 300° the nickel can be seen to swell up into a voluminous black material which finally fills the tube and chokes it up: all the ethylene disappears and a gas is obtained containing ethane, methane, and hydrogen. The proportion of ethane is less with higher temperatures of the metal: at a dull red only traces of it are left.

In contact with nickel, ethylene is decomposed into carbon and hydrogen, but the latter is taken up immediately by a portion of the ethylene to form ethane which is more and more broken down to methane at higher temperatures. The nickel is found diffused in the carbon that is formed.¹¹

Propylene suffers an analogous destruction but more slowly and without the voluminous swelling of the metal. The decomposition is appreciable at 210° and is clean at 350°. The escaping gas contains propylene, propane, ethylene, ethane, methane and hydrogen.¹²

All other unsaturated hydrocarbons give analogous results, e.g. the

SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 435 (1905).

SLATER, J. Chem. Soc., 109, 160 (1916).

¹⁶ SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 435 (1905).

¹¹ SABATIER and SENDERENS, Compt. rend., 124, 616 and 1358 (1897).

¹² Sabatier and Senderens, Compt. rend., 134, 1128 (1902).

vapors of trimethyl-ethylene give along with a deposit of carbon, the saturated hydrocarbon with the whole series of lower hydrocarbons.

Cobalt acts in a similar manner but less actively than nickel. With ethylene at 360° and even at 425° there is slow carbonization without rapid swelling and much ethylene survives.

Iron does not act till above 350° and gives a still slower decomposition.

Platinum (black or sponge) and reduced copper do not have any appreciable action on propylene or ethylene.¹³

- 913. Acetylene Hydrocarbons. Similar dehydrogenating actions, but less intense, are exercised by the finely divided metals on the aletylene hydrocarbons, especially acetylene. The action can be divided into two entirely distinct kinds, which coexist. One of these is easily observed with platinum or iron, the other particularly with copper, while nickel superimposes the two effects.
- 914. First Kind of Reaction. Pure acetylene when heated with platinum to 150°, is rapidly decomposed into carbon and hydrogen: the heat evolved by this decomposition heats the metal to incandescence which accelerates the destruction giving rise to a great carbonaceous swelling, and which causes the polymerization of the remaining acetylene into benzene, styrene, and hydrides of naphthalene and anthracene as in the celebrated synthesis of Berthelot. This phenomenon was observed by Moissan and Moureu;¹⁴ it is complicated by an important consecutive action, which escaped these chemists but which Sabatier and Senderens have studied.¹⁵

The hydrogen resulting from the decomposition of one portion of the gas can act on another portion, in the presence of platinum, to form *ethylene* and *ethane*. The liquid collected is small in amount and is chiefly benzene. This is the composition by volume of the gases evolved:

Acetylene								66.2%
Benzene (vapor)								2.8
Ethylene								
Ethane								
Hydrogen								

915. A much greater destructive activity belongs to reduced iron (obtained at about 450°) which is raised by acetylene from room temperature to incandescence. If the tube containing the iron is not heated, the reaction almost stops with the local decomposition due to

¹³ SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 436 (1905).

¹⁴ Moissan and Moureu, Compt. rend., 122, 1241 (1896).

¹⁵ SABATIER and SENDERENS, Compt. rend., 131, 40, 187 and 267 (1900).

the incandescence, the formation of black voluminous carbon in which the iron is disseminated and of brown liquids, almost entirely aromatic. The gases remaining are little but surviving acetylene and hydrogen saturated with benzene vapor. But if the entire iron tube is kept at above 180°, the hydrogenation of the acetylene is carried on by the metal beyond the incandescent portion so that little acetylene is left and the gas is only hydrogen, ethylene and ethane with the vapors of higher hydrocarbons.

916. Second kind of Reaction. This is caused by copper.

If a current of acetylene is passed over light copper (obtained by reduction at a low temperature) at 180° the copper is seen to turn brown at once and the pressure diminishes greatly on account of the rapid condensation of the acetylene in contact with the metal. Sometimes the current of acetylene which was 20 cc. per minute is entirely taken up for more than 20 minutes and then slowly begins to pass. At this moment the copper is seen to swell rapidly taking on a lighter tint of brown and soon filling the tube so as to stop the flow of gas.

The condensed liquid is a mixture of unsaturated and aromatic hydrocarbons (benzene, styrene etc.), the presence of the styrene causing partial solidification after a time. The small amount of gas that passes out contains, with a small amount of acetylene, hydrogen, ethane and particularly the unsaturated hydrocarbons, ethylene, propylene and butylene, which constitute more than two thirds of it.

The copper is found disseminated in the entire brown solid material formed. If a small portion of this is placed as a layer in another tube and heated to 180-250° in a current of acetylene, the material swells up, again filling the tube. One can start anew with a portion of this material and fill another tube. After three or four such swellings, a material is obtained which is no longer changed when heated in acetylene. This is a lighter or darker brown solid which appears under the microscope to be a thick felt of very fine filaments. It is light and fluffy and may be agglomerated into masses resembling tinder. It is a hydrocarbon of the empyrical formula C_7H_6 in which is found diffused a little copper (about 1.5%) which has caused its formation: this is cuprene. It is composition is identical with that of the condensed hydrocarbon formed by the decomposition of benzyl chloride by metallic chlorides (880) or by the dehydration of benzyl alcohol (714), and is perhaps hexaphenyl-cyclohexane, $C_6H_6(C_6H_8)_6$. 18

¹⁶ On account of polymerization of the styrene to a solid. — E. E. R.

¹⁷ SABATIER and SENDERENS, Bull. Soc. Chim. (3), 21, 530 (1899). — Compt. rend., 130, 250 (1900). — SABATIER, 3rd Congress on Acetylene, Paris, 1900, 345 and 4th Cong. Applied Chem., Paris, 1900, 3, 134.

¹⁸ SABATIER and MAILHE, Ann. Chim. Phys. (8), 20, 298 (1918).

The formation of cuprene is doubtless due to the formation of an unstable acetylide capable of reacting with acetylene to form a new condensed molecule, thus:

and
$$nC_2H_2 + nCu = nC_2Cu + nH_2$$

$$nC_2Cu + 6nC_2H_2 = \underbrace{(C_0H_7)_{2n}}_{\text{cuprene}} + nCu.$$

The regenerated metal is capable of repeating the reaction indefinitely. The hydrogen set free combines with a portion of the acetylene over the copper to give chiefly ethylene hydrocarbons.

Compact copper, in sheet or wire, gives a similar formation when heated in acetylene to 200-50° and covers itself with a brown coating which becomes more yellow as its thickness increases.

917. Superposition of the Two Kinds of Reaction. If over a layer of reduced copper heated at its middle portion to above 250° a rapid current of acetylene be passed, incandescence accompanied by intense swelling is observed at this point and there is simultaneous production of carbon and of cuprene formed by the superposition of the two reactions.

918. Reduced nickel usually causes both reactions. If the reduced nickel is entirely freed from the hydrogen absorbed by its particles, it no longer becomes spontaneously incandescent in acetylene and can be heated to 150° before it causes a reaction. It is only above 180°, that a slow reaction takes place, without incandescence, and this reaction remains thus if the passage of the gas is not too rapid. The metal turns black and swells a little, covering itself with a solid brownish fibrous silky hydrocarbon which suggest cuprene; but this formation is slow and if one tries to accelerate it by passing the acelylene more rapidly or by elevating the temperature, incandescence appears bringing rapid decomposition with charring.

Usually when acetylene is passed over a layer of recently reduced, nickel without precautions, there is immediate spontaneous incandescence, brought on by the occluded hydrogen, and carbonizing decomposition takes place always followed by the hydrogenation of the acetylene and of a part of the aromatic hydrocarbons resulting from the incandescence because the nickel is capable of effecting such hydrogenations.

Summing up, nickel acting on acetylene at 180° produces a triple effect:

1st. Rapid decomposition into carbon and hydrogen with polymerization to aromatic hydrocarbons.

2nd. Slow condensation into a solid hydrocarbon doubtless identical with cuprene.

3rd. Hydrogenation of the acetylene and of the aromatic hydrocarbons with production of aliphatic, unsaturated and cyclo-aliphatic hydrocarbons.

With a tube that is not externally heated, where the incandescence is intense and localized at a single point, the first effect is the greatest, the velocity of the gas rendering the subsequent hydrogenation unimportant. These are the conditions studied by Moissan and Moureu.

919. With cobalt quite free from nickel and reduced from oxide at below 350°, incandescence is not obtained, starting with the tube cold, but is readily started by heating some point on the tube, and is easily maintained if the tube is heated to 200°. The action is intermediate between that of iron and that of nickel. The tube is filled with a black mass consisting of carbon in which the cobalt is diffused and traces of a hydrocarbon analogous to cuprene can be seen.¹⁹

920. The effects of nickel, iron, cobalt and copper are much less intense when these metals are employed in the form of sheets and have appeared to many observers negligible even up to 600°. On the contrary magnesium powder acting at 600° on methane, ethane, ethylene and acetylene causes a 95% decomposition. Aluminum powder, at near the fusion point of the metal, causes a total decomposition while, platinum decomposes only 80%.²⁰

Hexane, under high pressure, is attacked energetically at 650-700° in an iron tube in the presence of alumina.²¹

921. Cyclic Hydrocarbons. As has been said above (640), the hydro-cyclic hydrocarbons in contact with finely divided metals form the corresponding aromatic hydrocarbons by loss of hydrogen; but the cyclic hydrocarbons, benzene, its homologs, naphthalene, anthracene etc. are themselves attacked, and tend to be resolved into CH₂ and CH groups like those furnished by the aliphatic hydrocarbons. Metallic oxides also can catalyze decompositions of this sort.

Finely divided nickel, iron and cobalt act energetically above 400° and especially at a dull red heat, on the hydro-cyclic hydrocarbons, among which are the terpenes, and cause, along with dehydrogenations which take place at lower temperatures (640), decompositions more and more serious as the temperature is raised, and accompanied by carbonaceous deposits which increase at the same time. The charring is less intense with *copper*.

The aromatic hydrocarbons, benzene and its homologs, are much

- 19 SABATTER and SENDERENS, Ann. Chim. Phys. (8), 4, 430 (1905).
- * Kusnetzow, Berichte, 40, 2871 (1907).
- ²¹ IPATIEF and DOVGELEVICH, J. Russian Phys. Chem. Soc., 43, 1431 (1911); C. A., 6, 736 (1912).

less affected by the action of finely divided metals than when they are acted on in the nascent state, that is when they are being formed by the dehydrogenation of cyclohexane or terpene hydrocarbons.

922. The Case of Pinene. The action of heat on pinene when its vapors are passed through a red hot tube has been described above (909). The tube being of iron and sometimes filled with broken pumice or porcelain the peculiar influence of the metal or of the filling may enter into the reaction.

By passing these vapors over very light finely divided copper (59), in a glass tube heated to 600-30°, a rapid evolution of gas of high illuminating power consisting of hydrogen charged with the vapors of lower hydrocarbons, is obtained. By conducting the operation very slowly, 100 cc. of pinene gave 81 cc. of condensate which contained:

13.3 cc. passing over below 95° to 150° 150° to 185° to 185° over from 9.3 cc. passing over from 9.3 cc. passing over above 185°.

Treatment with sulphuric acid, which dissolves the terpenes and the ethylenic and di-ethylenic hydrocarbons, reduced the volume to 31.5 cc. of hydrocarbons almost entirely nitrifiable and consisting of about 19 cc. cymene and methyl-ethyl-benzene, 10 cc. m. xylene and toluene and a small amount of benzene. In the most volatile portion of the hydrocarbons is found some isoprene, hardly more than 2 cc. The terpenes remaining in the product have no effect on polarized light.

923. Reduced *nickel* acts more violently than copper at 600° and causes intense carbonization in consequence of its destructive action on ethylenic and di-ethylenic hydrocarbons (912). The gas is richer in hydrogen, the liquids condensed are less and contain a considerable proportion of saturated hydrocarbons resulting from the hydrogenating action of the metal on the unsaturated hydrocarbons and unattacked by either sulphuric or nitric acid.²²

Reactions carried out in the Presence of Hydrogen

924. The decompositions of the hydrocarbons by the metals correspond to an elimination of hydrogen of which a portion is utilized for hydrogenating the fragments. It seemed probable that the presence of hydrogen with the hydrocarbon molecules would stabilize them or would favor the hydrogenation of the fragments resulting from their decomposition. The stabilization is actually realized in the

²² SABATIER, MAILHE and GAUDION, Compt. rend., 168, 826 (1919).

case of the cyclohexane hydrocarbons which are preserved to a great extent (640), in the case of the aromatic hydrocarbons derived from the terpenes (644). Hydrogenation carried out at temperatures at which the hydrocarbons are broken up would necessarily lead to the hydrogenation of the fragments that would be formed in the absence of the hydrogen.

925. Acetylene. We have seen (423) that the direct hydrogenation of acetylene, carried out over cold nickel or at a low temperature, gives ethane accompanied by a certain amount of higher aliphatic hydrocarbons, both gaseous and liquid: the reason for the formation of these by-products being the breaking up of the molecule HC: CH, which takes place at near room temperature, thereby liberating the CH groups which are hydrogenated to methane, CH4, or to the groups CH₂ and CH₃, the groups CH₂, CH₂ and CH being able to unite in various ways to give, in the cold, more or less complex aliphatic hydrocarbons. By operating continuously for 24 hours with nickel maintained at 200°, Sabatier and Senderens condensed about 20 cc. of a clear yellow liquid with a splendid fluorescence and an odor quite similar to that of rectified petroleum. It began to boil at about 45° and half of it passed over below 150°, while at 250° there remained a small quantity of very fluorescent orange vellow liquid, certainly containing polycyclic hydrocarbons. The original liquid had a density of 0.791 at 0° and was slightly attacked by the nitro-sulphuric acid mixture which extracted a small amount of aromatic hydrocarbons. The remaining oil had a density of 0.753 at 0° and was composed almost entirely of aliphatic hydrocarbons (pentane, hexane, heptane, octane, nonane, decane, undecane etc.) which were associated in the original product with unsaturated hydrocarbons, soluble in slightly diluted sulphuric acid, and with traces of aromatic hydrocarbons. The composition, odor, density and fluorescence class this liquid with Pennsylvania petroleums.

926. If through a tube containing reduced nickel and kept between 200 and 300°, a rapid current of pure acetylene is passed, without hydrogen, a lively incandescence is obtained on account of the decomposition of the acetylene into carbon and hydrogen (918). A portion of the acetylene thus carried to a high temperature condenses to benzene and other aromatic hydrocarbons according to the reaction discovered by Berthelot; another portion breaks up into CH groups which can be hydrogenated along with the aromatic hydrocarbons by the portion of the nickel layer which remains at 200–300°. In the receiver is collected a considerable amount of liquid, greenish by reflected light, reddish by transmitted, the appearance of which greatly

resembles crude petroleum. If this liquid is hydrogenated directly over nickel at 200° a colorless liquid is obtained which is only slightly attacked by the nitro-sulphuric acid reagent and which, on fractionation, gives a whole series of liquids of densities similar to those of the corresponding fraction of Caucasian petroleum. The chief constituents, as in the petroleum fractions, are the polymethylene hydrocarbons resulting from the hydrogenation of the aromatic hydrocarbons formed by the incandescence. As in the Caucasian petroleum there are certain amounts of aliphatic hydrocarbons resulting from the hydrogenation of the CH groups which are set free and then reunited in various fashions.

927. By causing incandescence in mixtures of acetylene and hydrogen, the proportion of the aliphatic hydrocarbons is increased and the poly-methylenes diminished and *intermediate petroleums* are obtained.

If the hydrogenation following the incandescence takes place at about 300°, the cyclohexane hydrocarbons are formed only incompletely and are accompanied by certain proportions of untransformed aromatic hydrocarbons: we have *Galician petroleum*.

928. Analogous reaction can be effected by finely divided cobalt and to a certain extent by iron. Sabatier and Senderens, who found out the above facts, have based on them a sim le theory of the genesis of natural petroleum. There are doubtless far down in the earth's crust large masses of alkaline and alkaline earth metals as well as of the carbides of these metals. Water penetrating through fissures in the rocks and coming in contact with these materials will evolve hydrogen and acetylene, in proportions which will doubtless vary greatly.

If the hydrogen is in large excess, the gaseous mixture, coming in contact with nickel, cobalt or iron disseminated in adjacent rocks at temperatures which may be lower than 200°, gives rise to American petroleum and at the same time to large quantities of combustible gases in which are found, as in the natural gas of the Pittsburgh district, much methane, ethane and free hydrogen.²³

Use of Anhydrous Aluminum Chloride

929. Anhydrous aluminum chloride heated with aliphatic hydrocarbons, tends to decompose them into lower and higher hydrocarbons. Amylene gives methane at the same time as hexane and still more condensed hydrocarbons.²⁴

930. More important and more regular effects are observed with SABATIER and SENDERENS, Ann. Chim. Phys. (8), 4, 445 (1905). — SABATIER,

Rev. Mois, 1, 257 (1906).

M ASCHAN, Annalen, 324, 1 (1902).

aromatic hydrocarbons, as has been stated above (887), tending to their degradation and building up at the same time. Ethyl-benzene heated with aluminum chloride is degraded to benzene, while diethyl-benzene is formed to compensate (888).

A xylene (in which the meta predominated) boiled for 5 minutes with 2% anhydrous aluminum chloride in an apparatus with a mechanical stirrer gave 29% of hydrocarbons boiling below 135°. Prolonged boiling raised the yield only to 34%, nor does increasing the amount of the chloride increase it sensibly. Benzene is formed chiefly with a little toluene. The proportion of toluene is no better when the operation is carried on under 18 atmospheres pressure.²⁵

Cymene heated with a third of its weight of aluminum chloride gives a mixture which contains: 2% benzene, 42% toluene, and 7% xylene (chiefly meta) with a little di-isopropyl-benzene and methyl-di-isopropyl-benzene, increasing the amount of the catalyst increases the amount of the benzene and decreases the toluene.²⁶

931. Naphthalene heated in an autoclave at 330° under 10 atmospheres pressure, for 20 minutes with 4% anhydrous aluminum chloride gives, along with a carbonaceous and tarry material, 32% of a liquid hydrocarbon of which about half is dihydro-naphthalene resulting from the hydrogenation of one part of the naphthalene at the expense of another.²⁷

Under the action of anhydrous aluminum chloride, pinene gives pentane and its homologs as well as cyclohexene hydrocarbons.²⁸

Application to the Treatment of Petroleums

932. The use of catalysts enables us to improve greatly the operation of cracking (906) for the purpose of increasing the volatile portion of petroleum, since it lowers greatly the temperature at which the reaction takes place thus permitting the survival of molecules resulting from the decomposition which would otherwise be attacked at the higher temperatures.

If over finely divided metals, such as powdered iron or reduced copper, maintained at a temperature between 400° and a dull red, the vapors of a crude petroleum (from any source) or of petroleum previously stripped of its gasoline, there is partial decomposition into a mixture of hydrogen and gaseous hydrocarbons and liquids of which a considerable proportion distils below 150° and may be separated.

F. Fischer and Niggemann, Berichte, 49, 1475 (1916).

²⁶ Schorger, J. Amer. Chem. Soc., 39, 2671 (1917).

²⁷ F. Fischer, Berichte, 49, 252 (1916).

²⁸ STEINKOFF and FREUND, Berichte, 47, 411 (1914).

When the residue is again submitted to the action of finely divided metals, a new amount of volatile liquids is formed, and so on.

The gases evolved are quite abundant and are composed of saturated and unsaturated hydrocarbons having high calorific and illuminating power.

Iron has the inconvenience that it causes an abundant deposit of carbon on its surface. Copper causes much less of this but requires a higher temperature, near to 600°; temperatures below 550° give poor results while above 800° there is intense carbonization with diminution of the yield of gasoline.

Thus starting with an American petroleum containing nothing boiling below 150°, by a single passage over copper at 600°, 1 l. gave 225 cc. gasoline boiling below 150°. At the same time 120 l. of gas was evolved with high illuminating power and having a heating power of 15,000 calories per cu. m.

After some time the copper becomes too much fouled with carbonaceous materials and does not have sufficient activity. In order to regenerate it all that is necessary is to pass over it a current of steam which causes the carbon to disappear without altering the metal while producing water gas which may be used for heating the apparatus.

The liquids thus obtained are composed in part of saturated and aromatic hydrocarbons and in part of hydrocarbons containing one or two double bonds. These are oxidisable and polymerizable and have a disagreeable odor. In the experiment cited above their proportion was 40%.

In order to transform them into saturated compounds without disagreeable odors it is sufficient to hydrogenate their vapors in the presence of finely divided metals (particularly reduced nickel) between 150° and 300°. A hydrocarbon is thus obtained that may be used as gasoline. Furthermore, the two phases of the process can be combined so as to transform continuously a crude petroleum or petroleum residue into gasoline, of which as high as 75% may be obtained.²⁹

933. Numerous patents have been taken out for processes of this sort. One proposes to use finely divided metals at 600° under 6 atmospheres pressure.²⁰

In another patent, the gas issuing from the catalytic cracking is charged with ammonia and thus modified is used to carry along the vapors of the hydrocarbon over metal oxides which can be reduced to the metal. The nascent hydrogen set free by the decomposition of the ammonia by the metal, saturates the hydrocarbons and diminishes the

²⁹ SABATIER, French patent, 400,141, May, 1909.

²⁰ Hall, English patent, 17,121, of 1913; J. S. C. I., 33, 1149 (1914).

amount of the carbonaceous deposits. By this process gasoline free from sulphur is obtained even from Mexican petroleum containing 5% of sulphur.⁸¹

934. Catalytic oxides (titania, alumina, and zinc oxide) can also be utilized for such transformations, particularly for changing Russian cyclohexane petroleums into aromatic hydrocarbons (benzene, toluene and homologs). A benzine from Baku (98 to 102°) gave 30% of aromatic hydrocarbons of which over half was toluene.

The use of iron retorts is to be avoided on account of the intense carbonization which this metal causes and the rapid deterioration which results therefrom.

935. Aluminum chloride enables us to carry out analogous reactions at much lower temperatures (929).

Petroleum freed from water and gasoline is heated 24 to 48 hours with dry aluminum chloride. The products obtained are almost entirely saturated and it is unnecessary to treat them with sulphuric acid, washing with soda and then with water being sufficient to get rid of the hydrogen sulphide. The aluminum chloride is regenerated by submitting the residual coke to a current of chlorine at a red heat. The yield of gasoline from Oklahoma petroleum, which gives only 12.5% by the ordinary cracking process, is raised to 34.8% by this method.²²

936. Iron chlorides, although less active, may be substituted for the aluminum chloride, and with Russian oils, very poor in gasoline, a certain amount of hydrocarbons passing over from 40° to 140° is obtained. Of this about 35% is hexane and heptane while the rest is chiefly 7 and 8 carbon cyclic hydrocarbons. Heavy hydrocarbons the consistence of which resembles asphalt, are produced at the same time.¹⁴

⁸¹ Valfy and Lucas, English potent 20,470 of 1913 and 2,838 of 1914; J. Soc. Chem. Ind., 34, 71 (1915).

²⁵ ZELINSKI, J. Russian Phys. Chem. Soc., 47, 1807 (1915).

McAfee, J. Ind. Eng. Chem., 7, 737 (1915).

²⁴ PICTET and LERCEYNSKA, Bull. Soc. Chim. (4), 19 (1916).

SUPPLEMENT TO CHAPTERS XI AND XII HYDROGENATION OF LIQUID FATS

937. The liquid fats, oils of various origins, contain along with the neutral glycerine esters of the saturated acids, $(C_nH_{2n}O_2)$, palmitic, margaric, stearic, arachidic etc., a considerable proportion of the glycerine esters of the unsaturated acids, either ethylenic acids, $(C_nH_{2n-2}O_2)$, hypogaic, oleic, elaidic, erucic etc., or diethylenic, $(CH_{2n-4}O_2)$ as linoleic, or unsaturated hydroxy as ricinoleic, or still more unsaturated acids as linolenic, $C_{12}H_{20}O_2$ and clupadonic, $C_{12}H_{22}O_2$. The unsaturated acids and their glycerine esters have much lower melting points than the corresponding saturated compounds, thus:

Stearic acid,	$C_{18}H_{36}O_{2}$			melts at	71°
Oleic acid,	$C_{18}H_{34}O_{2}$			melts at	14°
Ricinoleic acid,	C18H24O2			melts at	26°
Linoleic acid.	C18H209			melts below	-18°.

Stearine, or glycerine tristearate, melts at 71.5° while oleine, or glycerine trioleate, is liquid at the ordinary temperature. In some cases these unsaturated compounds have disagreeable odors. The presence of clupadonic acid is responsible for the repulsive odor of fish oils.

938. The absorption of *iodine* by fats gives an exact measure of the amounts of unsaturated acids that enter into their constitution.

By the term iodine number we mean the amount of iodine absorbed by 100 parts of the fat.¹

The following table gives the average value of the iodine number for a number of different fats:

Cocoa butter									36	
Mutton or bee	f to	rolla	V						35 to	47
Hog lard									44 to	70
Goose fat									77	
Olive oil									82	
Colza oil	_		_	_					100	

¹ Höbl, Dingler's Polytech. J., 253, 281 (1884).

Almond oil							98
Castor oil							84
Peanut oil							97
Cottonseed oil .							109
Sesame oil							108
Poppy seed oil .							133 to 158
Whale oil							127
Cod liver oil							140 to 180
Linseed oil							180
Clupadonic acid							365 to 370.

939. By adding hydrogen and thereby transforming unsaturated glycerides into saturated, the bad odors of certain oils (fish and co-coanut) disappear and the melting points are greatly raised.

When applied to the oils themselves, hydrogenation changes them into solid fats, i.e. more and more solid the more the *oleine* is transformed into *stearine*.

Sabatier and Senderens showed between 1897 and 1902 that hydrogen is easily added to ethylene bonds in the presence of reduced nickel at temperatures below 250° and it was desired to apply this method to the hydrogenation of the liquid fatty acids or to the oils themselves. It is possible by dragging the vapors of oleic acid by a violent current of hydrogen over nickel at 280° to transform it completely into stearic acid. A column of reduced copper can effect the same reaction at 300° and in this case the hydrogen may be replaced by water gas.²

In the patents of Bedford, the fatty material vaporized in hydrogen under reduced pressure is hydrogenated while traversing a vertical cylinder filled with nickeled pumice heated to 200°, but the lowering of the pressure of the gas is unfavorable to its fixation.

940. The difficulty of volatilizing the liquid fatty acids and the practical impossibility of volatilizing the oils themselves led to the abandonment of the reaction on the vapors and to attempts to effect it in the liquid material. The patent of Norman of 1903 comprehended hydrogenation of the vapor and of the liquid.⁴

It is to hydrogenation in the liquid medium that almost all of the very numerous patents applying to this important industrial problem relate: more than 200 have been taken out on the choice, preparation and method of using catalysts or for apparatus.

² Sabatier, French patent, 394,957 (1907).

³ BEDFORD and WILLIAMS, English patent, 9,112 of 1908. — BEDFORD, U. S. patent, 949,954 (1910).

⁴ NORMAN, English patent, 1,515 of 1903.

941. Catalysts. Nickel is most frequently employed being used alone in the finely divided state as is obtained by the reduction of the oxide with hydrogen, or more commonly incorporated with an inert material intended to disseminate it and to increase the useful surface of contact with the hydrogen and oil. For this purpose have been used nickeled pumice, kieselguhr, or infusorial earth, impregnated with nickel, nickeled asbestos, and wood charcoal impregnated with nickel. The method of incorporating the nickel with the carrier may vary: for example, the nickel is dissolved in sulphuric acid and double its weight of siliceous material is added (pumice, kaolin, asbestos etc.); the metal is precipitated as the carbonate which is calcined to form the oxide which is thus distributed over every fragment of the porous material, and the oxide is reduced by hydrogen at 350°.9

942. It has been suggested to use the finely divided nickel formed in the decomposition of nickel carbonyl by heat. Finely divided nickel is kept suspended in oil at above 180° and a current of carbon monoxide or of water gas is passed through. This transforms the metal into nickel carbonyl which immediately breaks down into carbon dioxide, carbon and finely divided nickel which remains suspended in the liquid and is ready to realize its hydrogenation at a somewhat higher temperature, around 220–40°. Practically the nickel thus formed may be only 0.1% of the oil to be hydrogenated.

It has been proposed to impregnate pumice or kieselguhr with nickel carbonyl and then heat it to liberate the metal which should be perfectly spread over the porous material: the catalyst thus prepared is incorporated in the oil to be treated without coming in contact with the air.¹¹

943. The substitution of *nickel oxides* for metallic nickel has put over against existing patents, other patents which could not be of value if the oxide does not act until after it has been reduced to the metal, as various investigations seem to have established (598).

NORMAN, English patent, 1,515 of 1903. — BEDFORD and WILLIAMS, English patent, 9,142 of 1908. — ERDMANN, German patents, 211,669, C., 1910 (1),1908, and 222,890 (1907), C. A., 4, 2715 (1910).

⁶ KAYSEE, U. S. patents, 1,004,035 and 1,008,474 (1911), J. S. C. I., 30, 1266 and 1461 (1911). —WILBUSCHEWITCH, French patent, 426,343 (1910), J. S. C. I., 30,966 (1911). — CROSSFIELD and MARKEL, French patent, 435,249 (1911), J. S. C.I., 3,346 (1912).

⁷ Schwoerer, German patent, 199,909 (1906)).

ELLIS, U. S. patent, 1,060,673, (1913), C. A., 7, 2132 (1913). — ITTNER, Mat. grasses. 1918, 4964.

[•] WILBUSCHEWITCH, English patent, 15,439 of 1911, J. S. C. I., 30, 1170 (1911)

¹⁰ Shukoff, German patent, 241,823 (1910), C., 1912 (1), 175.

¹¹ SCHICHT, Mat. grasses, 1916, 4634.

This substitution, inspired by the work of Ipatief (584), has been advised by Bedford and Erdman, who believe that the most active catalyst is a sub-oxide such as N₂O,¹² and has been frequently applied to the hydrogenation of oils,¹³ 14

944. Various salts of nickel have been proposed to replace the oxide as catalyst. Nickel *borate* recommended by Schönfeld ¹⁸ as being very active, has been found by other chemists absolutely useless unless broken down to the oxide by a temperature of above 260°, the presence of the boric acid appearing to be unfavorable. ¹⁶

The nickel salts of organic acids, acetate, lactate, and especially the formate have shown themselves useful.¹⁷ The product produced by heating nickel formate in a current of nitrogen has been advised.¹⁸

- 945. The other common metals near to nickel, *iron*, *cobalt* and *copper* have been rarely used although they figure along with nickel in a large number of patents.¹⁹ The same is true of *platinum* on account of its high price which is not compensated for by any special activity.
- 946. Palladium on the contrary, has been recommended as a catalyst for oils in spite of its high cost because of its remarkable activity, 1 part of metal effecting the hydrogenation of 10,000 ²⁰ parts of oil below 100°. It is advantageously employed at about 80° under 2 or 4 atmospheres of hydrogen. ²¹ It is best to use the palladium precipitated on an inert carrier, either animal charcoal or a metallic oxide or
 - ¹³ Bedford and Erdmann, J. prakt. Chem. (2), 87, 425 (1913).
- ¹³ BEDFORD and WILLIAMS, French patents, 418,355 (1910); 436,295 (1911); J. S. C. I., 31, 444 (1912). English patent, 29,612 of 1910, J. S. C. I., 31, 398 (1912). U. S. patent, 1,026,339 (1912), J. S. C. I., 31, 593 (1912).— BEDFORD and ERDMANN, French patent, 451,155 (1912),—J. S. C. I., 32, 602 (1913).
- 14 The exact comparative experiments of WILLSTATTER and WALDSCHMIDT-LETTZ (Berichte, 54, 131, (1921)) go far towards proving that nickel is entirely inactive unless it contains some oxygen. Using 0.2 g. nickel in 20 cc. water with 1 g. sodium cinnamate, no hydrogen was taken up in 1 hour at 60° but the catalyst was activated by shaking with oxygen for 15 minutes. A number of similar experiments are cited. A quantitative experiment showed that the amount of oxygen absorbed by a sample of nickel, exposed to the air, was not weighable yet the nickel was activated by this exposure. E. E. R.
 - SCHÖNFELD, Zeit. f. angew. Chem., 27 (2), 601 (1914), C. A., 8, 3868 (1914).
 ERDMANN and RACK, Zeit. f. angew. Chem., 28, 220 (1915), C. A., 9, 1255
- (1915).

 17 Winder and Higgins, French patent, 441,097 (1912), J. S. C. I., 31, 826 (1912).
 - 18 HIGGINS, Mat. grasses, 1917, 4760.
- ¹⁰ NORMAN, English patent, 1,515 of 1903, J. S. C. I., 23, 26 (1904). WIL-BUSCHEWITCH, French patent, 426,343 (1910), J. S. C. I., 30, 966 (1911).
- ²⁰ Hydrogenation of cottonseed oil may be carried on at 180° with this proportion of nickel on a carrier. E. E. R.
 - ²¹ DAY, U. S. patent, 826,089 (1906), J. S. C. I., 25, 1035 (1906).

carbonate or magnesium or nickel, the use for this purpose of iron, lead, zinc or aluminum being unfavorable.22

The chief disadvantage in the use of palladium is its excessive cost since some loss of metal is inevitable, the cost according to experience amounting to 1.60 francs per 100 kg. oil treated.

947. Life of Catalysts. Certain substances when found in even small amounts in the oils, paralyze the activity of catalysts and do not take long to render them inactive (112). The most to be feared in the hydrogenation of oils are sulphur compounds.

Hydrogen sulphide immediately renders inactive 100 times its weight of nickel and pulverized sulphur is half as effective. The action is less rapid with the same proportion of sodium sulphide. On the contrary, sulphates, sodium nitrate, and nickel chloride have no harmful effect. Free chlorine kills the nickel instantly.^{22 24}

In contact with fish oil and whale oil the catalyst is quickly killed; the toxic material is fixed by the metal since if a fresh catalyst is added hydrogenation takes place. Consequently a practical method of purification of these oils is to agitate them with a spent catalyst which abstracts the harmful substances.

948. Oils frequently contain free fatty acids which attack the nickel to the detriment of its catalytic activity. Hence it is best to neutralize them by agitation with pulverized calcium carbonate or with a small amount of dilute cold sodium carbonate solution. The neutral oil thus obtained may be effectively freed from its toxic materials by agitating it hot with freshly precipitated cupric hydroxide.²⁵

949. The presence of moisture in the oil or in the hydrogen can lead to a certain amount of saponification at the elevated temperature at which the reaction is carried out, hence it is important to avoid the presence of water and to dry the gas before using it, e.g., by refrigeration to $-20^{\circ}.26$

- ²² VERBINIGTE CHEM. WERKE, German patent, 236,488 (1910), C. A., 5, 3633 (1911). French patents, 427,729 and 434,927 (1911), J. S. C. I., 30, 1022 and 31,346 (1912). English patent, 18,642 of 1911, C. A., 7, 555 (1913).
 - MOORE, RICHTER, and ARSDEL, J. Ind. Eng. Chem., 9, 451 (1917).
- ²⁴ It is suggested by Willstätter and Waldschmidt-Leftz (Berichte, 54, 127. (1921), that the poisoning of catalysts, by certain substances, at least, may be due to the fact that they deprive the catalysts of their oxygen content and thereby render them inactive. They show that thiophene removes the oxygen from platinum black. In an experiment in which 1.9 g. bensene was being hydrogenated in acetic acid solution by means of 0.5 g. platinum black, 1.5 mg. thiophene was added whereby the activity of the catalyst was completely destroyed. The catalyst recovered 87% of its original activity by treatment with oxygen for 2 hours. E. E. R.
 - ELLIS and WELLS, Mat. grasses, 1917, 4760.
 - 24 Soc. DE STEARIN. ET SAVON. DE LYON, French patent, 485,414 (1917).

950. Nickel catalysts that have been rendered inactive by long use are regenerated by degreasing and treating with nitric acid and calcining the nitrated material thus obtained.

951. Amount of Catalyst. The weight of catalyst can vary much according to the work it has to accomplish. The rapidity of the reaction is nearly proportional to the amount of catalyst used. It is usually best not to cut down the amount of the catalyst since it is convenient to shorten the time of the hydrogenation as much as possible. Usually 2 or 3% of nickel distributed on an inert material is employed. With palladium the amount of catalyst may be much smaller.*

952. Temperatures. The temperatures most frequently employed for hydrogenations with nickel are around 180° but may sometimes be as low as 150° and are frequently raised to 200–50° especially when the oxide is used.

Much lower temperatures suffice with palladium, usually 80 to 100°.

Elevation of temperature increases the speed of the reaction greatly. In the neighborhood of 170–80° with nickel, raising the temperature 10° increases the velocity about 20%.28

953. Hydrogen. The hydrogen for hydrogenations may be prepared electrolytically or may be produced as a by-product in the manufacture of caustic soda.

It can also be obtained by the action of steam on incandescent coke, the water gas thus formed, after absorption of the carbon dioxide, being partially liquefied to eliminate the carbon monoxide. But it is more frequently prepared by the decomposition of water by metallic iron, the iron oxide so formed being reduced at a red heat by water gas.

This decomposition can take place at a red heat alternating with the reduction of the iron oxide which is thus formed; but under these conditions it is difficult to avoid the presence of a certain amount of carbon monoxide which it is important not to admit in appreciable amount.

Amounts of 0.25 to 2% of carbon monoxide produce a serious diminution in the activity of the catalyst; 6 or 8% prevent any hydrogenation by the nickel either by forming a deposit of carbon which covers the catalyst (614), or because the carbon monoxide turns the catalytic activity of the nickel to its own use in transforming itself

²⁷ Good results on cottonseed oil may be obtained with 0.1% of nickel distributed on 10 parts of carrier.— E. E. R.

28 MOORE, RICHTER and VAN ARSDEL, J. Ind. Eng. Chem., 9, 451 (1917). — Mat. grasses, 1918, 5018.

to methane.²⁹ This toxicity of carbon monoxide is all the more peculiar since *nickel carbonyl* has no harmful effect when it is sent into the oil with the hydrogen, even in large amount, and since the nickel resulting from its decomposition is, up to a certain limit, superior to reduced nickel.²⁰

The result is that water gas containing about equal volumes of hydrogen and carbon monoxide with a little carbon dioxide and nitrogen, which can give good results with copper as a catalyst (515), is proscribed in the hydrogenation of oils over nickel.

954. According to Bergius, the formation of hydrogen by water and iron can be very advantageously carried out by operating with water kept in the *liquid* form by high pressures. By working under 300 atmospheres at 300-40°, the reaction:

$$3\text{Fe} + 4\text{H}_2\text{O} = 4\text{H}_2 + \text{Fe}_2\text{O}_4$$

takes place completely and can be greatly accelerated by the presence of sodium chloride or *ferrous chloride* along with metallic copper. Under exactly the same experimental conditions, the amounts of hydrogen evolved per hour were:

Iron and water alone	at 300°			230 cc.
Iron, water and FeCl ₂	at 300°			1390
Iron, water, FeCl ₂ and Cu	at 300°			1930
Iron, water, FeCl. and Cu	at 340°			3450

An apparatus holding 45 l. can produce 102 cu. m. per day. The iron oxide obtained is in fine powder and easy to reduce to metallic iron by water gas.

The hydrogen thus prepared is very pure since the iron carbides and sulphides which are in the iron are not attacked by liquid water. The hydrogen evolved under a pressure of 300 atmospheres can be stored in steel cylinders without further compression.³¹

955. The volume of hydrogen required for hydrogenation varies with the proportion and nature of the unsaturated acids which enter into the composition of the oils.

For pure *oletc acid* about 79 cu. m. per 100 k. of acid are required while *linoletc* acid requires twice and *clupadonic acid* four times this amount.

²⁹ MAXTED, Trans. Faraday Soc., 13, 36 (1918).

³⁰ MAXTED, Trans. Faraday Soc., 13, 201 (1918).

²¹ Brigius, J. Soc. Chem. Ind., 32, 463 (1913). — German patents, 254,593 and 286,961.

The corresponding glycerides require somewhat less, about 76 cu. m. for oleine. The amount required by an oil is proportional to its iodine number: linseed oil requires 150 cu. m. per 1000 k.

956. Pressure. It is advantageous to operate under pressures higher than atmospheric, the velocity of the hydrogenation being, at least up to a certain limit, proportional to the pressure of the hydrogen. In practice, pressures of 2 to 15 atmospheres are used.

957. Apparatus. A large number of forms of apparatus, many of which differ only in details, have been devised for the hydrogenation of oils. Contrary to the general impression, it is not necessary to agitate the oil and the catalyst very violently with the hydrogen; the agitation should especially have the effect of replacing hydrogenated portions of the oil in contact with the catalyst by portions not yet acted on.

The various forms of apparatus may be divided into four distinct types:

958. First Type. The oil and hydrogen are simultaneously projected on to a catalytic surface.

This is the principle of the apparatus of Erdmann which is composed of a vertical nickel cylinder in the centre of which a vertical concentric terra cotta cylinder covered with a layer of catalyst with a nickel base, turns slowly; the apparatus is heated to 180° and the oil driven by compressed hydrogen is projected onto the surface of the cylinder and runs down after it is acted upon.²⁵

959. In the apparatus of Schwoerer designed to hydrogenate oleic acid, the acid carried along by steam superheated to 250–70° and mixed with hydrogen is projected on to a helicoidal surface covered with nickeled asbestos.²⁴

960. The apparatus of Schlinck belongs in this class; it is composed of a centrifuge which turns on a vertical axis in a closed cylinder at the top of which oil and compressed hydrogen are introduced together. The basket of the centrifuge is furnished with asbestos impregnated with catalyst (specially palladium). The oil on account of the rapid rotation, the hydrogen on account of its pressure traverse

ss The volume of hydrogen required to saturate any oil is readily calculated from its iodine number. Thus 1 K of oil whose iodine number is 1, requires 882.0 cc., or 1000 K requires 0.8820 cu.m., of hydrogen measured at 0° C. and 760 mm. Hence multiply these figures by the iodine number of the oil in question. 1 K cottonseed oil with iodine number 110 would take 97.02 1. to saturate it completely or 35.28 1. of hydrogen to bring it down to an iodine number of 70. — E. E. R.

^{*} ERDMANN, German patent, 211,669 (1907), C. A., 3, 2732.

²⁴ Schwoerer, German patent, 199,909 (1906).

this layer simultaneously and partially combine. The partially hydrogenated oil runs out at the bottom; the partially expanded hydrogen passes out at the side of the top of the cylinder and is recompressed to be sent into another cylinder along with the partially treated oil. After passing through a sufficient number of cylinders exactly alike the oil is completely hydrogenated.²⁵

961. Second Type. The oil mixed with the catalyst is atomized in an atmosphere of hydrogen which is kept at a suitable temperature by steam heat.

The apparatus of Wilbuschewitch which seems to have given good results belongs here. It is composed of an autoclave in the form of an elongated vertical cylinder the lower end of which terminates in a 60° cone which is kept at 160°. The oil to which the pulverulent catalyst has previously been added and which is kept mixed by a suitable rotating apparatus, is atomized at the top of the cylinder where the falling droplets encounter an ascending current of hydrogen. This enters at the lower tip of the cone through a circular chamber the top of which is perforated with holes, passes through the oil which has accumlated in the cone, then up the cylinder where it encounters the droplets of oil with the catalyst and passes out at the top of the cylinder to be used again. The partially hydrogenated oil which accumulates in the cone is sent with the catalyst which it carries into a second autoclave like the first where the hydrogenation is carried further and so on into other cylinders till the desired hydrogenation is obtained.³⁶

962. Third Type. The hydrogen is atomized into oil holding the catalyst in suspension and heated to a known temperature.

This is the principle of one form of apparatus of Ellis, which consists of a conical heating vessel with vertical axis having its apex at the bottom and heated by circulation of high pressure steam in a double jacket. It is filled with oil for two thirds of its height. The catalyst is added through a hole in the top and the hydrogen admitted at the desired pressure is circulated by means of a pump, being drawn from the top and forced in at the bottom of the cone rising through the oil which it agitates and which it hydrogenates thanks to the catalyst which is suspended in it. The passage of the gas is continued till the desired degree of hydrogenation is attained. At this moment a horizontal circular filtering disc which is operated by a rod which occupies the axis of the cone, is lowered till it rests on the walls of the cone near the apex. The oil is filtered through this disc leaving the

²⁶ SCHLINCK, German patent, 252,320 (1911), C. A., 7, 910 (1913). — English patent, 8,147 of 1911, C. A., 6, 2858 (1912).

^{**} WILBUSCHEWITCH, French patent, 426,343 (1910), J. S. C. I., 30, 966 (1911).

— English patent, 30,014 of 1910, J. S. C. I., 31, 443 (1912).

catalyst. The apparatus can be charged with a fresh portion of oil which takes up the same catalyst.*7

963. Fourth Type. A vigorous agitation brings the oil, catalyst and hydrogen together in the same vessel.

Kayser's apparatus consists of an autoclave heated to 150-60° and filled one fourth full of oil mixed with a pulverulent nickel catalyst under hydrogen introduced at the desired pressure. An agitator consisting of six vanes of metal cloth mounted on a metal frame perpendicular to a horizontal axis, can revolve rapidly and thus cause an intimate mixture of gas, oil and solid catalyst.²⁸

964. The apparatus of Kimura is very similar.³⁹ In the apparatus of Woltmann the agitator rotates on a horizontal axis and carries perforated arms through which the hydrogen is sent in under pressure corresponding to the rate of its fixation by the oil.⁴⁰

965. Results. The hydrogenation of oil is carried on in quite a large number of plants, more than 24 in 1916.

It enables us to obtain from oils of very inferior quality, such as whale oil, fatty materials with odors that are not disagreeable, possessing a remarkable consistence along with high melting points. A regulated hydrogenation enables us to prepare at will products intermediate between the oils and the solid fats.

The fixation of 1% by weight of hydrogen is sufficient to transform cottonseed oil and other oils of that class into substances with the consistency of lard. This result may be attained directly by means of hydrogenation of the whole mass of the oil and stopped at the desired hardness, the operation being carried on at as low a temperature as possible so as not to alter the qualities of the oil. But the desired end can be more surely attained by hydrogenating a portion of the oil to the limit and then mixing this with untreated oil to obtain the desired hardness.

966. The table below gives the melting points of the fats obtained by complete hydrogenation of the oils or fats.⁴¹

⁸⁷ Ellis, J. Soc. Chem. Ind., 31, 1155 (1912).

^{**} KAYSER, U. S. patente, 1,004,035 and 1,008,474 (1911), J. S. C. I., 30, 1266 and 1461 (1911).

⁸⁸ KIMURA, French patent, 486,621 (1918).

⁴⁰ WOLTMAN, English patent, 112,293 (1916), C. A., 12, 1006 (1918).

a Mannich and Thiele, Mat. grasses, 1917, 4676.

					Melting point	ats
					Hydrogenated oil	Original
Olive oil .					70°	6°
Almond oil					72°	-10°
Peanut oil				,	64.5°	- 3
Sesame oil					63.5°	-5°
Poppy seed	oi	1			70.5°	-18°
Linseed oil					68°	-16°
Codliver oil					68°	-10°
Cocoa butte	r				64°	23°
Tallow					62°	35°
Lard oil .					64°	28°

The iodine number becomes very small in every case.

967. The commercial grades do not correspond to such complete hydrogenation. They exhale a peculiar, very persistent aromatic odor which resists saponification and distillation under reduced pressure.

Such hardened oils are known under the French trade names of duratol, talgol, candelite and synthetic tallow.

Below are given some of the characteristics of such products, melting point, iodine number and melting point of the fatty acids: 42 45

	M.p.	I. No.	M.p. of Acids.
Talgol	35–37°	86.1	38.5°
Talgol extra	42 -44 °	63.9	45.5°
Candelite	48-50°	10.4	48.5°
Candelite extra	51-52°	10.5	51.8°

- 968. Castor oil which does not solidify till below -18°, gives on careful hydrogenation a very white hard solid which melts above 80° and which is advantageously employed as an electric insulator.
- 969. The question of the use of deodorized and hardened oils as fats in food has not been completely settled as yet, because we are not altogether certain about the toxicity of the small amounts of nickel which remain in the materials, amounts that are hardly more than 0.000002% if the oils treated were entirely neutral.
 - ⁴⁶ Gärth, Seif. Zeit., 39, 1277 (1912).
- on in the United States by about 60 concerns and hydrogenated oils are important constituents in some 92 brands of shortening. Formerly these were regarded as lard substitutes but they have made an independent position for themselves as "vegetable shortening" and have found favor with many who object to lard.

For edible products cottonseed oil is the chief oil that is hydrogenated. The aim is to prepare a product that will not be too hard in winter or too soft in summer. Sometimes the whole of the oil is hydrogenated. The chief products thus made

for the American market are Crisco, Selex, MFB, and Fairco. These melt at 33 to 37° and have iodine numbers running from 76 to 86. By varying the mode of hydrogenation, products with nearly the same melting points but with iodine numbers varying as much as 10 points may be obtained.

By hydrogenating down to an iodine number of 10 to 20 and mixing this very hard fat with untreated cottonseed oil the desired consistency may be obtained with a much higher average iodine number. This is the most common practice as much less hydrogen is required and the fraction of the oil that has to be hydrogenated is small.

The melting points and iodine numbers of some leading brands are as follows:

	Melting Point	Iodine Number
Scoco	. 44.4	89.2
Snowdrift	. 44.1	89.3
Armstrong White Cloud	. 45.8	99.4
Armstrong Bob White	. 39.2	93.5
Fairbanks Boar's head	. 41.8	100
Morris Purity	. 42.0	97.5
P. and G. Flake white	. 42.8	90.6
P. and G. White flake	. 47.8	87.4
Swift Jewel	. 45.5	97.0
Wilson Advance	. 44.2	96.9
Kream Krisp	. 45.5	97.0

Highly hydrogenated cottonseed oil is a hard, white, brittle solid and does not become rancid. These properties make it a suitable constituent for prepared cake flours.

Vast amounts of fish oils are hydrogenated to be used in making soaps.

E. E. R.

AUTHOR INDEX

(References are to Paragraphs: a number followed by "n" designates a note.)

Abakumobskava. Miss L. N., with Nametkin, 478 Aboulenc, Jean, with Senderens, 598, 758, 759, 760 Acree, S. F., and Johnson, J. M., 202 Adam, Paul, 241, 896 Adams, Roger; Kamm, O., and Marvel. C. S., 306, 696, 713 Adkins, Homer, 797n, 861n Ador, E., with Rilliet, 291 Adrianowsky, 297 Akunoff, J., with Lunge, 445 Alexyef, 899 Allen, William, and Kölliker, Alfred, 890 Almedingen, 212 Aloy, J., and Brustier, 385, 661 Altmayer with Mayer, 411 Amberger, C., with Paal, 69, 70, 545 Ambrey, A., with Bourquelot, 18 Amouroux, G., 385, 435 Amouroux, G., with Mailhe, 739 Amouroux, G., with Murat, 414, 415 Andrews, C. E., with Boehner, 811 Anschütz, Richard, 885, 889, 897 Anschütz, R., and Immendorff, H., 888 Antropoff, A. von, 180a Antropoff, A. von, with Bredig, 180a Arbusof, A. E., and Friauf, A. P., 633 Arbusof, A. E., and Khrutzkii, N. E., 633 Arbusof, A. E., and Tichwinsky, W. M., 611, 633, 635 Armstrong, H. E., and Tilden, W. A., 198 Aronheim, B., 286 Arrhenius, Svante, 178, 319, 324 Asahina, Y., 571 Aschan, Ossian, 929 Atkinson, R. H., Heycock, C. T., and Pope, W. J., 282n Auger, Victor, 893 Auger, V., and Behal, A., 280 Austerweil, Geza, 260n

Baborovsky, G., and Kusma, B., 276 Badische, A. S. Fb., 180s, 215, 270, 273, 511, 730, 876 Backeland, Leo H., 792 Baeyer, Adolf, 90, 893 Baeyer, Adolf, and Drewsen, Viggo, 798 Baker, H. B., 73 Balv, E. C. C., and Krulla, Rudolf, 180 Bancroft, W. D., 116n, 180a, 180g, 180s Barbaglia, G. A., 224 Barbier, Ph., and Locquin, R., 565 Bardt, A. Y., with Doroshevskii, 268, Barendrecht, H. P., 180i Bartels, A., with Jannasch, 817 Bartels, G., with Meigen, 598 Bauer, A., 211 Bauer, Maurice, with Brochet, 601 Baumann, E., 150, 233 Bayer, A. G., 232 Bayer and Co., 104, 298 Bayley, 180a Becker, C., with Semmler, 570 Beckmann, E., 185, 189 Bedford, Fred, 939 Bedford, F., and Erdmann, E., 598, 943 Bedford, F., and Williams, C. E., 939, 941. 943 Béhal, A., 192, 308 Béhal, A., with Auger, 280 Behn, Richard, 892 Beilstein, F., and Geitner, P., 278 Beilstein, F., and Kuhlberg, A., 287 Belloni, E., with Carrasco, 250 Bergen, J. von, with Skita, 554 Bergius, Friedrich, 954 Bergreen, Henry, 894 Berl, E., 180r Berliner, 1800 Bernthsen, August, 296 Berthelot, Marcellin, 21, 60, 84, 148, 165,

160, 180q, 308, 325, 340, 409, 477, 616, 637, 650, 749, 750n, 751, 752, 757, 758, 767, 768, 770, 822, 905, 909, 914, 926 Berthelot and Jungfleisch, 650, 749 Berthelot and St. Gilles, 758n Bertrand, Gab., 153, 264 Berselius, J., 4, 129 Bevan, E. J., with Cross, 268 Bialobi zeski, M., 900 Biehler, F., with Paal, 70 Blackadder, Thomas, 822 Blaise, 304 Blanck, F. C., with Tingle, 269n Blanes, J. S., with Madinaveitia, 569, Blanksma, J. J., with van Ekenstein, 186 Bodenstein, Max. 8 Bodenstein, Max, and Fink, Colin G., Bodländer, G., Koppen, K., 180r Bodroux, F., 751, 752, 757 Bodroux, F., and Taboury, F., 420 Boedtker, Eyrind, 819 Boedtker, Eyrind, and Halse, O. M., 888 Boehner, R. S., and Andrews, C. E., 811 Boehner, R. S., and Ward, A. L., 811 Böeseken, J., 81, 87, 224, 643 Böeseken, J., and Schimmel, A., 224 Böeseken, J., van der Scheer, J., and de Vogt, J. G., 879 Böeseken, J., and van Senden. G. H.. 664 Böeseken, J., van de Weide, O. B., and Mom, C. P., 117, 546 Boessneck, P., 89 Boettger, B., 62, 165 Bone, W. A., and Jerdan, D. S., 409 Bornwater, J. T., and Holleman, A. F., 284 Borodin, A., 795 Borsche, W., and Heimbürger, G., 546 Borsche, W., and Wollemann, J., 546 Bösler, M., 220 Böters, O., with Wolffenstein, 269 Bouchardat, G., 212 Boudet, F., 184 Boudouard, O., 615 Bougault, J., 203 Boullay, J. F. G., 691 Bourquelot, Em., and Aubry, A., 18 Bouveault, L., 654, 656, 663, 717 Bouveault, L., and Locquin, René, 663

Boyd, Robert, with Henderson, 459 Braun, J. von. and Deutsch, H., 897 Braune, H., 12 Bredig, G., 68 Bredig, G., and V. Antropff, 180a Bredig, G., and Brown, John Wesley, 272 Bredig, G., and Carter, S. R., 574 Bredig, G., and Fraenkel, W., 12 Bredig, G., and Ikeda, K., 116 Bredig, G., and Joyner, R. A., 836 Breteau, Pierre, 484, 536, 562, 571, 579 Breuer, Aug., and Zincke, Th., 220 Brochet, André, 30, 596, 598, 599, 600, 603 Brochet, André, and Bauer, Maurice, 601 Brochet, André, and Cabaret, André, 601, 602 Bromberg, O., with Fischer, 187 Brooks, Beni. T., and Humphrey, Irwin, 210, 306 Brown, J. W., with Bredig, 272 Brown, O. W., and Carrick, L. L., 512 Bruce, James, with Willstätter, 293, 473 Brunel, L., 126, 349, 443, 459, 464 Bruner, L., 291 Brunner, W., with Skita, 561 Brustier, V., with Aloy, 385 Bugarszky, Stefan, 178 Bum, Friedrich, with Kohn, 293 Bunsen, R., 180r Burrows, George J., 324 Burstert, H., with Claus, 285 Burstyn, Walther, 234 Butlerow, A., 210, 306

Cabaret, André, with Brochet, 601, 602 Caldwell, G. C., and Grössmann, A., 184 Calm, A., 89 Calvert, F. C., 48, 180b Cannizzaro, S., 880 Carpenter, C. C., 372 Carrasco, O., and Belloni, E., 250 Carrasco, O., and Padoa, M., 497, 684 Carrick, L. L., with Brown, 512 Carter, S. R., with Bredig, 574 Carughia, A., with Padoa, 489 Cathcart, W. R., Jr., and Meyer, Victor, 893 Chauvin, A. C., 268 Chelintsev, V. V., and Trunov, B. V., 805

Ch. Fab. auf Actien (E. Schering), 215 Chem. Fabr. Buckau, 881 Chiaves, C., with Padoa, 490 Chichibabin, A. E., 310, 686, 807, 810 Chichibabin, A. E., and Ryumshin, P. F., 901 Chowdhuri, T. C., with Neogi, 382 Ciamician, G., 647 Ciamician, G., and Silber, P., 150 Claisen, L., 783, 799, 804 Claisen, L., and Claparède, A., 798 Claisen, L., and Crismer, L., 106 Claisen, L., and Ponder, A. C., 798 Claparède, with Claisen, L., 798 Clark, Latham, and Jones, W. N., 414 Claus, Ad., 207, 893 Claus, Ad., and Burstert, H., 285 Clement and Desormes, 324 Cohen, Ernest, 8 Cohen, J. B., and Dakin, Henry, D., Cohen, Lillian, with Harding, 298 Colin, H., and Sénéchal, A., 11 Colson, Albert, 858 Commercial Research Co., 269n Consortium f. Electroch. Ind., 228 Cooke, Stephen, 166 Copisarow, Maurice, 893 Corenwinder, B., 15, 342 Cornubert, R., 602 Couroy, James T., 161, 837 Crafts, J. M., 82 Crafts, J. M., with Friedel, 87, 173, 263, 295, 297, 883, 889, 890, 893 Crismer, L., with Claisen, 106 Cross, C., F., Bevan, E. J., and Heiberg, Th., 268 Crossfield, J., and Sons, and Markel, K. E., 941 Curtius, Th., and Foersterling, H. A., 196 Curtius, Th., and Lang, J., 332 Cusmano, Guido, 564, 571

Dakin, H. D., with Cohen, 293
Damoiseau, O., 48, 180b, 282
Daniels, E. A., with Frankforter, 239, 806
Darzens, Georges, 30, 56, 243, 360, 389, 417, 420, 476, 488

Davv. E., 4 Davy, Sir Humphrey, 4 Day, D. T., 946 Deacon, H., 103, 180r Debray, H., with St. Claire-Deville, 64, Debus, Heinrich, 180d, 180f, 342, 528 Delepine, Marcel, 795 Delisle, Alfred, 183 Demole, E., 890 Demtschenko, S., 224 Demuth, R., and Dittrich, M., 893 Deniges, Georges, 268 Dennstedt, M., 742 Dennstedt, M., and Hassler, F., 257 Desormes, with Clement, 324 Deutsch, H., with v. Braun, 897 Deuss, J. J. B., 629 Deussen, Ernst, 560 Deville, St. Clair, 346 Dewar, James, 132, 136, 165 Dey, M. L., with Ray, 815n Dimroth, Otto, and W. von Schmaedel, 816 Dittrich, M., with Demuth, 893 Dits, Hugo, 272 Dixon, Harold B., 73 Doebereiner, J. W., 62 Doroshevskii, A. G., and Bardt, A. Y., 268, 275 Douris, Roger, 208, 419 Dovgelevich, N., with Ipatief, 920 Downes, Helen R., with Reimer, 340n Downs, C. R., with Weiss, 260n Drachussow, with Ipatief, 594 Drewsen, Viggo, with Baeyer, 798 Drevfus, Henri, 255, 261, 309 Douris, Roger, 487 Dubois, H., with Müller, A., 285 Dubrumfaut, 188 Ducellies, F., Gay, L., and Raynaud, A., Duclaux, Jacques, 139 Dulk, L., with Meyer, 224 Dulong, and Thénard, 637 Dumas, J., and Péligot, E., 691 Dupont, Georges, 195, 565, 577 Earle, R. B., and Kyriakides, L. P., to

Hood Rubber Co., 802

Earle, R. B., with Kyriakides, 723

Easterfield, T. H., and Taylor, Miss Clara, M., 843 Egloff, Gustav, and Moore, Robert J., 909 Egloff, Gustav, and Twomey, T. J., 908 Egloff, Gustav, with Zanetti, 907 Eijkman, J. F., 392, 452, 454, 474 Ekl, Elizabeth, with Klemenc, 269n Elbs, Karl, 605 Ellis, Carleton, 941, 962 Ellis, C., and Rabinovitz, Louis, 601 Ellis, C., and Wells, A. A., 949 Engel, R., and de Girard, 780 Engelder, C. J., 108q, 694, 708, 710 Engler, C., 150 Engler, C., and Wild, W., 150 Engler, C., and Wöhler, Lothar, 137, 154 Enklaar, C. J., 415, 416 Erdmann, E. O., 188, 598, 754, 941, 958 Erdmann, E., with Bedford, 598, 943 Erdmann, E., and Rack, E., 944 Erlenmeyer, Emil, 321, 696 Erlenmeyer, E. Jr., 203 Espil, Léo, with Sabatier, 12, 16, 56, 80, 113, 114, 125, 338, 346, 358, 492, 598 Euler, A., with Euler, H., 225 Euler, Hans, 324 Euler, H., and Euler, A., 225 Evans, E. V., 372 Evans, P. N., and Sutton, Lena M., 691 Fabinyi, R., 90 Fabris, Ugo, with Padoa, 484, 491, 642, 643 Fahlberg, List and Co., 285 Faillebin, with Vavon, 565 Fajans, Kasimir, 836 Faraday, Michael, 1800 Farbw. Meister, Lucius, and Brüning, Farbf., v. F. Bayer and Co., 816 Fassek, W., 795 Farre, P. A., and Silbermann, J. T., 131

Faworsky, Al., 192

268

Fenton, H. J. H., 268

Filippov, O. G., 577

Fenton, H. J. H., and Jackson, Henry,

Fenton, H. J. H., and Jones, H. O., 268

Filippov, O. G., with Ipatief, 589, 590

Fink, C. G., with Bodenstine, 180r

Fischer, Emil, 81, 187, 220, 754, 756, 758 Fischer, E., and Bromberg, O., 187 Fischer, Emil, and Fischer, O., 890 Fischer, E., and Giebe, Georg, 782 Fischer, E., and von Mechel, Lucas, 793 Fischer, E., and Morell, R. S., 187 Fischer, E., and Passmore, F., 221 Fischer, E., and Piloty, O., 187 Fischer, E., and Speier, Arthur, 753 Fischer, E., and Tafel, J., 237 Fischer, Ernst, with Schmidt, 571 Fischer, Franz, 931 Fischer, Franz, and Niggemann, Hermann 930 Fischer, O., with Fischer, E., 890 Fischer, O., and Körner, G., 89 Fittig, Rudolph, 11, 183, 203, 293, 797 Fittig, R., and Köhl, Wilhelm, 183 Fittig, R., and Langworthy, C., F., 183 Fleitmann, Th., 270 Foesterling, H. A., with Curtius, 196 Fokin, S., 12, 252, 254, 266, 526, 556, 562, 587 Formin, W., with Tchougaeff, 570 Fossek, W., 224 Fournier, H., 567 Fownes, G., 194 Fraenkel, W., with Bredig, 12 Franchimont, A. P. N., 761 Franke, Adolf, 226 Franke, Adolf, and Kohn, Leopold, 234 Franke, Adolf, and Kohn, Morits, 227 Franke, Adolf, and Wozelka, Hermann, 223 Frankforter, G. B., and Daniels, E. A., 239, 806 Frankforter, G. B., and Kokatnur, V. R., Frankforter, G. B., and Kritchevsky, W., 806 Frankland, E., and Kolbe, H., 232 Freas, Raymond, and Reid, E. Emmet, 758nFrébault, A., 428 Frerichs, G., 598 Freund, Michael, with Sleinkopf, 931 Friauf, A. P., with Arbuzof, 633 Friedel, C., and Crafts, J. M., 87, 173. 263, 295, 297, 883, 889, 890, 893 Friedmann, T. E., with Huston, 728

Haag, J., 233

Gabriel, S., and Neumann, A., 107 Gambier, with Trillat, 781 Gangloff, W. C., and Henderson, W. E., Garrand, S. F., with Weismann, 654n Gärth, Johann, 967 Gattermann, Ludwig, 606, 610, 895 Gattermann, L., with Stockhausen, 886 Gattermann, L., and Koch, J. A., 298 Gaudion, Georges, 382, 429, 513, 741 Gaudion, Georges, with Sabatier, 631, 634, 641, 643, 644, 645, 680, 681, 726, 923 Gay, L., with Ducelliez, 292 Geigy, R., and Koenigs, W., 893 Geilmann, W., with Mannich, 656 Geitner, P., with Beilstein, 278 Genieser, Ad., with Wilgerodt, 238 Genveresse, P., 890 Gerum, J., 576 Gerum, J., with Paal, 72, 546, 556 Geuther, A., 387, 780 Gibbs, H. D., 244n, 249n, 254n, 257n, 260n, 262n, 273n Gibello, with Seyewetz, 221 Giebe, Georg, with Fischer, 782 Girard, de, with Engel, 780 Gladstone, J. H., and Tribe, Alfred, 165, 166, 169, 785 Glinka, N., with Zelinski, 648, 822 Godchot, Marcel, 29, 363, 390, 392, 453, 482, 483 Godchot, Marcel, and Taboury, Félix. 390, 421, 436, 856 Godon, F. de, with Mailhe, 539, 682, 740, 772n, 808, 814 Godon, F. de, with Sabatier, 801 Goldberg, Irma, 901 Goldschmidt, Heinrich, and Larsen, Halfdan, 283, 288 Goldsmith, J. N., 661 Gössmann, A., with Caldwell, 184 Gottlieb, J., 183 Gottlob, Kurt, with Harries, 235 Graebe, C., 272 Graebe, C., and Guye, Ph., 107 Graebe, C., Liebermann, C., 328 Graham, Thomas, 65, 536 Grassi, G., 496 Greene, W. H., with LeBel, 691 Griesheim Elektron, with Johnson, 260

Grignard, Victor, 11, 104
Grigoreff, 702
Grillet, 339
Grimaux, Édouard, 246, 680
Gross, K. F. L., 215
Grube, G., and Krüger, J., 233
Grucarevic, S., and Mers, V., 899
Grün, A., with G. Schicht, Akt.-Ges., 846
Gustavson, G., 174, 199, 290, 293
Guthrie, F., 691
Guttmann, O., with Stock, 8
Guye; Ph., with Graebe, 107
Guyot, A., with Haller, 893

Haarmann, Wilh., with Tiemann, 329 Hall, W. A., 933 Haller, A., 334, 341 Haller, A., and Guyot, A., 893 Haller, A., and Lassieur, A., 435 Haller, A., and Martine, C., 416, 421, 436, 476, 478 Haller and Youssouffian, 341 Halse, O. M., 569 Halse, O. M., with Boedtker, 888 Hamonet, J., 902 Hantzsch, A., 893 Harbeck, E., and Lunge, G., 1800 Harding, E. P., and Cohen, Lillian, 298 Harned, Herbert, S., 180q Harries, C., 213 Harries, C., and Gottlob, Kurt, 235 Hartmann, Wilhelm, with Paal, 1800 Hartwich, Frank, with Wolffenstein, 893 Hassler, F., with Dennstedt, 257 Hatt, Daniel, with Willstätter, 569, 571 Hauser, O., and Klotz, A., 778 Haussknecht, Otto, 184 Hautefeuille, P., 15 Heckel, W., with Knoevenagel, 650, 669, 692, 720 Heidelberger, M., with Willstätter, 571 Heilberg, Th., with Cross, 268 Heimbürger, G., with Borsche, 546 Heinemann, A., 815 Helfrich, O. B., and Reid, E. E., 278n Heller, Gustav, and Schülke, Kurt, 889n, 898 Hemptinne, A. de, 1800

Henderson, G. G., and Boyd, Robert, Henderson, G. G., and Sutherland, Maggie M., 463 Henderson, W. E., with Gangloff, 899 Henrard, J. Th., 418 Henri, Victor, 180j, 180r Henry, L., 236 Henseling, 410 Hersenstein, Anna, with Zelinski, 649 Hess, K., and Liebbrandt, F., 561 Heycock, C. T., with Pope, 282n Hibbert, Harold, 699 Higgins, E. B., 944 Higgins, E. B., with Wimmer, 944 Hobohm, K., with Vorländer, 799 Hoffmann, F., and La Roche and Co., 555 Hofmann, A. W., 63, 232, 287 Hofmann, K. A., 262 Hofmann, K. A., and Schibsted, Helge, Hofmann, K. A., and Schumpelt, K., 271 Hohenegger, C., with Paal, 212, 548 Holdermann, K., 269n Holleman, A. F., with Bornwater, 284 Holtzwart, Rudolf, 231 Hood Rubber Co., with Earle, 802 Hoppe, Eduard, 795 Hoppe-Seyler, F., 150 Houben, J., and Pfau, A., 569 Hübl, Baron, 938 Hübner, H., and Majest, W., 278 Hüfner, G., 180d Hugershoff, A., 630 Humphrey, Irwin, 210, 306 Husemann, Aug., and Marmé, Wilh., 330 Huston, R. C., and Friedmann, T. E., 728 Hutin, Albert, 792

Ikeda, K., with Bredig, 116
Iljinsky, M., 816
Immendorff, H., with Anschütz, 888
Imray, O., from Fbw. Meister, Lucius and Brüning, 228
Ingle, Harry, with Mackey, 266
Ipatief, Vladimir N., 78, 180g, 1807, 190, 211, 232, 542, 543, 584, 585, 586, 587, 588, 589, 590, 591, 592, 593, 594, 595, 598, 667, 670, 694, 702, 706, 711, 714, 717, 722, 724, 943

Ipatief and Dovgelevich, N., 920
Ipatief and Drachussow, 594
Ipatief, V., Jakowlew, W., and Rakitin, W., 592
Ipatief and Leontowitch, W., 200
Ipatief and Louvogoi, 589
Ipatief, V., and Matow, N., 591, 722
Ipatief, V., and Filippov, O., 589, 590
Ipatief, V., and Rutala, O., 211, 714
Ipatief, V., and Schulman, G. G., 838
Ipatief, V., and Sdsitowecky, W., 713
Ittner, Martin H., 54n, 941n

Jackson, Henry, with Fenton, 268

Jacobson, Oscar, 291, 887 Jacquet, D., with Willstätter, 563, 569 Jahn, Hans, 678 Jakowlew, W., with Ipatief, 592 Jannasch, P., and Bartels, A., 817 Jennings, H. S., 121n Jerdan, D. S., with Bone, 409 Joannis, J., 267 Job, André, 153 Johnson, F. M. G., 180g Johnson, G. W., from Griesheim Elek., Johnson, J. M., with Acree, 202 Jonas, K. G., with Semmler, 570 Jones, H. O., with Fenton, 268 Jones, W. N., with Clark, 414 Jorissen, W. P., and Reicher, L. Th., 100 Joyner, R. A., with Bredig, 836 Jungfleisch, Émile, 278 Jungfleisch, with Berthelot, 650, 749

Kametka, T., with Willstätter, 197, 472, 479

Kamm, O., with Adams, and Marvel, 306, 696, 713

Kastle, J. H., and Loevenhart, A. S., 180s

Kaschirski, M., 200

Kawalier, A., 328

Kayser, E. C., 941, 963

Keghel, Maurice de, 481n

Keiser, E. H., with Remsen, 150

Kekulé, Aug., 182, 183, 795, 796

Kekulé, A., and Schrotter, H., 199

Kekulé, A., and Strecker, Otto, 182

Kekulé, A., and Zincke, Th., 222

Kelbasinski, S. S., with Ostromuisslenskii, 784 Kelber, C., 598, 599 Kelber, C., and Schwartz, A., 69, 548 Kempf, R., 276 Kenner, James, with Knoevenagel, 297 Keres, Conrad, 877 Khrutzkii, N. E., with Arbuzof, 633 Kimura, Kanesuke, 964 Kipping, F. Stanley, 799 Kijner, N., 444 King, A. T., and Mason, F. A., 782 King, V. L., with Willstätter, 569, 571 Kjeldahl, 272 Kirchof, 4 Kishner, N., 611, 612 Klemenc, Alfons, and Ekl, Elizabeth, Klever, H. W., with Staudinger, 235 Klots, A., with Hauser, 778 Kluge, Paul, 278 Knoevenagel, E., 240, 296, 632, 729, 790, 804 Knoevenagel, E., and Heckel, W., 650, 669, 692, 720 Knoevenagel, E., and Kenner, J., 297 Koch, Erwin, 278 Koch, J. A., with Gattermann, 298 Koelichen, Karl, 229 Koenigs, W., with Geigy, 893 Koerner, G., and Menozzi, A., 312 Köhl, Wilhelm, with Fittig, 183 Kohn, Leopold, with Franke, 234 Kohn, Morits, 697 Kohn, Moritz, with Franke, 227 Kohn, Moritz, and Bum, Friedrich, 293 Kohn, M., and Müller, N. L., 293 Kokatnur, V. R., with Frankforter, 806 Kolbe, H., with Frankland, 232 Kolbe, H., and Saytzeff, Michael, 165, Kölliker, Alfred, with Allen, 890 Konsortium f. Elektrochemische Industrie, 228 Kopp, Adolph, with Michael, 219 Koppen, K., with Bodländer, 180r Körner, G., with O. Fischer, 89 Koshelev, F. F., with Ostromuislenskii, Köts, A., and Schaeffer, 550 Koursanof, N. J., 889

Krämer, G., and Spilker, A., 217 Kramer, R. L., and Reid, E. E., 707n, 708n, 744n 4015 Catalysis 8-8-10 JM 18 gal 6 Krassuski, K., 200 Kraut, K., 329 Krestinsky, V., and Nikitine, N., 713 Kritchevsky, W., with Frankforter, 806 Krüger, A., 278 Krüger, F., 180j Krüger, J., with Grube, 233 Krüger, Paul, with Tiemann, 198, 800 Krulla, Rudolf, with Baly, 180i Kuhlberg, A., with Bielstein, 287 Kuhlmann, F., 342, 529 Kutscheroff, M., 309 Kuzma, B., with Babarovsky, 276 Kusnetzov, M. I., 623, 920 Kvapishevskii, K. V., with Zal'kind, 548, 566 Kyriakides, L. P., 726 Kyriakides, L. P., and Earle, R. B., 723, 802

Lafont, J., 216 Laming, 146 Landolph, F., 211 Lang, J., with Curtius, 332 Langer, C., with Mond, 614 Langmuir, Irving, 180d, 180e, 180f, 180p Langworthy, C. F., with Fittig, 183 Larsen, Halfdan, with Goldschmidt, 283, 288 Lassieur, A., 435 Lassieur, A., with Haller, 435 Laurent, A., 184 Lasarew, 293 Lebach, H., 792 Le Bel and Greene, W. H., 691 Le Chatelier, 131 Leeds, Albert R., 150 Lehmann, F., 583 Lemonie, Georges, 2, 11, 20, 32, 34, 38, 49, 63, 77, 679 Leontowitch, W., with Ipatief, 200 Leprince, and Siveke, 542 Lerczynska, Miss I., with Pictet, 936 Leroux, Henri, 481 Leroy, A. J., 293 Lescoeur, H., and Rigaut, A., 230 Lespieau, R., 566

Lespieau, R., and Vavon, G., 566 Lewis, W. C. McC., 180j, 180r Lewkowitsch, 314, 318 Libavius, 4 Lieben, Adolf, 104, 222, 321, 795 Liebermann, C., with Graebe, 328 Liebig, Justus von. 11, 312 Liebig, Justus von, with Wöhler, 220 Liebrandt, F., with Hess, 561 Limpricht, H., 278, 320, 851 Lindenbaum, Ernst, with Naumann, 260, 269 Lineburger, C. E., 890 Lipp, Peter, 478 Lippmann, Edmund O. von. 324 Livache, Ach., 266 Lobry de Bruyn, C. A., 186 Locquin, R., with Barbier, 565 Locquin, René, with Bouevault, 663 Loevenhart, A. S., with Kastle, 180s Loew, O., 62, 221, 562, 621 Longman, J., 298 Lorin, 822 Louise; E., 797 Louvogoi, with Ipatief, 589 Löwenherz, Richard, 315 Lucas, O. D., with Valpey, 933 Ludwig, H., 329 Lunge, G., and Akunoff, J., 445 Lunge, G., with Harbeck, 1800

McAfee, A. M., 935 Mackey, W. McD., and Ingle, Harry, Madinaveitia, A., 117, 580 Madinaveitia, A., and Blanes, J. S., 569, Mahl, with Wöhler, 75 Mailhe, Alphonse, 383, 386, 435, 514, 735, 745, 833, 842, 843, 849 Mailhe, Alphonse, with Sabatier, 75, 77, 78, 112, 127, 162, 169, 170, 258, 337, 343, 347, 363, 385, 387, 391, 404, 406, 407, 420, 422, 430, 431, 437, 438, 442, 457, 458, 461, 470, 475, 486, 495, 521, 617, 621, 628, 641, 651, 655, 660, 672, 673, 674, 677, 689, 693, 702, 704, 706, 708, 709, 714, 715, 717, 731, 733, 734, 735, 737, 739, 743, 744, 745, 746, 762, 766, 769, 771, 772, 774, 777, 785, 786, 787, 788, 789, 791, 794, 822, 823, 824,

834, 839, 842, 843, 844, 845, 849, 850, 852, 853, 856, 857, 858, 866, 873, 876, 878, 916, 923 Mailhe, A., and Amouroux, 739 Mailhe, Alph., and de Godon, F., 539, 682, 740, 772n, 808, 814 Mailhe, A., and Murat, Marcel, 384. 385, 494 Majest, W., with Hübner, 278 Mamontoff, W., 691 Mannich, C., 646 Mannich, C., and Geilmann, W., 656 Mannich, C., and Thiele, 966 Mansfield, Johannes, with Scholl, 685 Markaryan, Miss V., with Zal'kind 548, 566 Markel, K. E., with Crossfield, 941 Marmé, Wilh., with Husemann, 330 Martine, C., with Haller, 416, 421, 436, 476, 478 Mason, F. A., with King, 782 Mason, John E., and Wilson, John, 262 Masson, A., 691 Matignon, C., and Trannoy, 75, 259 Matow, N., with Ipatief, 591, 722 Matthews, F. E., and Strange, E. H., 213 Maxted, Edward B., 10, 180o, 954 Mayer, E. W., with Willstätter, 565, 569 Mayer, Max, and Altmayer, 411 Mechel, Lukas von. with E. Fischer. 793 Meigen, W., and Bartels, G., 598 Meissel, M., 899 Meissl, E., 325 Meister, Lucius, and Brüning, Farbw, 261, 299 Meister, Lucius, and Brüning, with Imray, 228 Melsens, 48 Menozzi, A., with Koerner, 312 Menschutkin, N., 38, 768, 769 Mereshovski, B. K., 192, 193, 472 Merz, V., with Grucarevic, 899 Metzger, R., with Schmidt, 454, 484 Meyer, Ernst von, 231, 232 Meyer, Lothar, 294 Meyer, Richard, and Tanzen, August, 683 Meyer, Victor, with Cathcart, 893 Meyer, Victor, and Dulk, L., 224

Meyer, W. A., with Skita, 69, 545, 551, 552, 554, 557, 559, 560 Michael, Arthur, 104, 239 Michael, Arthur, and Kopp, Adolph. Michael, A., Scharf, E., and Voigt, K., 200 Michiels, Louis, 874 Mignonac, Georges, 380, 512, 809 Millar, W. S., 12 Miller, W. Lash, 180a Milligan, C. H., 538n, 706n, 772n, 778n Millon, E., with Reiset, 637 Moeser, Ludwig, with Naumann, 260, 269 Moissan, Henri, 73, 136, 147 Moissan, H., and Moureu, Ch., 637, 914 Mom, C. P., with Böeseken, 117, 546 Mond, Ludwig, Langer, C., and Quincke, F., 614 Mond, Ludwig, Ramsay, William, and Shields, John, 136, 137 Moore, H. K., Richter, G. A., and Van Arsdel, W. B., 947, 952 Moore, R. T., with Egloff, 909 Morrell, R. S., with Fischer, 187 Mouneyrat, A., 199, 284, 289 Mouneyrat, A., and Pouret, Ch., 284 Moureu, Ch., with Moissan, 637 Mulder, E., 246 Müller, A., and Dubois, H., 285 Müller, Hugo, 278, 795 Müller, N. L., with Kohn, 293 Müller, 287 Müntz, A., 325 Murat, Marcel, 475 Murat, Marcel, with Mailhe, 384, 385, 494 Murat, Marcel, with Sabatier, 343, 348, 362, 364, 369, 389, 415, 449, 452, 453, 455, 471, 475, 488, 523, 538, 714, 720, 721 Murat, M., and Amouroux, G., 414, 415

Nametkin, S. S., and Abakumovskaya, L. N., 478 Naumann, Alex., Moser, Ludwig, and Lindenbaum, Ernst, 260, 269 Neuberg, Carl, 268 Neuberg, C., with Wohl, 237

Mylo, B., with Wohl, 725

Nencki, M., 899
Neogi, P., and Chowdhuri, T., C., 382
Neumann, A., with Gabriel, 107
Neumann, G., 137
Niederhäusern, Heinrich V., 787
Niggemann, H., with Fischer, F., 930
Nikitine, N., with Krestinsky, 713
Nord, F., with Skita, 555
Norman, W., 542, 598, 940, 941, 945
Norman, W., and Pungs, W., 598
Norman, W., and Schlick, F., 80, 583
Norton, L. M., and Prescott, C. O., 691

Geonomides, S., 795
Oehme, H., with Paal, 555
Oelsner, K., with Semmler, 570
Oldenberg, Babette, with Oldenberg, 272
Oldenberg, Hermann, and Oldenberg,
Babette, 572
Olivier, S. C. J., 893
Orloff, E. I., 253, 254, 256
Orndorf, W. R., 223
Ostromuislenskii, I. I., and Kelbasinski, S. S., 784
Ostromuislenskii, I. I., and Kosheler,
I. I., 214
Ostwald, W., 8, 37, 140, 178, 180, 336
Overton, B., 893

Paal, C., 69, 72, 248, 251, 542, 544, 547,

555

Paal, C., with Skita, 555 Paal, C., and Amberger, C., 69, 70, 545 Paal, C., Biehler, F., and Steyer, H., 70 Paal, C., and Gerum, J., 72, 546, 556 Paal, C., and Hartmann, Wilhelm, 1800, 546 Paal, C., and Hohenegger, C., 212, 548 Paal. C., and Oehme, H., 555 Paal, and Schwars, A., 548, 558 Paar, W., with Wolffenstein, 269n Padoa, Maurisio, 485 Padoa, M., with Carrasco, 497, 684 Padoa, M., and Carughi, A., 489 Padoa, M., and Chiaves, C., 490 Padoa, M., and Fabris, U., 484, 491, 642, 643 Padoa, M., and Ponti, 371, 434, 487, 619 Padoa, M., and Scagliarini, G., 647 Parcus, E., and Tollens, B., 188

Pardee, A. M., and Reid, E. E., 340n

Parker, H. K., 285n Passmore, F., with Fischer, 221 Paterno, E., 282 Patrick, W. A., 75n, 180c Patry, E., with Pictet, 270 Pauwels, Joseph, 236 Peachey, S. J., 11, 104, 108 Péligot, E., with Dumas, 691 Perkin, W. H., 107 Perkin, W. H. Jr., 224, 795 Perrin, Jean, 180j Peter, Arnold H., 691 Pétricou, 288 Petri, Camille, 182 Pétrie, 75 Pfau, A., with Houben, 569 Phillips, P., 4 Pictet, Aimé, and Lerczynska, 936 Pictet, Aimé, and Patry, E., 270 Piloty, O., with Fischer, 187 Pinkney, 261n Piria, R., 328, 329, 851 Pishchikov, P. V., with Zal'kind, 38 Plattner, 75 Plotnikov, V. A., 284 Plummerer, Rudolf, with Wilstätter, 835 Ponder, A. C., with Claisen, 798 Ponti, with Padoa, 371, 434, 487, 619 Pope, Wm. J., with Atkinson, 282n Potter, H. M., with Rosanoff, 324 Prescott, C. O., with Norton, 691 Priebs, Bernhard, 89, 803 Pring, John N., 525 Prins, H. J., 198, 216, 242, 625, 903 Pungs, W., with Norman, 598 Purgotti, A., and Zanichelli, L., 563

Quincke, F., with Mond, 614

Rabinovits, Louis, with Ellis, 601
Rack, E., with Erdmann, 944
Radziewanowski, Cornelius, 886, 888
Rai, Hashmat, 265
Rakitin, W., with Ipatief, 592
Ramsay, Wm., with Mond, 136, 137
Rather, J. B., and Reid, E. E., 601
Raupenstrauch, G. A., 795
Rây, J. N., and Dey, M. L., 815n
Raynaud, A., with Ducelliez, 292
Reboul, E., 212
Reformatski, A., 298

Regnault, V., 131 Reich, 75 Reicher, L. T., 178 Reicher, L. T., with Jarissen, 100 Reid, E. Emmet, 10n, 241n, 285n, 340n, 538n, 696n, 772n, 816n, 778n, 947n Reid, E. Emmet, with Freas, 758n Reid, E. Emmet, with Helfrich, 278 Reid, E. Emmet, with Kramer, 707n, 708n, 744n Reid, E. Emmet, with Pardee, 340n Reid, E. Emmet, with Rather, 601 Reid, E. Emmet, with Van Epps, 812 Reimer, Marie, and Downes, Helen R., 340n Reiset, J., and Millon, E., 637 Remsen, Ira, and Keiser, E. H., 150 Riban, J., 216, 795 Richardson, A. S., 269n Riché, 397 Richter, G. A., with Moore, 947, 952 Richter, W., with Semmler, 570 Rideal, E. K., 180j Rideal, E. K., and Taylor, H. S., 180s Rigaud, L., 328 Rigaut, A., with Lescoeur, 230 Rilliet, A., and Ador, E., 291 Risse, F., with Semmler, 570 Ritter, H., with Skita, 549, 642 Rochleder, F., 340 Roenisch, P., with Semmler, 570 Röntgen, W. C., 324 Rosanoff, M. A., and Potter, H. M., 324 Rosenmund, K. W., 575 Rosenmund, K. W., and Zetsche, F., 545 Rossel, Arnold, 220 Rost, H., with Darzens, 390, 459, 476 Rothmund, Victor, 324 Roux, Léon, 199, 293 Rosanov, N. A., 193, 472 Russanow, A., 280 Rutala, O., with Ipatief, 211, 714 Ryumshin, P. F., with Chichibabin, 901

Sabatier, P., 10, 146, 180, 364, 397, 399, 400, 402, 416, 418, 511, 515, 590, 932, 939
Sabatier, P., and Espil, Léo, 12, 16, 56, 80, 113, 114, 125, 338, 346, 358, 492,

598

Schicht, G., Akt.-Ges., 942

Sabatier, P., and Gaudion, G., 631, 634. 641, 643, 644, 645, 680, 681, 726 Sabatier, P., and de Godon, F., 801 Sabatier, P., and Mailhe, Alphonse, 75, 77, 78, 112, 127, 162, 169, 170, 258, 337, 343, 347, 363, 385, 887, 391, 404, 406, 407, 420, 422, 430, 431, 487, 438, 442, 457, 458, 461, 470, 475, 486, 495, 521, 617, 621, 628, 641, 651, 655, 660, 672, 673, 674, 677, 689, 692, 702, 704, 706, 708, 709, 714, 715, 717, 781, 733, 734, 735, 737, 739, 743, 744, 745, 746, 762, 766, 769, 771, 772, 774, 777, 785, 786, 787, 788, 789, 791, 794, 822, 823, 824, 834, 839, 842, 843, 844, 845, 849 850, 852, 853, 856, 857, 858, 866, 873, 876, 878, 916 Sabatier, P., Mailhe, Alph., and Gaudion, G., 923 Sabatier, P., and Murat, M., 343, 348, 362, 364, 369, 389, 415, 449, 452, 453, 455, 471, 475, 488, 523, 538, 714, 720, Sabatier, P., and Senderens, J. B., 26, 111, 208, 343, 351, 362, 368, 370, 374, 375, 376, 377, 578, 379, 388, 394, 396, 410, 413, 414, 419, 424, 425, 427, 433, 435, 436, 446, 449, 451, 456, 460, 464, 466, 467, 468, 469, 471, 475, 477, 481, 482, 494, 497, 500, 501, 502, 503, 504, 506, 508, 509, 510, 511, 512, 515, 517, 518, 519, 520, 521, 526, 527, 530, 531, 533, 534, 542, 614, 616, 619, 622, 637, 652, 654, 656, 658, 659, 660, 664, 666, 668, 683, 701, 832, 912, 914, 916, 919, 920, 925, 928, 939 Sainte-Claire-Deville, H., 216 Sainte-Claire-Deville, H., and Debray, H., 64, 822 Sandmeyer, T., 91, 607, 608, 609 Sastry, S. G., 265 Sayre, R., with Schorger, 235 Saytseff, Michael, 536, 306 Saytseff, Michael, with Kolbe, 165, 536 Scagliarini, G., with Padoa, 647 Schaeffer, with Köts, 550 Scharf, E., with Michael, 200 Scheiber, Johannes, 893 Scheufelen, Adolf, 293 Schiaparelli, Cesare, 293 Schibsted, Helge, with Hofmann, 824

Schicht, G., Akt.-Ges., and Grün, A., 846 Schiel, J., 282 Schimmel, A., with Böeseken, 224 Schlick, F., with Norman, 80 Schlinck, H., and Co., 960 Schmaedel, W, von, with Dimroth, 816 Schmidt, C., with Schraube, 206 Schmidt, J., and Fischer, Ernst. 571 Schmidt, J., and Metzger, R., 454, 484 Schneider, W. von, 210 Schnellenberg, Albert, with Sonn, 565 Schoenbein, C. F., 1800 Scholl, Roland, and Mansfield, Johannes, 685 Scholl, Roland, and Seer, Christian, 685 Scholl, R. Seer, Chr., and Wellsenbook. R., 685 Scholtz, M., 799 Schöne, Em., 160 Schönfeld, H., 944 Schorger, A. W., 930 Schorger, A. W., and Sayre, R., 235 Schraube, C., and Schmidt, C., 206 Schrohe, A., 212 Schrotter, H., with Kekulé, 199 Schülke, Kurt, with Heller, 889n, 898 Schulman, G. G., with Instief, 838 Schultze, Paul, 320 Schumpelt, K., with Hofmann, 271 Schützenberger, 637 Schwartz, A., with Kelber, 69, 548 Schwars, A., with Paal, 548, 558 Schwoerer, 941, 959 Sdzitowecky, W., with Ipatief, 713 Seelig, E., 283, 285, 286 Seer, Christian, with Scholl, 685 Seiichideno, 115 Seligman, R., and Williams, P., 12 Semmler, F., W., and Becker, C., 570 Semmler, F., W., Jonas, K. G., and Oelsner, K., 570 Semmler, F. W., Jonas, K. G., and Richter, W., 570 Semmler, F. W., Jonas, K. G., and Roenisch, P., 570 Semmler, F. W., and Risse, F., 570 Senden, G. H. von with Bösseken, 664 Senderens, J. B., 78, 511, 694, 696, 700, 713, 718, 719, 725, 840, 844, 849, 855, 858, 873, 874, 878, 881

Senderens, J. B., with Sabatier, 26, 111, 208, 343, 351, 362, 368, 370, 374, 375, 376, 377, 378, 379, 388, 394, 396, 410, 413, 414, 419, 424, 425, 427, 433, 435, 436, 446, 449, 451, 456, 460, 464, 466, 467, 468, 469, 471, 475, 477, 481, 482, 494, 497, 500, 501, 502, 503, 504, 506, 508, 509, 510, 511, 512, 515, 517, 518, 519, 520, 521, 526, 527, 530, 531, 533, 534, 542, 570, 614, 616, 619, 622, 637, 652, 654, 656, 658, 659, 660, 664, 666, 668, 683, 701, 832, 912, 914, 916, 919, 920, 925, 928, 939 Senderens, J. B., and Aboulenc, Jean, 598, 758, 759, 760 Sénéchal, A., with Colin, 11 Seyewetz, A., and Gibello, 221 Shaw, T. W. A., 601 Shroeter, G., 481n Shukow, A. A., 942 Siemens, 397 Silber, P., with Ciamician, 150 Silbermann, J. T., with Favre, 131 Silva, R. D., 890 Siveke, with Leprince, 542 Skita, A., 71, 420, 546, 548, 553, 559, Skita, A., and Brunner, W., 561 Skita, A., and Meyer, W. A., 69, 545, 551, 552, 557, 559, 560 Skita, A., Meyer, W. A., and Bergen, J. von, 554 Skita, A., and Nord, F., 555 Skita, A., and Paal, C., 555 Skita, A., and Ritter, H., 549, 642 Skraup, H., 182 Smirnof, V. A., 369, 465 Snethlage, H. C. S., 12 Soc. de Stéarinerie, and Savonnerie de Lyon, 949 Sommelet, Marcel, 818, 889, 899 Sommer, Rudolf, 268 Sonn, Adolf, and Schnellenberg, Albert, Sonnenfeld, Eugen, with Willstätter, 251 Spier, Arthur, with Fischer, 753 Spilker, A., with Krämer, 217 Spohr, J., 324 Sponagel, Paul, with Ullmann, 904 Sprent, C., 413, 713 Squibb, Edward R., 161, 180a, 837

Steinkopf, Wilhelm, and Freund, Michael, 931 Steyer, H., with Paal, 70 St. Gilles, Pean de, with Berthelot, 758n Stock, A., and Guttmann, O., 8 Stockhausen, F., and Gattermann, L., Stollé, R., 201 Stone, W. E., and Tollens, B., 727 Strange, E. H., with Matthews, 213 Strecker, Adolph, 240 Stricker, Otto, with Kekulé, 182 Strutt, R. J., 180c Sulsberg, N., with Thiele, 203 Sutherland, M. J., with Henderson, 463 Suto, K., 268 Sutton, Lens M., with Evans, 691 Taboury, Félix, with Bodroux, 421 Taboury, Félix, with Godchot, 390, 421, 436, 856 Tafel, J., with Fischer, 237 Tanatar, S., 182, 193 Tanret, C., 188 Tanzen, August, with Meyer, 683 Taylor, Clara M., with Easterfield, 843 Taylor, H. S., 180b, 180o, 180q Taylor, H. S., with Rideal, 180s Tchougaeff, L., and Fomin, W., 570 Teuchert, R., 312 Thénard, with Dulong, 637 Thiele, with Mannich, 966 Thiele, Johannes, and Sulsberger, N., 203 Tiemann, Ferd., 191 Tiemann, Ferd., and Haarmann, Wilh., Tiemann, F., and Krüger, Paul, 198, 800 Tikhvinskii, V. M., with Arbusow, 611. 633, 635 Tilden, W. A., with Armstrong, 198 Tingle, J. B., and Blanck, F. C., 269n Tischenko, V. E., 228, 299 Tollens, B., with Parcus, 188 Tollens, B., with Stone, 727

Tollens, B., with Yoder, 727 Tollens, B., and Wigand, P., 83

Trannoy, with Matignon, 75, 259

Tornthwaite, 75

Traube, M., 73

Staudinger, H., and Klever, H. W., 235

Trey, H., 317
Tribe, Alfred, with Gladstone, 165, 166, 169, 785
Trillat, A., 73, 249, 253, 256
Trillat, A., and Gambier, 781
Trunov, B. V., with Chelintzev, 805
Tschelinzeff, W., 301
Turbaba, D., 22
Turner, Edward, 10, 116
Twomey, T. J., with Egloff, 908
Tyndall, 180j

Uklonskaja, Miss, N., with Zelinski, 649 Ullmann, Fritz, 901 Ullmann, F., and Sponagel, Paul, 904

Valpy, O. H., and Lucas, O. D., 933 Van Arsdel, W. B., with Moore, 947, 952 Van der Scheer, J., with Böeseken, 879 Van der Weide, O. B., with Böeseken, 117, 546

Van Ekenstein, W. A., with Lobry de Bruyn, 186

Van Ekenstein, W. A., and Blanksma, J. J., 186

Van Epps, G. D., and Reid, E. Emmett, 812

Van't Hoff, 139, 175

Varet, Raoul, and Vienne, G., 241 Vavon, G., 63, 567, 568, 570

Vavon, with Lespieau, 566 Vavon and Faillebin, 565

Veley, V. H., 8, 269n

Veraguth, H., with Willstätter, 480

Verein, Chininfabr. Zimmer, and Co., 572, 604

Verein, f. Chem. Ind., 254 Verinigte Ch. Werke, 946

Vienne, G., with Varet, 241 Vignon, Léo, 269n, 540

Villiers, A., 153

Vogt, J. D. de, with Böeseken, 879

Voigt, with Michael, 200

Vorländer, D., and Hobohm, K., 799

Wacker, L., 274

Waldschmidt-Leits, Ernst, with Willstätter, 62n, 167n, 562n, 563n, 573n, 943n, 947n.

Wallach, O., 198, 205, 546, 552, 797 Wallach, O., and Wüsten, M., 97 Ward, A. L., with Boehner, 811 Waser, E., with Willstätter, 480, 535, 571

Weismann, Charles, and Garrand, S. F., 654n

Weiss, J. M., and Downs, C. R., 260n Wells, A. A., with Ellis, 949

Wellsenbock, R., with Scholl, 685

Wesson, David, 967n

Wilbaut, J. P., 683

Wieland, Heinrich, 251

Wieland, H., and Wishart, R. S., 579

Wigand, P., with Tollens, 83

Wiggers, A., 307

Wilbuschewitch, M., 941, 945, 961

Wild, W., with Engler, 150

Wilde, M. P. von. 342, 526

Wilfarth, H., 272

Willgerodt, C., 283

Willgerodt, C., and Genieser, Ad., 238
Williams, C. E., with Bedford, 939, 941.

943

Williams, P., with Seligman, 12 Williams, R. R., and Gibbs, H. D., 254n

Williamson, Alexander, 159, 169, 848, 851

Willstätter, Richard, 542, 562, 563

Willstätter, Richard, and Bruce, James, 293, 473

Willstätter, R., and Hatt, D., 569, 571 Willstätter, R., and Heidelberger, M.,

Willstätter, R., and Jacquet, D., 563, 569, 571

Willstätter, R., and Kametaka, T., 197, 472, 479

Willstätter, R., and King, V., 569, 571
Willstätter, R., and Mayer, E. W., 565,
569

Willstätter, R., and Pummerer, Rudolf, 835

Willstätter, R., and Sonnenfeld, Eugen, 251

Willstätter, R., and Veraguth, H., 480 Willstätter, R., and Waser, E., 480, 535, 571

Willstätter, Richard, and Waldschmidt-Leits, Ernst, 62n, 167n, 562n, 563n, 573n, 943n, 947n

Wilsmore, N. T. M., 829

Wilson, John, with Mason, 262

AUTHOR INDEX

Wimmer, K. H., and Higgins, E. B., 944 Winkler, Clemens, 73 Wippermann, R., 230 Wischnegradsky, A., 210 Wishart, R. S., with Wieland, 579 Witsemann, Edgar, J., 725 Wohl, A., 324 Wohl, A., and Mylo, B., 725 Wöhler, F., and Liebig, Justus, 220 Wöhler, F., and Mahla, 75 Wöhler, Lothar, with Engler, 137, 154 Wohl, A., 94 Wohl, A., and Neuberg, C., 237 Woker, Miss Gertrude, 180i Wolffenstein, R., and Böters, O., 269 Wolffenstein, Richard, and Hartwich, Frank, 893 Wolffenstein, R., and Paar, W., 269n Wollemann, J., with Borsche, 546 Wollrath, A., 278 Woltman, A., 964 Woog, Paul, 257, 259 Woselka, Hermann, with Frank, 223 Wurtz, Adolphe, 11, 219, 307, 334, 605 Wüsten, M., with Wallach, 97

Yoder, P. A., and Tollens, B., 727 Youssouffian, with Haller, 341

Zagumenni, A., 692 Zal'kind, Y. S., 548, 566 Zal'kind, Y. S., and Kvapishevskii. K. V., 548, 566 Zal'kind, Y. S., and Miss Markarvan, V., 548, 566 Zal'kind, Y. S., and Pishchokov, P. V., 38 Zanetti, C. U., 742 Zanetti, J. E., and Egloff, G., 907 Zanichelli, L., with Purgotti, 563 Zdrawkowitch, Milan R., 62 Zeise, 62 Zelinski, H., 302 Zelinsky, N. D., 390, 472, 478, 534, 535, 649, 906, 934 Zelinsky, N. D., and Glinka, N., 648. 822 Zelinsky, N. D., and Herzenstein, Miss Anna, 649 Zelinsky, N., and Uklonskaja, Miss N., 649 Zetsche, F., with Rosenmund, 545 Zincke, Th., with Breuer, 220 Zincke, Th., with Kekulé, 222 Zinin, N., 203, 220 Zsigmondy, R., 180i

SUBJECT INDEX

(References are to Paragraphs: a number followed by "n" designates a note.)

A1	£ 1 1001
Absorption of gases, 131, 135	formed, 180b
Acenaphthene, hydrogenated, 482	into nitrile, 812
Acetaldehyde, 219, 222, 261, 700, 724,	by oxidation, 48, 150, 255, 257, 261
725	retarder, 11
into acetals, 780, 782	Acetols, 783
from acetylene, 92, 309, 310	Acetone, into acetol, 783
as catalyst, 105, 310, 312	with chloroform, 238
condensed, 52, 592, 780, 782, 795, 796, 801, 807	condensed, 238, 783, 797, 797n, 798, 800, 801, 805
crotonized, 795, 801	condensed with aldehydes, 798
decomposed by Pd., 623, 680	condensed with benzaldehyde, 798
into ester, 228	condensed with citral, 800
formed, 180g, 200	condensed with o.nitrobensaldehyde,
hydrogenated, 432, 439, 494, 538,	798
593, 664, 668, 670, 673	crotonized, 797, 798
with hydrogen sulphide, 810	decomposed, 620, 659, 665, 668
by oxidation, 249, 254, 256, 268	formed, 161, 180a, 249, 309, 831, 837,
oxidised, 255, 260, 261	844
preparation of, 309	hydrogenated, 435, Co 503, Cu 594,
Acetal, by oxidation, 249	Fe 593, Ni 588, 598, Pt 567, Zn 595
Acetals, 81, 97, 175, 249, 305, 779-783	by oxidation, 249, 254a
formed, 81, 106, 249	preparation of, 161
hydrolysis of, 322, 323	Acetone-oxime hydrogenated, 383
preparation of, 782	Acetonitrile as catalyst, 108, 605
Acetamide, dehydrogenated, 811	formed, 871
hydrolyzed, 336, 386	polymerized, 50, 231, 427
Acetanhydride, 107	Acetonyl-acetone hydrogenated, 440
decomposed, 829	Acetophenone, 848, 849
in esterification, 761	catalytic solvent, 38
into ketone, 857	condensed, 799
Acetanilide in synthesis, 901	by Friedel and Crafts reaction, 891,
Acetic acid, into acetone, 161, 180a,	893
840-845	hydrogenated, 389, 455, 538, 539,
from acetylene, 255	568
from alcohol, 48, 150, 255, 257,	Acetophenone-oxime hydrogenated, 384
261	Acet-oxime hydrogenated, 383
as catalyst, 106, 215, 687, 780	Acetyl-acetone hydrogenated, 439, 595
chlorinated, 280	Acetyl-brom-glucose condensed, 793
decomposed, 831, 843	Acetyl chloride formed, 280
esterified, 750, 758, 760, 771	in F. and C. reaction, 891–893
esters of decomposed, 863	Acetylation, 81, 240

Acetyl-chloramino-benzene isom., 202 Acroleine, 101, 249, 658, 680, 713, 725, Acetyl-cyclo-hexene hydrogenated, 476 Acetyl-diphenyl-amine, syn., 901 hydrogenated, 419 Acetylene, 38, 102, 308-310 Acrylic acid hydrogenated, 417 condensed. 686. Co 928, Cu 916, Fe by oxidation, 249 928. Ni 925-928 Activated charcoal as catalyst, 282n condensed with benzene, 241 Active alkaloids as catalysts, 836 decomposed, 637, 913-920 Active modifications, 180i formed, 409 Adipic acid, 251 hydrated, 27, 92, 308 Adsorption theory, 180c hydrogenated by Co, 501, Cu 518, Fe Aeration of Pt. black, 563 506, Ni 423, 424, Pd 548, 558, Pt Agitation in catalysis, 541 342, 527 in hydrogenation, 957 polymerized, 212, 914 Albumin, stabiliser for, Pd. 69 Acetylene bond, migration of, 192 Alcohol eliminated, 817 Acetylene compounds hydrogenated. as source of H. 537 423-425, 518, 527, 548, 558, 566, as solvent in hydrogenation, 599 577, 601 toxic to Pt. black, 117 Acetylene glycols hydrogenated. 548. Alcoholates, 299 566, 577 Alcohols with aldehydes, 784 Acetylene hydrocarbons decomposed, decomposed, 180g, 538, CdO 674, 913-919 C 679, SnO 673, Zn 678 hydrated, 308 dehydrated, 28, 75, 98, 99, 138, 180h dehydrogenated, 28, 650-679 polym., 212 Acetylene tetrabromide, 289, 897 by hydration of hydrocar., 305 Acetylene tetrachloride, 199 hydrogenated, 416 Acetyl-vanilline hydrogenated, 568 oxidised, 246, 249, 254, 268 Acid amides hydrolyzed, 331 Alcohols, aromatic, hydrogenated, 369, Acid anhydrides formed, 872 465 hydrogenated, 392 Alcoholysis, 340, 340n Aldehydes, 236, 653, 655, 668, 701, 723 into ketones, 857 Acid chlorides, 243 acetylated, 240 with ammonia, 813 into acetals, 780-783 decomposed by AlCla, 625 from alcohols, 15, 48, 246, 650 in F. and C. reaction, 891-894 condensed, 90, 237, 239, 240, 794hydrogenated, 575 810 Acids, in formation of acetals, 783 crotonised, 794-801 from alcohols, 150, 246, 275 decomposed, 618-623, 532, 549 catalysts, 17, 81 dehydrated, 794-802 decomposed, 171, 820-856 by depolymerisation, 234 in depolymerisation, 234 into esters, 225-228 in esterification, 748-756 formed, 15, 28, 31, 48, 75, 142, 200, hydrogenated, 422 208, 866 in hydrolysis, 175, 305 hydrogenated, 388, 419, 432-434, 568, Co 503, Cu 522, Fe 506, 593, in hydrolysis of glucosides, 175 in inversion of sugar, 175 Ni 588, 602, Pt 567 in isomerisation 181-182 by oxidation, 15, 48, 258-261, 268, Aconitic acid hydrogenated, 558 275 Acridine hydrogenated, 491 from oximes, 268, 332 oxidised, 270 with phenols, 792 Acridone, 270 polymerized, 82, 106, 218-228

SUBJECT INDEX

	41 1 1 200
preparation of, 851–854	Almond oil, 938
Aldolization, 82, 83, 95, 97, 219, 221	Alumina as carrier, 127
Aldols depolymerized, 234	as catalyst, 75, 78, 169, 180j, 540,
formed, 219	586, 624, 676, 693, 694, 700, 714,
Aldoximes dehydrated, 814	721, 722, 732, 740, 784, 797n, 807-
hydrogenated, 383, 384, 514	814, 825
isomerized, 204	in cracking, 906, 934
Alicyclic ketones hydrogenated, 436	decomp. chlorides, 881
Aliphatic hydrocarbons cracked, 911,	decomp. esters, 861n, 866, 872-874
912	decomp. hexane, 920
Alisarine, 274, 328	dehydration catalyst, 142, 180g,
Alkalies in hydrolysis, 305	180h, 651, 686, 743, 784, 791
Alkaline bisulphates in esterif., 748,	esterification cat., 764
759, 760	isomerises unsat. hydroc., 190, 200
carbonates as cats., 97	ketone cat., 840, 849
halides elim., 904	life of, 708
Alkaloids as cats., 836	mercaptan cat., 746
Alkyl-anilines, 729	polymeris. unsat. hydroc., 211
Alkylation of arom. hydrocar., 884-890	preparation of, 77, 706, 713, 714
Alkyl bromides decomposed, 876	Aluminum cat., 284, 886, 901
in F. and C. syn., 885	in cracking, 906
in Grignard reaction, 302	dec. hydrocarb., 920
isomerized, 199, 200	influence on, Pd, 946
Alkyl chlorides decomposed, 86, 876-	Aluminum alcoholates, 12, 299
882	amalgam, 293
in syn., 883–889	catalyst, 51
Alkyl halides, 104, 300-304, 883-885	dissolved in alcohols, 12
isomerized, 876	oxidation cat., 254
Alkyl iodides, 304	Aluminum bromide, bromination cat.,
decomposed, 876	93, 293
in F. and C. reaction, 885	catalyst, 298
Alkyl phenol ethers, 494, 789	chlorination cat., 289
hydrogenated, 464	isomeris. cat., 119
Alkyl piperidines, 741	in F. and C. reaction, 893
Alkyl sulphides, 626–628, 743, 744	Aluminum chloride, bromination cat.,
Alkyl ureas, 431	293
Allenic hydrocarb. hydrated, 308	with carbon monoxide, 298
isomerized, 192	catalyst, 6, 33, 87, 173-174, 239-243,
Allyl alcohol 601, 680, 740	293–297, 687, 728, 795, 797, 803,
dehydrated, 713	806, 817–819, 877–879, 88 4–9 00,
esterified, 757	903, 929–931, 935
isomerized, 208, 658	chlorination cat., 283, 284
oxidised, 249	
	condensation cat., 903
hydrogenated, 416, 558	cracking cat., 935
Allyl amine as cat., 836 Allyl bensoate, 766	on cymene, 930
•	dehydrogenation cat., 638, 685
Allylene hydrated, 309	in F. and C. reaction, 896–900
polymerized, 212	on hydrocarb., 929–931
Allyl iodide, 605	on naphthalene, 931
Allyl memory 744	on pinene, 931
Allyl mercaptan, 744	polymerises hydrocarb., 211, 216

regenerated, 935	Amino-nitrobensene, syn. of, 901
in sulphonation, 295	Amino-phenol, 630, 632
with sulphur dioxide, 297	Amino-phenols, 381, 536
theory of action, 173, 687, 728	Amino-succinic acid, 312
on thiophenol, 629	Ammonia into amines, 901
on xylene, 930	condensed with alds., 807-809, 812,
Aluminum ethylate polymer. alds., 228	813
preparation, 299	condensed with ketones, 809
Aluminum phosphate, 99, 719, 726	
	condensation cat., 803, 804
Aluminum powder cat., 901	in cracking, 933
Aluminum salts dehydration cat., 717 in nitration, 269n	decomposition cat., 637 eliminated, 611, 631–633
	with esters, 871
Aluminum sulphate cat., 99, 696, 706,	
717, 725, 759, 760 Aluminum turnings chlorination cat.,	by hydrogenation, 377
284	from nitrie oxide, 368, 374
	oxidised, 249, 256
in F. and C. reaction, 886	polymeris. cat., 104
with mercuric chloride, 886	syn. of, 180s, 180t, 180u, 342
Amarine formed, 194	Ammonium alum, 784
American hardened oils, 967s	Ammonium chloride cat., 97, 783
American petroleum, 908, 928, 932	Ammonium isosulphocyanate, 104, 207
Amides dehydrated, 811 hydrogenated, 386, Cu 514, Ni 386	Ammonium nitrate, 97, 256, 375, 376 783
hydrolised, 305, 311, 331, 335, 336	Ammonium nitrite, 375
from nitriles, 311	Ammonium sulphete 07 783
from oximes, 204, 205	Amount of catalyst 508 051
polymerized, 233	Amount of earlyst, 598, 951
syn. by F. and C. reaction, 895	Amphotoric hydroxides 850
Amines, 170, 426, 430, 431, 513, 514,	Amphoteric hydroxides, 859
600, 682	Amylamina 486 631 681
cond. cat., 803 decomp. Ni, 681	Amyl-amine, 486, 631, 681 Amyl alcohol, 150, 673
• • • • • • • • • • • • • • • • • • • •	
dehydrogenated, 681, 682 formed 15, 170, 382, 383, 731–742	Amylene 626 606 746 871 929
formed, 15, 170, 382, 383, 731–742	Amylene, 626, 696, 746, 871, 929
hydrogenated, 496	dec. by AlCla, 929
by hydrogenation, 382, 383 oxidised, 268	hydrated, 306
from oximes, 383	hydrogenated, 558, 565 preparation of, 706n
secondary, 682	Amyl nitrite, tox. to cats., 116
suphurized, 296	Amyl oleate hydrogenated, 526
syn. of, 683, 731–742	Amyl stearate, 526
in vulcanis. of rubber, 104	Anethol hydrogenated, Ni 590, 601
Amino-acetophenone, 545, 557	Anhydrides hydrogenated, 392
Amino-benzoic acid hydrogenated, 569	Aniline, 469, 495, 497, 531, 536, 538,
Amino-caproic acid into lactam, 205	545, 554, 557, 578, 630-632, 683,
Amino-cyclo-hexanes dehydrogenated,	790
642	alkylated, 729, 740
Amino-cyclo-hexane-carbonic acid, 569	eliminated, 634, 635
Amino-ethyl alcohol oxid., 268	formed, 165, 277, 378, 380, 381
Amino-malonic nitrile, 230	hydrogenated, 466, 467, 569
Amino-naphthol, 564	by hydrogenation, 277, 378, 380, 381
AMMO WONTHING, OVE	by my mongementally arrivers, or o, ood, oot

SUBJECT INDEX

manufacture, 378, 511	Araboketose by oxid., 268
	Arabonic acid, 187
	Arachidic esters, 937
	Arbutin hydrolysed, 328
	Argon abs. by C, 180d
7 . ·	Aromatic acids dec., 830
Aniline black, 260n, 271	esterified, 758, 766
Aniline hydrobromide as cat., 726	hydrogenated, 471
	Aromatic alcohols condensed, 728
Animal charcoal, carrier, 946	hydrogenated, 369, 465
catalyst, 48, 282	reduced, 369
	Aromatic aldehydes hydrogenated, 388,
Anisaldehyde condensed, 808	568
	Aromatic amides syn. by F. and C.,
polymerised, 220	895
	Aromatic amines with alcohols, 740
Anisolne, 220	hydrogenated, 466
Anisol condensed, 806	syn., 901
	thioureas, 630
hydrogenated, 464, 494, 589 neg. cat., 11, 303	Aromatic bromides from diaso., 608
	Aromatic chlorides from diaso, 607
	Aromatic ethers, 494
	Aromatic hydrocarbons from C ₂ H ₂ , 926
from acetylene, 914 condensed, 806	alkylated, 877–890 brominated, 291–293
and the second s	by cracking, 932
by cracking, 908, 909	
decomposed, 921 by F. and C. reaction, 897	decomp., 921, 930
_ · · · · · · · · · · · · · · · · · · ·	hydrogenated, 446
hydrogenated, 29, 363, 483, 592	oxidised, 269
oxidised, 257, 280n, 262, 262n, 269, 271	Aromatic ketones hydrogenated, 389, 455, 523, 590
	Aromatic nitro-comps. hydrogenated,
Anthracene hydrides dehyd., 642, 728	511, 600
	Aromatic nucleus hydrogenated, 444-
269, 271, 839, 846	456, Cu 594, Fe 593, Ni 589, Pd
sulphonated, 816	556, 578, Pt 534, 560, 569
T 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1	Aromatic rings hydrogenated, 444–456,
Anthraquinone disulphonic acids, 816	534
	Arsenic toxic to Pt, 116
	Arsenic acid, 691
	Arsenious oxide toxic to Pt, 116
chlorination cat., 283, 287	transformed, 73
	Arsine cat. poison, 1800
Apparatus for dehydration, 717	decomp. of, 8
A 11 1	Asbestos as carrier, 126, 941
	Asparagine formed, 312
	Aspartic imide hydrogenated, 312
	Asymmetric dec. of acids, 836
	Atoms, migration of, 199
· · · · · · · · · · · · · · · · · · ·	Auto catalysis, 8
	Autoclave for, H ₂ , 597
•	

Auto oxidation, 150, 151 444-447, 534, 589, 947n, Co 502, Asobensene, by hydrogenation, 511 Pt 560, 569 hydrogenated, 497, 554 from hydrogenation, 370, 378, 388 Aso-compounds hydrogenated, 600 neg. cat., 11 Asoxy-compounds hydrogenated, 600 oxidised, 260n, 263, 268, 276 sulphonated, 815 Bakelite, 792 from xylene, 930 Baku petroleum, 444 Bensene homologs from acetylene, 518 cracked, 934 hydrogenated, 447-450, Co 502, Pt Barium carbonate cat., 98, 838 569 Barium chloride cat., 86, 876, 880 Bensene ring hydrogenated, Ni 603 Barium hydroxide cond. agt., 800 Benzene sulphinic acid. 297 Barium peroxide poly. cat., 214 Benshydrol, 538, 728, 745 Barium salts decomp., 837 to amine, 736 dehydrated, 688, 692, 720 Barium soaps neg. cat., 115 Bases as cats., 83 dehydrogenated, 650, 662, 720, Pd in hydrolysis, 175, 178 669 Bauxite cat., 706, 706n hydrogenated, 369 Benshydryl amine, 736 Beef tallow, 938 Benzal-acetone, 798 mercaptan, 745 Bensal chloride, 320, 890 Benzidene, 202 Bensaldehyde, into acetal, 783 Benzoic acid from aldehyde, 225 acetalated, 240 into aldehyde, 853 from bensal chloride, 320 from bensotrichloride, 320 decomposed, 830, 834, 839, 840-845 condensed, 89, 798, 799, 803, 804, 807, 808 esterified, 757, 758, 766, rate 758n decomposed, Ni 620, Pd 623 hydrogenated, Ni 590, Pd 551, Pt into ester, 225 559, 560 formed, 165, 575, 657, 674 into ketone, 848-850 by F. and C. react., 297, 298 by oxidation, 257 by hydrolysis, 329 sulphonated, 815n, 816 by oxidation, 249, 257, 259, 260n, 268 from toluene, 150 Benzoic esters, decomp., 864 hydrogenated, Fe 593, Ni 388, Pd 549, 560, Pt 568, thoria 538 hydrogenated, 471 polymerized, 220 Bensoine, 220, 234, 590 Bensaldoximes transformed, 185 hydrogenated, 391 Bensamide by F. and C. syn., 895 Bensonitrile by dehydration of amine, Benzanthrone, 685 681 Bensene, 518, 593, 620, 641, 643, 649, by diaso reaction, 605 657, 674 from esters, 871 from C₂H₂, 914-916 hydrogenated, 428 brominated, 292 polymerized, 232 chlorinated, 278, 284, 285 Benzophenone, 538, 650, 662, 669, condensed, 806, 817-819 720, 845, 846, 890 by cracking, 908, 909 condensed, 809 in cracking, 907 formed, 839, 891, 893, 899 from cymene, 930 by F. and C. syn., 891, 893, 899 decomposed, 907, 921 hydrogenated, 389, 539, 590, Cu in F. and C. syn., 894, 897 523, Pt 560 in Grignard syn., 300 Bensophenone oxime, 384 hydrogenated, 26, 344, 361, 362, Bensotrichloride, 320

Bensoyl-bensoic acid, syn. of, 893 Bensoyl chloride in F. and C. syn., 893, 899 hydrogenated, Pd 675 into nitrile, 813 Bensoyl-proxide cat., 214 Bensoyl-propionic acid, 203 Bensoyl-acitophic acid, 203 Bensyl-acitophic acid, 203 Bensoyl-acitophic acid, 203 Bensoyl-acitophy 203 Boronic acid dec. esters, 844 Boron fluoride polymer. eat., 84, 211 Borneol from acetate, 340 from camplor, 591, 722 dehydrated, 714 dehydration cat., 274 boria cid ec. esters, 844 dehydratogenated, 661 esters of, 340 ordideod, 249 Broads of hardened fate 967, 967n Brass cat., 254n, 670 Broadside acid, 601 Broad	Bensoyl-acetone hydrogenated, 391	dehydration cat., 791
Bensoyl chloride in F. and C. syn., 893, 899 hydrogensted, Pd 575 into nitrile, 813 Bensoyl peroxide cat., 214 Bensoyl-propionic acid, 203 Bensoyl-vanilline hydrogensted, 568 Bensoyl-salicylic ald. hydrogensted, 568 Bensyl-acetone hydrogensted, 389 Bensyl alcohol, 549, 560, 715, 729, 740 from aldehyde, 225 into amine, 734 cat. solvent, 38 condensed, 728 dehydrated, 688, 714 dehydrogensted, 688, 714 dehydrogensted, 688, 714 dehydrogensted, 688, 714 dehydrogensted, 687, 717, 773 hydrogensted, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 catalyst, 836 dehydrogensted, 470, 496, 560 Bensyl-bensylidene-acetone, 547 Bensyl-holride dechlorinsted, 605 desomposed, 880, 916 formed, 231, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl domate, 773 Bensylidene-hydrindone, 799 Bensylidene-hydrindone, 799 Bensylidene-bydrindone, 799 Bensylidene-hydrindone, 799 Bensylidene-hydrogensted, 578 Betuloh hydrogensted, 570 Bi-cyclo-octane hydrogensted, 480 Bi-cyclo-octane hydrogensted, 568 Bensoyl-acetic, 765, 767, 702, 778, 828 Bensyl-acetic, 568 Bensoyl-acetic, 568 Bensoyl-acetic, 568 Bensoyl-acetic, 568 Bensoyl-acetic, 587 Boric aid dec cesters, 864 dehydration cat., 274 toxic to cats., 115 Borne floring acetate, 340 from camphor, 591, 722 dehydration cat., 274 toxic to cats., 115 Borne floring acetate, 340 from camphor, 591, 722 dehydration cat., 274 toxic to cats., 115 Borne floring acetate, 340 Framds of hurdened fats 967, 967n Brass lock furnace, 248 Brassidic acid formed, 184 into ketone, 34	_ '	
883, 899 hydrogenated, Pd 575 into nitrile, 813 Bensoyl peroxide cat., 214 Bensoyl-propionic acid, 203 Bensoyl-salicytic ald. hydrogenated, 568 Bensyl-sactone hydrogenated, 568 Bensyl-scotone hydrogenated, 389 Bensyl alcohol, 549, 560, 715, 729, 740 from aldehyde, 225 into amine, 734 cat. solvent, 38 condensed, 728 dehydrated, 688, 714 dehydrogenated, 688, 714 dehydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 catalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-hensylidene-acotone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 231, 818, 839 in F. and C. syn., 90, 901 Bensyl-loencylohotylohotylohotylohotylohotylohotylohotylohotylohotylohotylohotylohotylohotylohotylohotylohotylohotyloho		
hydrogenated, Pd 575 into nitrile, 813 Bensoyl-propionic soid, 203 Bensoyl-salicylic ald, hydrogenated, 568 Bensyl-acetone hydrogenated, 568 Bensyl-acetone hydrogenated, 389 Bensyl alochol, 549, 560, 715, 729, 740 from aldehyde, 225 into amine, 734 eat. solvent, 38 eondensed, 728 dehydrated, 688, 714 dehydrogenated, 688, 714 dehydrogenated, 688, 714 dehydrogenated, 689, 715, 729 dehydrated, 683, 714 dehydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 eatalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-anilline, 729 Bensyl-lensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 889, 901 Bensyl formate, 773 Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene-hydrindone, 799 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene-hydrindone, 799 Bensylidene-hydrindone, 799 Bensylidene-hydrindone, 799 Bensylidene-hydrindone, 799 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene acetate formed, 240 Bensing formed, 808 Bromaniline, 708 Bromhensov chioride in F. and C. syn., 893 Bromines cat., 436 Bromhensov chioride in F. and C.		
into nitrile, 813 Bensoyl-propionic acid, 203 Bensoyl-scotone hydrogenated, 568 Bensoyl-scotone hydrogenated, 389 Bensyl alcohol, 549, 560, 715, 729, 740 from aldehyde, 225 into amine, 734 cat. solvent, 38 condensed, 728 dehydrated, 688, 714 dehydrogenated, 369, 771, 773 hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 catalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 380, 916 formed, 281, 818, 889 in F. and C. syn., 839 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl-dyclohexyl-amine, 739 Bensyl idene acetate formed, 240 Bensylidene-malonic acid, 804 Bensyl-pyridines syn., 901 Berylidene-malonic acid, 804 Bensyl-pyridines syn., 901 Berylidene-malonic acid, 804 Bensyl-pyridines syn., 901 Berylidene-fyridines, 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 2897 Blue oxide of molybdenum cat., 675, Bensyl-acetone hydrogenated, 568 Bensoyl-acetone hydrogenated, 568 Bensoyl-acetone bydrogenated, 568 Bensoyl-acetone hydrogenated, 568 Bensyl-acetone hydrogenated, 568 Bensyl-acetone hydrogenated, 568 Bensyl-acetone bydrogenated, 568 Bensyl-acetone hydrogenated, 568 Bensyl-acetone bydrogenated, 569 Bromaniline cat., 687, 687 britanion cat., 746 bridahydration cat., 749 bridahydration cat., 741 toxic to cate., 115 Borne alidydration cat., 687 bror individual preparation, 944 toxic to cate., 115 Borne alidydration cat., 274 toxic to cate., 115 Borne alidydration cat., 274 toxic t	hydrogeneted Pd 575	
Bensoyl-propionic acid, 203 Bensoyl-salicylic ald. hydrogenated, 568 Bensoyl-actone hydrogenated, 568 Bensyl-acetone hydrogenated, 389 Bensyl alcohol, 549, 560, 715, 729, 740 from aldehyde, 225 into amine, 734 cat. solvent, 38 condensed, 728 dehydrated, 688, 714 dehydrogenated, 667, 711, 773 hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 catalyst, 836 dehydrogenated, 681 hydrogenated, 681 hydrogenated, 681 hydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-bensylidene-acetone, 547 Bensyl-bensylidene-acetone, 547 Bensyl-byroidnes, 799 Bensyl formate, 773 Bensyl-cyclohevyl-amine, 739 Bensyl formate, 773 Bensyl-gyroidnes acetate formed, 240 Bensyl-gyroidnes acetate formed, 240 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific, cat., 746 preparation of, 715 Boric acid dec. esters, 844 dehydration cat., 687 influence in hydrogenation, 944 oxidation cat., 274 toxic to cats., 115 Bornon fluoride polymer. cat., 84, 211 Bornon fluoride polymer. cat., 84 Bornal sol c		
Bensoyl-propionic acid, 203 Bensoyl-propionic acid, 203 Bensoyl-vanilline hydrogenated, 568 Bensyl-acetone hydrogenated, 389 Bensyl alcohol, 549, 660, 715, 729, 740 from aldehyde, 225 into amine, 734 eat. solvent, 38 condensed, 728 dehydrated, 688, 714 dehydrogenated, 630, 715, 729, 740 from aldehyde, 225 into amine, 734 eat. solvent, 38 condensed, 728 dehydrated, 688, 714 dehydrogenated, 630, 715, 729, 740 from aldehyde, 225 into amine, 734 eat. solvent, 38 condensed, 728 dehydrated, 688, 714 dehydrogenated, 630, 717, 773 hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 catalyst, 836 dehydrogenated, 661 esters of, 340 oxidised, 249 Bensyl-bensylidene-acetone, 547 Bensyl-choride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl-dyrindone, 799 Bensylidene-malonic acid, 804 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific, cat., 778 Betulol hydrogenated, 450 Bi-cyclo-octane, 430 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Bromise acid ec. esters, 864 dehydration cat., 687 influence in hydrogenation, 944 oxidation cat., 274 toxic to cats., 115 Borne fluoride polymer. cat., 84, 211 Borneol from acetate, 340 oxidation eat., 274 toxic to cats., 115 Borneol from acetate, 340 oxidation eat., 274 dehydrated, 714 dehydrogenated, 661 esters of, 340 oxidised, 257, 280n Bornyl acetate, 340 Brands of hardened fate 967, 967n Brass block furnace, 348 Brassidic acid formed, 184 into ketone cat., 242 dehydrated, 714 dehydrogenated, 661 esters of, 340 oxidiston eat., 274 toxic to cats., 115 Borneol from cats., 274 dehydrated, 714 dehydrogenated, 661 esters of, 340 oxidiston, 271 Bensulof, 716, 729, 760 Brass block furnace, 348 Bromines red., 405 Bromines red., 405 Bromines cats., 84 Bromines in hydrogenated, 405, 545 sulphonated, 370 bro		
Bensoyl-salicylic ald. hydrogenated, 568 Bensoyl-vanilline hydrogenated, 568 Bensoyl-vanilline hydrogenated, 568 Bensyl alcohol, 549, 560, 715, 729, 740 from aldehyde, 225 into amine, 734 eat. solvent, 38 condensed, 728 dehydrated, 688, 714 dehydrogenated, 630, 674, Cu 657, MnO 673 esterified, 771, 773 hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 catalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-bensylidene-acetone, 547 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 231, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl dynaide, 605, 871 Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene-hydrindone, 799 Bensyl formate, 773 Bensylidene-malonic acid, 804 Bensylidene-bydrindone, 799 Bensyl formate, 778 Betulol hydrogenated, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Blue oxide of molybdenum cat., 675, Blue oxide of molybdenum cat., 675, Borning ded. ded., dehydrogenation, 944 oxidation cat., 684 benorion in hydrogenation, 944 oxidation cat., 684 oxidation cat., 274 toxic to cats., 115 Borno fluoride polymer. cat., 84, 211 exit solvent, 38 ondensed, 728 dehydrated, 714 dehydrogenated, 661 exters of, 340 oxidised, 257, 260n Bornyl acetate, 340 from camphor, 591, 722 dehydrated, 714 dehydrogenated, 661 exters of, 340 oxidised, 257, 260n Bornyl acetate, 340 From camphor, 591, 722 dehydrated, 714 dehydrogenated, 661 exters of, 340 oxidised, 257, 260n Bornyl acetate, 340 From lacetal, 661 exters of, 340 oxidised, 257, 260n Bornyl acetate, 340 From lacetal, 661 exters of, 340 oxidised, 257, 260n Bornyl acetate, 340 From camphor, 591, 722 dehydrated, 714 dehydrogenated, 661 exters of, 340 oxidised, 257, 260n Bornyl acetate, 340 From lacetal, 345 Bromalitine, 729 Brass block furnace, 348 Bromalitine, 729 Brass block furnace, 348 Bromalitine, 729 Bromal cond., 806 Bromanilines red., 405, 675 into acid, 605 From lovide dechlorinated, 6		
Bensyl-acetone hydrogenated, 568 Bensyl-acetone hydrogenated, 389 Bensyl alcohol, 549, 560, 715, 729, 740 from aldehyde, 225 into amine, 734 eat. solvent, 38 condensed, 728 dehydrated, 688, 714 dehydrogenated, 680, 716, 729, 740 from camplor, 591, 722 dehydrated, 681, 771, 773 hydrogenated, 681, 771, 773 hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 eatalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-aniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl cyanide, 605, 871 Bensyl dyanide, 605, 871 Bensyl dyanide, 605, 871 Bensyl dene-acetate formed, 240 Bensyl-pyridines syn., 899 in F. and C. syn., 889 in F. and C. syn., 899 Bensylidene-acetate formed, 240 Bensylidene-bydrindone, 799 Bensylidene-cate, 685, 716, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 480 Bis-cyclo-octane, 480 Bis-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bimuth oxidised, 289n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Blue oxide of molybdenum cat., 675, Blue oxide of molybdenum cat., 675,		
Bensyl-acetone hydrogenated, 389 Bensyl alcohol, 549, 560, 715, 729, 740 from aldehyde, 225 into amine, 734 eat. solvent, 38 condensed, 728 dehydrated, 688, 714 dehydrogenated, 680, 871, 773 hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 eatalyst, 836 dehydrogenated, 470, 496, 560 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 830, 916 formed, 281, 818, 839 in F. and C. syn., 889 in syn., 899, 901 Bensyl dyndred, 605, 871 Bensyl-cyclohexyl-amine, 739 Bensyl-dyclohexyl-amine, 739 Bensyl-dyclohexyl-amine, 739 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 450 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Blue oxide of molybdenum cat., 675		
Bensyl alcohol, 549, 560, 715, 729, 740 from aldehyde, 225 into amine, 734 est. solvent, 38 condensed, 728 dehydrated, 688, 714 dehydrogenated, 688, 714 dehydrogenated, 686, 714 dehydrogenated, 687, 714, 773 hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 estalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-aniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl choride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl-cyclohexyl-amine, 739 Bensylidene-malonic acid, 804 Bensyl-pyridines syn., 901 Bensyl-pyridines syn., 901 Bensyl-pyridines syn., 901 Bensyl-pyridines syn., 901 Berylia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bi-cyclo-octane hydrogenated, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 299n Bleaching of oils, 285 Blue oxide of molybdenum cat., 675, Blue oxide of molybdenum cat., 675 Blue oxide of molybdenum cat., 675 Blue oxide of molybdenum cat., 675, Boron fluoride polymer. cat., 34, 211 Borneol from acetate, 340 from camphor, 591, 722 dehydrated, 714 dehydrated, 714 dehydrated, 714 dehydrogenated, 661 esters of, 340 oxidised, 257, 260n Bornyl acetate, 340 Brands of hardened fats 967, 967n Brass clat., 254n, 670 Brass block furnace, 348 Broschet's apparatus, 597 Bromal cond., 806 Brombensene formed, 292, 293 hydrogenated, 405, 545 sulphonated, 815 toxic to cats., 115 boron fluoride polymer. cat., 84, 211 dehydrated, 714 dehydrated, 7		
from aldehyde, 225 into amine, 734 cat. solvent, 38 condensed, 728 dehydrated, 688, 714 dehydrogenated, CqO 674, Cu 657, MnO 673 ceterified, 771, 773 hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 catalyst, 836 dehydrogenated, 481 hydrogenated, 470, 496, 560 Bensyl-eniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene-malonic acid, 804 Bensyl-mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 ceterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Brominic cat., 284, 211 Boron fluoride polymer. cat., 34, 211 dehydrogenated, 340 from camphor, 591, 722 dehydrated, 714 dehydrogenated, 661 esters of, 340 oxidised, 257, 260n Bornyl acetate, 340 Brands of hardened fats 967, 967n Brass block furnace, 348 Brassidic acid formed, 184 into ketone, 843 Brochet's apparatus, 597 Bromal cond., 806 Bromanilines red., 405 Brombensene formed, 292, 293 hydrogenated, 405, 545 sulphonated, 815n in syn., 901, 904 Brombensoyl chloride in F. and C. syn., 893 Bromine, cat., 43 chlorination cat., 279 elim, 405, 407, 605 isomerisation cat., 182 toxic to cats., 116, 359 Bromine, cat., 43 chlorination cat., 279 elim, 405, 407, 605 isomerisation cat., 182 toxic to cats., 116, 359 Bromine, 228, 631, 734 catalyst, 836 catelyst, 836 Bornyl acetate, 840 Brands of hardened fats 967, 967n Brass cat., 254n, 670 Brass block furnace, 348 Broschet's apparatus, 597 Bromal cond., 806 Bromanilines red., 405, 405, 545 sulphonated, 815n in syn., 901, 904 Brombensone formed, 292, 293 hydrogenated, 406, 545 sulphonated, 815n in syn., 904 Brombensone formed, 292, 293 hydrogenated, 406, 545 sulphonated, 815n in syn., 904 Brombensone formed, 206 Bromanilines red., 405 Brombensone		
into amine, 734 eat. solvent, 38 condensed, 728 dehydrated, 688, 714 dehydrogenated, CdO 674, Cu 657, MnO 673 esterified, 771, 773 hydrogenated, 389, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 eatalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-aniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 231, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl dyanide, 605, 871 Bensyl dyanide, 605, 871 Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene-malonic acid, 804 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 480 Bis-cyclo-octane, 480 Bis-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Blue oxide of molybdenum cat., 675, Bisoacetic acid hydrol., 332 Blue oxide of molybdenum cat., 675, Bornel from acetate, 340 from acetate, 340 eatleydrated, 714 dehydrogenated, 661 eaters of, 340 oxidised, 257, 280n Bornyl acetate, 340 Brass clat, 254n, 670 oxidised, 257, 280n Bornyl acetate, 340 Brass clat, 254n, 670 Brass clat, 254n, 670 Brass clock furnace, 348 Brassidic acid formed, 184 into ketone, 843 Bromalines red., 405 Bromanilines red., 405 Bromalines red., 405 Bromanilines red.		
cat. solvent, 38 condensed, 728 dehydrated, 688, 714 dehydrogenated, CdO 674, Cu 657, MnO 673 esterified, 771, 773 hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 catalyst, 836 dehydrogenated, 661 hydrogenated, 470, 496, 560 Bensyl-aniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene-malonic acid, 804 Bensyl-midiene-malonic acid, 804 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 480 Bis-cyclo-octene hydrogenated, 480 Bis-cyclo-octene hydrogenated, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Blue oxide of molybdenum cat., 675, Bromless cate, 340 oxidised, 27, 280n benyl acetate, 340 benyl acetate, 340 brom eamphor, 591, 722 dehydrated, 714 dehydrogenated, 361 esters of, 340 oxidised, 257, 280n Bromyl acetate, 340 Brandend fats 967, 967n Brass block furnace, 348 Bromal cond., 806 Bromanilines red., 405 Bromal cond., 806 Bromanilines red., 405 Brombensene formed, 292, 293 hydrogenated, 405, 545 sulphonated, 315n in syn., 901, 904 Bromanilines red., 405 Bromanilines red., 405 Bromanilines red., 405 Bromilian from amphorated, 611 ceters of, 340 oxidised, 257, 280n Bromyl acetate, 340 Brass cat., 254n, 670 Brass block furnace, 348 Bromeletrom exites, 340 Brandend fats 967, 967n Brass cat., 254n, 670 Brass block furnace, 348 Bromeletrom exites, 340 Brandend fats 967, 967n Brass cat., 254n, 670 Brass block furnace, 348 Bromeletrom exites, 940 Brass cat., 254n, 670 Brass block furnace, 348 Bromeletrom exites, 940 Brass cat., 254n, 670 Brass bloc		
condensed, 728 dehydrated, 688, 714 dehydrogenated, 688, 714 dehydrogenated, 771, 773 hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 catalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-aniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene-malonic acid, 804 Bensyl-imercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 480 Bis-cyclo-octane, 480 Bis-cyclo-octane, 480 Bis-cyclo-octane hydrogenated, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, from camphor, 591, 722 dehydrated, 714 dehydrogenated, 661 esters of, 340 oxidised, 257, 260n Bornyl acetate, 340 Brands of hardened fats 967, 967n Brass cat., 254n, 670 Brass block furnace, 348 Brassidic acid formed, 184 into ketone, 843 Bromaline, re70 Brass block furnace, 348 Brassidic acid formed, 184 into ketone, 843 Bromalines red., 405 Bromal cond., 806 Bromanilines red., 405 Brombensone formed, 292, 293 hydrogenated, 405, 545 sulphonated, 31, 81, 816 in syn., 901, 904 Bromsplace, 257, 260n Brands of hardened fats 967, 967n Brass cat., 254n, 670 Brass block furnace, 348 Bromelete, 340 Bromelete, 340 Bromelete, 267 Bromal cond., 806 Bromanilines red., 405 Brombensone formed, 292, 293 hydrogenated, 405, 545 sulphonated, 315 Brombensone formed, 292, 293 hydrogenated, 405, 545 sulphonated, 316 Brombensone formed, 292, 293 hydrogenated, 405, 645 sulphonated, 316 Brombensone formed, 292, 293 hydrogenated, 405, 645 sulphonated, 316 Brombensone formed, 292, 293 hydrogenated, 405, 545 sulphonated, 316 Brombensone formed, 292, 293 hydrogenated, 405, 545 sulphonated, 315n in syn., 901, 904 Brombensone formed, 292, 293 Bromine, cat., 43 Brombensone formed, 292, 293 bromine, cat., 43 Brombensone formed, 292, 405 isomerisat		
dehydrated, 688, 714 dehydrogenated, CdO 674, Cu 657, MnO 673 esterified, 771, 773 hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 eatalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-aniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl-dyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene-malonic acid, 804 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 285 Blue oxide of molybdenum cat., 675,		
dehydrogenated, CdO 674, Cu 657, MnO 673 esterified, 771, 773 hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 estalyst, 836 dehydrogenated, 470, 496, 560 Bensyl-aniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl denne-hydrindone, 799 Bensylidene-hydrindone, 799 Bromitation cat., 279 elim., 405, 407, 605 isomerisation cat., 182 toxic to cats., 116, 359 Bromnitrobensenes hydrogenated, 546 Bromnitrobensenes hydrogenated, 546 Bromnitrobensenes hydrogenated, 546 Bromnitrobensenes, 547 Bromhal caid, 661 Bromal cond., 861 Bromesidic acid formed, 240 Brassidic acid formed, 240 Brombensene fo		• • • . •
estersified, 771, 773 hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 catalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-aniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in F. and C. syn., 889 in syn., 990, 901 Bensyl cyanide, 605, 871 Bensyl dene-hydrindone, 799 Bensylidene-hydrindone, 799 Bromination, 290–293 Bromine, 242, 243 Bromine, 243, 265 Bromine, 244, 25 Bromine, 244, 25 Bromine, 244, 25 Bromine, 244, 25 Bromine, 243, 265 Brominitine, 240 Bro		
oxidised, 257, 260n hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 catalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-aniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl dene acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene-malonic acid, 804 Bensyl mercaptan, 744 Bensyling cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bi-oyclo-octane, 480 Bi-oyclo-octane hydrogenated, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Bensyl acetate, 340 Brands of hardened fats 967, 967n Brass block furnace, 348 Brassidic acid formed, 184 into ketone, 343 Brassidic acid formed, 184 into ketone, 343 Brosmalines red., 405 Brass block furnace, 348 Brassidic acid formed, 184 into ketone, 343 Brosmal cond., 806 Brass block furnace, 348 Brassidic acid formed, 184 into ketone, 343 Brosmal cond., 806 Bromalines of hardened fats 967, 967n Brass cat., 254n, 670 Brass block furnace, 348 Brassidic acid formed, 184 into ketone, 343 Brosmal cond., 806 Bromallines red., 405 Bromal cond., 806 Bromalines red., 405 Bromal cond., 806 Bromalines red., 405 Brassidic acid formed, 184 into ketone, 343 Brosmal cond., 806 Bromalines red., 405 Bromal cond., 806 Bromal cond., 8		
hydrogenated, 369, 465, 538, 593 oxidised, 249 Bensyl amine, 428, 631, 734 catalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-aniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensylidene-acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene-malonic acid, 804 Bensyl mercaptan, 744 Bensylingene-malonic acid, 804 Bensylidene-malonic acid, 804 Brominition cat., 279 elim., 405, 407, 605 isomerisation cat., 116, 359 Bromnation, 290-293 Brominition, 290-293 Bromnition, 290-293 Bromnit		
oxidised, 249 Bensyl amine, 428, 631, 734 catalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-aniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene-hydrindone, 799 Bensylidene-malonic acid, 804 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Brass cat., 254n, 670 Brass block furnace, 348 Brassidic acid formed, 184 into ketone, 343 Broadic acid formed, 184 into ketone, 343 Broadic acid formed, 184 into ketone, 348 Broadic acid formed, 184 i		
Bensyl amine, 428, 631, 734 catalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-bensylidene-acetone, 547 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl dene acetate formed, 240 Bensylidene-malonic acid, 804 Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bi-cyclo-octane, 480 Bi-cyclo-octane, 480 Bis-diassoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Brass cat., 254n, 670 Brass block furnace, 348 Brassidic acid formed, 184 into ketone, 843 Broschet's apparatus, 597 Bromal cond., 806 Bromalines red., 405 Brombensone formed, 292, 293 hydrogenated, 405, 545 sulphonated, 815n in syn., 901, 904 Brombensoyl chloride in F. and C. syn., 893 Bromides cats., 84 Brombensoyl chloride in F. and C. syn., 893 Bromides cats., 84 Brombensone formed, 292, 293 hydrogenated, 405, 545 sulphonated, 815n in syn., 901, 904 Brombensoyl chloride in F. and C. syn., 893 Bromides cats., 84 Brombensone formed, 292, 293 hydrogenated, 405, 545 sulphonated, 815n in syn., 901, 904 Brombensoyl chloride in F. and C. syn., 893 Bromides cats., 84 Brombensone formed, 290, 605 sulphonated, 815n in syn., 901, 904 Brombensoyl chloride in F. and C. syn., 893 Bromides cats., 84 Brombensone formed, 260 Bromanilines red., 405 Brombensone formed, 292, 293 hydrogenated, 405, 545 sulphonated, 815n in syn., 901, 904 Brombensoyl chloride in F. and C. syn., 893 Bromides cats., 84 Brombensoyl chloride in F. and C. syn., 893 Bromides cats., 84 Brombensoyl chloride in F. and C. syn., 893 Bromides cats., 84 Brombensoyl chloride in F. and C. syn., 893 Bromides cats., 84 Brombensoyl chloride in F. and C. syn., 893 Bromides cats., 84 Brombensoyl chloride in F. and C. syn., 893 Bromides cats., 84 Brombensoyl chloride in F. and C. syn., 893 Bromides		
catalyst, 836 dehydrogenated, 681 hydrogenated, 470, 496, 560 Bensyl-anilline, 729 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl cyanide, 605, 871 Bensyl formate, 773 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene-hydrindone, 799 Bensylidene-malonic acid, 804 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675,		
dehydrogenated, 470, 496, 580 Bensyl-aniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene-malonic acid, 804 Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bi-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675,		
hydrogenated, 470, 496, 560 Bensyl-aniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl-cyclohexyl-amine, 739 Bensyl-cyclohexyl-amine, 739 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene-malonic acid, 804 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675,		
Bensyl-aniline, 729 Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl-cyclohexyl-amine, 739 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensyl mercaptan, 744 Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675,		
Bensyl-bensylidene-acetone, 547 Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl-cyclohexyl-amine, 739 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene-malonic acid, 804 Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675,		
Bensyl chloride dechlorinated, 605 decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene-malonic acid, 804 Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Bi-cyclo-oxide of molybdenum cat., 675, Bi-cyc		
decomposed, 880, 916 formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675,		<u> </u>
formed, 281, 818, 889 in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, polym., 213 hydrogenated, 405, 545 sulphonated, 815n in syn., 901, 904 Brombensoyl chloride in F. and C. syn., 893 Bromines cats., 84 Bromination, 290-293 Bromine, cat., 43 chlorination cat., 279 elim., 405, 407, 605 isomerisation cat., 182 toxic to cats., 116, 359 Bromnaphthalenes isom., 199 Bromnstyrene hydrogenated, 405, 512 Bromstyrene hydrogenated, 546 Brucine hydrogenated, 555 Butadiene formed, 726, 784 polym., 213		
in F. and C. syn., 889 in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, polym., 213 sulphonated, 815n in syn., 901, 904 Brombensoyl chloride in F. and C. syn., 893 Bromines cats., 84 Bromination, 290-293 Bromine, cat., 43 chlorination cat., 279 elim., 405, 407, 605 isomerisation cat., 182 toxic to cats., 116, 359 Bromnaphthalenes isom., 199 Bromnstyrene hydrogenated, 405, 512 Bromstyrene hydrogenated, 546 Brucine hydrogenated, 555 Butadiene formed, 726, 784 polym., 213		
in syn., 899, 901 Bensyl cyanide, 605, 871 Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene-malonic acid, 804 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, 891 Bensyl cyanide, 605, 871 Brombensoyl chloride in F. and C. syn., 893 Bromides cats., 84 Bromination, 290-293 Bromine, cat., 43 chlorination cat., 279 elim., 405, 407, 605 isomerisation cat., 182 toxic to cats., 116, 359 Bromnaphthalenes isom., 199 Bromnaphthalenes isom., 199 Bromnstyrene hydrogenated, 405, 512 Bromstyrene hydrogenated, 546 Brucine hydrogenated, 555 Butadiene formed, 726, 784 polym., 213		
Bensyl cyanide, 605, 871 Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene-malonic acid, 804 Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Bensyl-cyclohexyl-amine, 739 Brombensoyl chloride in F. and C. syn., 893 Bromides cats., 84 Bromination, 290-293 Bromine, cat., 43 chlorination cat., 279 elim., 405, 407, 605 isomerisation cat., 182 toxic to cats., 116, 359 Bromnaphthalenes isom., 199 Bromnitrobensenes hydrogenated, 405, 512 Bromstyrene hydrogenated, 546 Bromtoluene, 293, 405 Brucine hydrogenated, 555 Butadiene formed, 726, 784 polym., 213	• • • • • • • • • • • • • • • • • • • •	•
Bensyl-cyclohexyl-amine, 739 Bensyl formate, 773 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene-malonic acid, 804 Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Bensyl-dydrindone, 799 Bromination, 290–293 Bromine, cat., 43 chlorination cat., 279 elim., 405, 407, 605 isomerisation cat., 182 toxic to cats., 116, 359 Bromnaphthalenes isom., 199 Bromnitrobensenes hydrogenated, 405, 512 Bromstyrene hydrogenated, 546 Bromstyrene hydrogenated, 546 Brucine hydrogenated, 555 Butadiene formed, 726, 784 polym., 213	Th. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.	
Bensyl formate, 773 Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene-malonic acid, 804 Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Bensylidene-hydrindone, 799 Bromination, 290-293 Bromine, cat., 43 chlorination cat., 279 elim., 405, 407, 605 isomerisation cat., 182 toxic to cats., 116, 359 Bromnaphthalenes isom., 199 Bromnitrobensenes hydrogenated, 405, 512 Bromstyrene hydrogenated, 546 Bromstyrene hydrogenated, 546 Brucine hydrogenated, 555 Butadiene formed, 726, 784 polym., 213		
Bensylidene acetate formed, 240 Bensylidene-hydrindone, 799 Bensylidene-malonic acid, 804 Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Bensylidene-hydrindone, 799 Bromination, 290-293 Bromine, cat., 43 chlorination cat., 279 elim., 405, 407, 605 isomerisation cat., 182 toxic to cats., 116, 359 Bromnaphthalenes isom., 199 Bromnitrobensenes hydrogenated, 405, 512 Bromstyrene hydrogenated, 546 Bromstyrene hydrogenated, 546 Brucine hydrogenated, 555 Butadiene formed, 726, 784 polym., 213		
Bensylidene-hydrindone, 799 Bensylidene-malonic acid, 804 Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Bromine, cat., 43 chlorination cat., 279 elim., 405, 407, 605 isomerisation cat., 182 toxic to cats., 116, 359 Bromnaphthalenes isom., 199 Bromnitrobensenes hydrogenated, 405, 512 Bromstyrene hydrogenated, 546 Brucine hydrogenated, 546 Brucine hydrogenated, 555 Butadiene formed, 726, 784 polym., 213		
Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Bensyl-pyridines syn., 901 Bis-diasoacetic acid, 675, 676, 702, 778, 828 isomerisation cat., 182 toxic to cats., 116, 359 Bromnaphthalenes isom., 199 Bromnitrobensenes hydrogenated, 405, 512 Bromstyrene hydrogenated, 546 Bromtoluene, 293, 405 Brucine hydrogenated, 555 Butadiene formed, 726, 784 polym., 213		
Bensyl mercaptan, 744 Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Belsimuth oxide of molybdenum cat., 675,		
Bensyl-pyridines syn., 901 Beryllia cat., 651, 675, 676, 702, 778, 828 esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Bromerisation cat., 182 toxic to cats., 116, 359 Bromnaphthalenes isom., 199 Bromnitrobensenes hydrogenated, 405, 512 Bromstyrene hydrogenated, 546 Bromtoluene, 293, 405 Brucine hydrogenated, 555 Butadiene formed, 726, 784 polym., 213		
esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bi-cyclo-octane hydrogenated, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Betulol hydrogenated, 570 Bromnitrobensenes hydrogenated, 405, 512 Bromstyrene hydrogenated, 546 Bromtoluene, 293, 405 Brucine hydrogenated, 555 Butadiene formed, 726, 784 polym., 213	Benzyl-pyridines syn., 901	
esterific. cat., 778 Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bi-cyclo-octane hydrogenated, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Betulol hydrogenated, 570 Bromnitrobensenes hydrogenated, 405, 512 Bromstyrene hydrogenated, 546 Bromtoluene, 293, 405 Brucine hydrogenated, 555 Butadiene formed, 726, 784 polym., 213		
Betulol hydrogenated, 570 Bi-cyclo-octane, 480 Bi-cyclo-octane hydrogenated, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Bromnitrobensenes hydrogenated, 405, 512 Bromstyrene hydrogenated, 546 Bromtoluene, 293, 405 Brucine hydrogenated, 555 Butadiene formed, 726, 784 polym., 213		
Bi-cyclo-octane, 480 Bi-cyclo-octene hydrogenated, 480 Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Bi-cyclo-octane, 480 512 Bromstyrene hydrogenated, 546 Bromtoluene, 293, 405 Brucine hydrogenated, 555 Butadiene formed, 726, 784 polym., 213	Betulol hydrogenated, 570	Bromnitrobensenes hydrogenated, 405,
Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Blue oxide of molybdenum cat., 675,	5. •	512
Bis-diasoacetic acid hydrol., 332 Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Blue oxide of molybdenum cat., 675,		Bromstyrene hydrogenated, 546
Bismuth oxidised, 269n Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Blue oxide of molybdenum cat., 675,		
Bleaching of oils, 265 Blue oxide of molybdenum cat., 675, Butadiene formed, 726, 784 polym., 213		
Blue oxide of molybdenum cat., 675, polym., 213	Bleaching of oils, 265	Butadiene formed, 726, 784
732, 827 Butaneal-ol (1, 3) formed, 219	Blue oxide of molybdenum cat., 675,	
	732, 827	Butaneal-ol (1, 3) formed, 219

Cadmium sulphate cat., 626
Calcium carbonate in acetone prep., 161
carrier for cat., 127
catalyst, 98
ketone cat., 161, 838, 839, 849, 857
to neut. oils, 848
Calcium hydroxide toxic to cats., 115
Calcium oxide cat., 83, 98
decomp. hydrocarb., 911
ketone cat., 840, 849
Calcium salts cats., 269n
decomposed, 837
Calcium sulphate cat., 98, 687, 718
Camphane, 477, 552, 594, 611, 722
Camphane from borneol, 722
dehydrogenated, 644
hydrogenated, Cu 594, Ni 477, 591,
Pd 552, Pt 570
Campholide by hydrogenation, 392
Camphor from borneol, 260n, 269n, 661
hydrogenated, 591, 722
by oxidation, 257, 260n, 269n
Camphor acids, 836
Camphor-hyrasone dec., 611
Camphoric acid by oxid., 257
Camphoric anhydride hydrogenated,
392
Camphorone hydrogenated, 421
Camphor-oxime hydrogenated, 385
Cane sugar oxidised, 269
Candelite, 967
Caoutchouc, syn. of, 214, 215
Caproic acid into aldehyde, 853
esterified, 771
into ketone, 845
Caproic aldehyde, 853
Caproic esters decomp., 871
Caprylic acid into ketone, 843
Caprylene hydrogenated, 414
Carbamic chloride in F. and C. syn.,
905
895
Carbasol hydrogenated, 490
Carbazol hydrogenated, 490 Carbides in earth, 928
Carbasol hydrogenated, 490 Carbides in earth, 928 Carbimides, 431
Carbasol hydrogenated, 490 Carbides in earth, 928 Carbimides, 431 Carbohydrates hydrogenated, 595
Carbasol hydrogenated, 490 Carbides in earth, 928 Carbimides, 431 Carbohydrates hydrogenated, 595 Carbon catalyst, 48, 49, 257, 257n,
Carbasol hydrogenated, 490 Carbides in earth, 928 Carbimides, 431 Carbohydrates hydrogenated, 595 Carbon catalyst, 48, 49, 257, 257n, 687, 700, 811, 828, 911
Carbasol hydrogenated, 490 Carbides in earth, 928 Carbimides, 431 Carbohydrates hydrogenated, 595 Carbon catalyst, 48, 49, 257, 257n, 687, 700, 811, 828, 911 cat. dec. alcohols, 679
Carbasol hydrogenated, 490 Carbides in earth, 928 Carbimides, 431 Carbohydrates hydrogenated, 595 Carbon catalyst, 48, 49, 257, 257n, 687, 700, 811, 828, 911 cat. dec. alcohols, 679 dehydrogenation cat., 638, 679
Carbasol hydrogenated, 490 Carbides in earth, 928 Carbimides, 431 Carbohydrates hydrogenated, 595 Carbon catalyst, 48, 49, 257, 257n, 687, 700, 811, 828, 911 cat. dec. alcohols, 679

separated, 613 hydrogenated, 968 Carbon dioxide eliminated, 831 iodine no., 938 hydrogenated, Co 504, Cu 508, Ni Catalysis, definition, 1, 3, 4, 140 395, 396, 586, Pt 533 at a distance, 180j history of, 4 Carbon disulphide chlorinated, 283 eliminated from ill. gas, 339, 372 mechanism of, 129-180u on F. and C. svn. 892, 893, 897 Catalysts, amounts of, 32, 951 hydrogenated, 372, 492 for cracking, 906, 910-912 hydrolyzed, 339 dehydration, 687 hydrogenation, 598, 941 negative cat., 303 life of, 111, 359, 947 reduced, 372 into thioureas, 630 orienting, 816 toxic to cats., 116 placing in tubes, 128 Carbon hexachloride formed, 289, 881 poisons for, 112, 113, 116, 359, 946 Carbon monoxide, 508, 593, 613-617, preparation, 54-56, 58, 59, 76-78, 821, 825-828, 866-869, 953 598, 606, 655, 704, 705, 707n, added, 298 861n. 941. 942 decomposed, 163, 614 regeneration of, 947n, 950 eliminated, 618-625 Caucasian petroleum, 926 from formic acid, 143, 172 Cellulose hydrolyzed, 323 in F. and C. syn., 297, 298 Ceria in drying oils, 266 hydrogenated, 540, Co 504, Ni 393, ketone cat., 849 oxidation cat., 259, 261 394, Pd 536, Pt 533 in hydrogenation, 953 promoter, 180¢ oxidised, 150, 180a, 248, 251 Cerium compounds cats., 153, 264, source of H in hydrogenation, 537 269n, 271 toxic to cats., 116, 180e Charcoal as absorbent, 180d, 180d Carbon suboxide, 873 animal as carrier, 946 Carbon tetrachloride added, 242 animal as cat., 282n, 700 formed, 180b, 278, 279, 282-285, carrier for, Ni 598, 941, 946 catalyst, 828 Carbylamines hydrogenated, 430, 521 chlorination cat., 282 hydrolyzed, 334 cat. for phosgene, 282n Carbonyl chloride in F. and C. syn., condenses gases, 131, 135 891, 893 dec. hydrocarb., 911 polymerize alds., 222 Chemical potential, 180g Chemical theory of cat., 145 preparation, 134, 282, 282n Carbonyl group hydrogenated, 432-442 Chloracetanilid, 202 Carboxy-camphor acids, 836 Chloracetates reduced, 407 Carriers for cats., 126, 941, 946 Chloracetic acid, 280, 281 Chloral cond., 239, 806 Carvacrol hydrogenated, 459 polymer., 224, 228 Carvacrylene, 789 Carvacryl ethers, 786, 789 Chloranilines, 512, 632 Carvomenthane, hydrogenated, 570 Chlorates oxidisers, 271 Carvomenthol, 567, 722 reduced by Pd. 165 Chlorbensenes formed, 278, 284, 285, Carvomenthone, 591 Carvone hydrogenated, 476, 552, 567, 404 sulphonated, 815n 591 Carvotenacetone, 567 in syn., 904 Chlorbensoic acid hydrogenated, 545 Caryophylene hydrogenated, 560 Castor oil in alcoholysis, 341 in syn., 901

Chlorbensoyl chloride in F. and C.	oxidation cat., 259
syn., 893 ω-Chlorbutyl-bensene in F. and C.	preparation of, 78 Cincheniding hydrogeneted Pd 555
	Cinchonidae hydrogenated, Pd 555
syn., 897 Chloreffeine hydrogeneted 545	Cinchonine hydrogenated, Pd 555, Pt 561
Chloreinnemie as hydrogeneted 245	
Chlor compounds by drolysad 220	Cincol debydrogeneted 645
Chlorestonia acid hydrogeneted 545	Cincol dehydrogenated, 645
Chloreredo horano 402	Cinnamene hydrogenated, 451
Chlorethal honome in F and C	Cinnamic acid esterified, 756, 757
β-Chlorethyl-benzene in F. and C.	formed, 107, 246
syn., 897	hydrogenated, 417, 581, 583, 604,
Chlorides cats., 84 et seq.	Cu 594, Ni 590, 601, Ru 580
Chlorides chlorination cats., 278	by hydrogenation, 548
oxidation cats., 263	Cinnamic alcohol oxidised, 249
Chlorination, 44, 58, 90, 156, 278–289	Cinnamic aldehyde from alcohol, 246
of acetic acid, 280	condensed, 799
catalysts, 283–285	hydrogenated, Pd 546, Pt 568, 560
Chlorine absorbed by C, 180b	by oxidation, 249
catalyst, 43	Citraconic acid hydrogenated, Pt 558
eliminated, 403, 404, 407, 605	isomerised, 183
produced, 103	Citral condensed, 800
toxic to cats., 359, 947	from geraniol, 658
on water, 257n	hydrogenated, Pd 595, Pt 567
Chlorketones produced, 243	Citric acid esterified, 756
Chlormethyl ethers cond., 818	retarder, 11
in F. and C. syn., 889, 899	toxic to cats., 115
Chlor-nitrobensenes hydrogenated, 404,	Citronellol, 416
512	Class of alcohol determined, 701
in syn., 901	Clay dehydration cat., 99, 700, 702, 717
Chlor-nitrobensoic acid in syn., 901	Clarifying solutions, 257n
Chloroform cond., 238	Clupadonic acid, 937, 938, 955
formed, 629, 879	esters of, 937
in F. and C. syn., 890	Coal oxidation cat., 257
negative cat., 11, 238	Cobalt on alcohols, 666
stabilised, 11	catalyst, 57, 167, 615
in syn., 890, 903	in cracking, 906
ω-Chlorpentyl-benzene in F. and C.	decomp. C ₂ H ₂ , 919, 920, 928
syn., 897	decomp. aromat. hydrocarb., 921
Chlorphenols by reduction, 404	decomp. hydrocarb., 906, 912
Chlorpicrin, 180q	dehydrogenation cat., 637, 651, 652,
Chlortoluene hydrogenated, 569	666
Cholesterine hydrogenated Pt, 565	deterioration of, 500
Chromic chloride, 357	hydrogenation cat., 344, 499-504, 945
Chromic oxide, cat., 75, 675, 676, 693,	in drying oils, 266
703, 732, 746, 791, 840, 849	oxidation cat., 254, 258
dehydration cat., 702, 791	Cobalt carbonyl, 616
dehydrogenation cat., 686	Cobalt chloride cat., 283, 876
in drying oils, 266	Cobalt oxide, oxidation cat., 75, 180a,
ketone cat., 840, 849	259
mercaptan cat., 746	Cobalt soap, 265
mixed cat., 702	Cocoa butter, alcoholysis of, 341

hardened, 966	in Deacon's proc., 103
iodine number, 938	Cottonseed oil, iodine number, 938
Cocoanut oil, hydrogenated, 939	hydrogenated, 587, 965, 967n
Codeine, hydrogenated, Pt 572	Coumanic acid dec., 835
Codliver oil, hardened, 966	Cracking, 906, 910-912, 929-936
iodine number, 938	by AlCl ₂ , 929–931, 935
Coke as catalyst, 48, 257	by cats., 910–912, 932, 934
Colchicine hydrogenated, Pd 555	discovery, 906
Collidines, 686	with oxide cats., 934
Collisions of molecules, 180e, 180f	Cresol ethers hydrogenated, 494
Colloidal metals, preparation of, 67	Cresols with aldehydes, 792
Colloidal palladium, 544–555, 604	ethers of, 386, 785, 786, 789
Colloidal platinum, 544, 556-561	formed, 386, 645, 660
Colophene from pinene, 216	hydrogenated, 457, 464
Colsa oil, 938	by oxidation, 263
Complex rings hydrogenated, Pt 571	Cresyl-carvacryl ether, 788
Condensation of aldehydes, 794–800	Cresyl-diamines by hydrogenation, 380
of ketones, 794–800	Cresyl oxides, 785
Coniferine hydrol., 329	Cresyl-propanes hydrogenated, 415
Coniferyl alcohol by hydrol., 329	Crisco, 967n
Contact process, 180r, 258	Crotonic acid into aldehyde, 853
Contra-valencies, 180h	esterified, 756, 771
Copper catalyst, 59, 269n, 538-540,	hydrogenated, 422, Pd 546, Pt 558
683, 692, 824, 831, 833-835, 901,	Crotonic aldehyde, 307, 794-796, 801
904, 920-922	condensed, 796
on alcohols, 142	formed, 52, 219, 308, 310
colloidal, 72	hydrogenated, 419, Pt 567
in cracking, 906, 932	Crotonisation, 81, 89, 107, 794-800
decomp. C ₂ H ₂ , 913, 916, 917, 920	of aldehydes, 795
decomp. aldehydes, 621	of ketones, 797
decomp. CO, 615	Crotonylene polym., 212
decomp. formic esters, 867	Cumene, 644
decomp. hydrocarb., 921	Cuminic aldehyde by oxidation, 249
decomp. pinene, 922	polymerized, 220
dehydrogenates amines, 681	Cuminoine, 220
dehydrogenation cat., 142, 636, 637,	Cuminyl alcohol oxid., 249
641, 646, 651-654, 656-663, 680,	Cuprene, 518, 916–918
681, 701, 720, 726, 824	Cupric hydroxide purif. of oils, 948
on diazo-comps., 606-610	Cuprous bromide cat., 608, 611, 633
hydrogenation cat., 344, 507-523,	in diaso reaction, 91
594, 939, 945	prep., 608
isomer. cat., 208	Cuprous chloride cat., 298, 611, 633,
oxidation cat., 15, 75, 162, 167,	879, 901
253, 254, 258	in diaso reaction, 91
preparation, 59, 606, 655	Cuprous iodide cat., 611, 901
Copper chloride cat., 635	in diaso reaction, 91
Copper oxide cat., 259, 269n	Cuprous oxide in diaso., 91
Copper powder cat., 606-610, 655	Cuprous salts cats., 611, 633
Copper salts in nitration, 269n	in diago reaction, 606-610
in oxidation, 271	Cyanethine, 232
Copper sulphate cat., 240, 272, 725	Cyanides cats., 95
Programme and and and and	-,,

C 1 1 4 1 000	O 11 11 11 110
Cyanogen hydrated, 312	Cyclohexendiol ether, 443
Cyaphenine, 232	Cyclohexenes by dehydration, 698
Cyclic acids dec., 830	dehydrogenated, 643
Cyclic hydrocarb. dec., 921	Cyclohexanol by oxidation, 251
polym., 216	Cyclohexenone hydrogenated, Pd 552
Cyclic ketones, 611	Cyclohexyl-acetic acid, 471, 476
Cyclic ketoximes, 205	Cyclohexyl alcohols dehydrogenated,
Cyclisation, 82, 194	714, 717
Cyclo-aliphatic ethers, 494	esterified, 757, 766
Cyclobutane, 473	Cyclohexyl-amine, 466, 469, 497, 560,
Cyclobutene hydrogenated, 473	569, 737, 739
Cyclobutene bromide, 293	dehydrogenated, Ni 642
Cycloheptane, 197, 479, 649	by hydrogenation, 378, 385
Cyclohexadienes dehydrogenated, 643	Cyclohexyl-amines, 739
Cyclohexadiols, 461	
	Cyclohexyl beneate 766
Cyclo-hexadione, 874	Cyclohexyl bensoate, 766
Cyclo-hexamethylene ring, 475	Cyclohexyl chlorides decomp., 876
Cyclohexane, 445, 452, 466, 468, 469,	in F. and C. syn., 889
471–475, 497, 534, 560, 587, 589,	Cyclohexyl-cyclohexene hydrogenated,
611, 643	475
decomp., 921	Cyclohexyl-diethyl-amine, 468
dehydrogenated, Fe 593, Ni 641	Cyclohexyl-ethyl-amine, 468
formed, 26, 55, 113, 361, 388, 389,	Cyclohexyl-heptane by hydrogenation,
408	414
oxidised, 251	Cyclohexyl mercaptan, 628, 745
prepared, 446	Cyclohexyl-methyl-amine, 468
purified, 446	Cyclohexyl oxide, 589
Cyclohexane alcohols, 698, 737	Cyclohexyl piperidine, 741
Cyclohexane hydrocarb. dehydrogen-	Cyclohexyl-propionic acid, 471, 580
ated, 641	581, 590
formed, 389	Cyclohexyl-propyl alcohol, 560
Cyclohexane petroleums cracked, 934	Cyclohexyl sulphide, 628
Cyclohexanol, 30, 120, 443, 460, 461,	Cyclo-octadiene, 480
560, 569, 589, 603, 739, 741	Cyclo-octane dec. Ni, 197
into amine, 737	by hydrogenation, 480, 571
dehydrated, 714	Cyclo-octanone, 571
dehydrogenated, 642, 660	Cyclo-octatetrene hydrogenated, 535,
Cyclohexanol homologs dehydrogenated,	571
642	Cyclo-octatriene hydrogenated, 571
	Cyclo-octenone hydrogenated, 571
Cyclohexanone, 456, 560, 642, 660	
crotonized, 797	Cyclo-paraffine oximes hydrogenated,
hydrogenated, 436, 567	385
hydrasone, 611	Cyclopentadiene hydrogenated, 474
oxime hydrogenated, 385	Cyclopentane, 436, 474, 649
Cyclohexatriol, 462	Cyclopentane-carbonic acid, 649
Cyclohexene, 456, 475, 515, 628, 643,	Cyclopentanol, 436
698, 714	Cyclopentanone, 874
dehydrogenated, 643	hydrogenated, 436, 567
hydrogenated, 587	oxime hydrogenated, 385
Cyclohexene acetic acid hydrogenated,	Cyclopentyl-amines by hydrogenation,
476	385

Cyclopental-bensene by F. and C. syn., 897

Cyclopentyl chlorides dec., 876

Cyclopentyl-cyclopentanone, 436

Cyclopropane hydrogenated, 472

Cyclopropane ring dec., 193

Cymene dec. by AlCl₂, 930

by dehydrogenation, 844, 645

hydrogenated, 448

from pinene, 922

Cymenes by hydrogenation, 369, 415

Deacon's process, 103, 180b, 257n Decahydro-acenaphthene, 563 Decahydro-anthracene, 592 Decahydro-fluorene, 454 dehydrogenated, 642 Decahydro-naphthalene, 481, 553, 571, Decahydro-naphthols, 481, 592, 714 Decahydro-quinaldine, 488 Decahydro-quinoline, 488, 555, 561, 592 Decane, 595 Decanol, 595 Decarbonization of CO, 614 Decomposition of acids, 820–856 Decomposition of esters, 858-872 Decomposition and cond. of hydrocarb., 905-936 Decompositions by Ni, 493 Decyclisations, 193 Dehydration, 687-727, 728-784, 785-816, 825 of alcohols, 138, 169, 688-727 of alcohols with acids, 747 of alcohol with alds., 779 of alcohols with amines, 729 of alcohols with ammonia, 729 of alcohols with hydrocarb., 728 of alcohols with hydrogen sulphide, 743 of alcohols with ketones, 779 of aldehydes, 794 of alds. with ammonia, 807 of alds. with hydrogen sulphide, 810 of alds. with ketones, 798 by alumina, 713 of amides, 811 apparatus, 717 of benshydrol, 720 by beryllia, 778

catalysts, 538, 651, 675, 687, 702, 825 in gas phase, 693, 694, 700-727, 731, 801 of glycerine, 760 with hydrogenation, 721, 722 by iodine, 699 of ketones, 797 in liquid medium, 691, 692, 695-699, by metal oxides, 702, 763 by mineral acids, 696, 749 of oximes, 814 of phenols, 785-793 of phenols with alcohols, 789 of phenols with amines, 790 of phenols with hydrogen sulphide, 791 of poly-alcohols, 723, 727 with ring formation, 727 theory of, 169, 689 by thoria, 716 by titania, 767 et seq. by sinc chloride, 698 Dehydroacetic acid formed, 387 Dehydrogenation, 15, 638-686, 807-809, 824, 910, 921 of alcohols, 31, 650-679 by aluminum chloride, 685 of amines, 681, 682 of anthracene hydrides, 642 apparatus, 654 by cadmium oxide, 674 by carbon, 679 catalysts, 636-638, 651, 675, 702, 824 classification, 638 by cobalt, 666 by copper, 653-663 in cracking, 910 of cyclohexane comps., 641 history of, 636 of hydro-aromatic hydrocarb., 640-649 of hydrocarbons, 639-649, 921 of hydrocyclic hydrocarbons, et seq. by iron, 667 by manganous oxide, 672 of methyl alcohol, 676 of naphthalene hydrides, 642 by nickel, 664, 684

by blue oxide of tungsten, 715

by oxides, 672-675, 686 by palladium, 649, 669 of piperidine, 647 by platinum, 668 of poly-alcohols, 680 of second. amines, 682 by stannous oxide, 673 of terpenes, 643 et sea. of tertiary amines, 682 theory of, 168 by various oxides, 675 by sinc. 670 Dehydromucic acid. 727 Dekalin, 481n Depolymerisation, 234, 235 Desoxybensoine hydrogenated, 389 Dextrine by hydrolysis, 326 hydrolyzed, 323, 326 Diacetonitrile, 230 Diacetonyl alcohol dehydrated, 698, 699 Diacetyl hydrogenated, 438 Diacetyl-dihydromorphine, 555 Diacetyl-morphine hydrogenated, 555. Pt 561 Diamines by hydrogenation, 380 Diamylene formed, 210, 211 Diastase, 758 Diazoacetic ester. 12 Diazobenzene, 59, 206, 606, 607 Diaso-compounds decomp., 59, 606-610 hydrogenated, 497 Dibasic acid chlorides in F. and C. syn., Dibasic acids decomposed, 855 esters of dec., 872-874 Dibensal-acetone, 798 Dibensovl hydrogenated, 391 Dibensyl hydrogenated, 452, 589 by hydrogenation, 389, 391, 415, 548, 590, 593 Dibenzyl-acetone, 547 Dibenzyl-amine, 428, 734 Dibensyl-aniline, 729 Dibenzyl-benzene, 728 Dibensylidene-acetone hydrogenated, 547 Dibensyl ketone hydrogenated, 455 Dibromethylene in F. and C. syn, 890 Dibrom-succinic acid, 182 Dibutyl ketone, 844 Dichloracetyl chloride dec., 625

Dichlorbensenes, 404 Dichlorevelohexane dec., 876 Dichlorethylene, 242 Dicyanamide formed, 233 Dicyanides hydrogenated, 429 Dicyclohexvl, 452, 475, 589 Dicyclohexyl-amine, 466, 469, 497, 569, 590, 642, 739 by hydrogenation, 385, 739 Dicyclohexyl-butanes, 452 Dicyclohexyl-ethanes, 452, 589 Dicyclohexyl-methane, 389, 453, 560 Dicyclohexyl-phenyl-methane, 453 Dicyclohexyl-propane, 455 Dicyclononane, 454 Diethyl-allylene formed, 50, 192 Diethyl-amine by hydrogenation, 377, 383, 386, 427 Diethyl-amine. HCl catalyst, 783 Diethyl-aniline dec. Ni, 634 formed, 729 hydrogenated, 468 Diethyl-benzene, 888, 930 Diethyl-carbinol into amine, 735 Diethyl-diphenyl formed, 241 Diethylene compounds hydrogenated, 547 Diethylenic acids, 937 Diethyl ketone formed, 838 hydrogenated, 435, 567 Diethyl-phenol hydrogenated, 459 Dihalogen compounds in F. and C. syn., 890 Diheptene, 519 Dihexahydrobenzyl-amine, 470 Dihydrobensene, 723, 876 Dihydrobrucine, 555 Dihydrocamphene, 570 Dihydrocamphorone, hydrogenated, 390 by hydrogenation, 421 Dihydrocinchonidine, 555 Dihydrocitronellol by hydrogenation, 416 Dihydrocodeine, 572 Dihydro-dimethyl-anthracene, 890 Dihydro-eugenol, 577 Dihvdro-indol. 571 Dihydro-ionones, 554 Dihydrolimonene, 517, 591 Dihydromorphine by hydrogenation, 572

by oxidation, 268	Di
Dihydronaphthalene, 571, 931	Di
Dihydrophytol, 565	
Dihydroquinine, 572	Di
Dihydropinene, 477, 570	Di
Dihydrophenanthrene, 484, 571, 592	Di
Dihydrophorone, 547	Di
Dihydrosafrol, 418, 565, 590	Di
Dihydrostrychnine, 555	Di
Dihydrotetrasines isom., 201	
Dihydroxy-acetone, 237, 246, 268	Di
Dihydroxy-diphenyl-amine, 632	Di
Di-isoamyl-amine, 682, 733	Di
Di-isobutyl-carbinol, 549, 567	Di
Di-isobutyl ketone, 435, 567, 840	Di
hydrogenated, 435	Di
Di-isopropyl-amine, 735	Di
Di-isopropyl-bensene from cymene, 930	Di
Di-isopropyl ketone formed, 844	Di
hydrogenated, 435	Di
Diketones by dehydrogenation, 663	Di
by F. and C. syn., 893	Di
hydrogenated, 391, 438-440	Di
Dimethyl-acetylene formed, 192	Di
Dimethyl-acrylic acid hydrogenated,	Di
417	Di
Dimethyl-allene formed, 192	
Dimethyl-allyl-carbinol hydrogenated,	Di
587	
Dimethyl-amine, 377, 430	Di
Dimethyl-aniline, 468, 729, 740	
dec. Ni, 634	
oxidised, 256	
Dimethyl-bensaldehyde, 298	Di
Dimethyl-butadiene, 726	
Dimethyl-butyl-phenol, 459	
Dimethyl-cyclohexane, 197, 449, 475,	
480	
dehydrogenated, Ni 641	
Dimethyl-cyclohexanols, 458, 660, 714	Di
Dimethyl-cyclohexene by dehydration	Di
714	\mathbf{D}_{i}
hydrogenated, 475	
Dimethyl-cyclohexyl-amine, 467	Di
Dimethyl-cyclopentyl-pentanones, 436	
Dimethyl-diethyl-butine-diol, 548	
Dimethyl-diphenyl-butine-diol, 548, 566	D
Dimethyl-diphenyl-methane hydrogen-	D
ated, 452	D
Dimethyl-ethyl-carbinol esterified, 757	Di
formed, 210, 306	D
• •	

imethylene-pentane hydrogenated, 414 methyl-heptane by hydrogenation, 414 imethyl-hexine hydrogenated Pd, 548 imethyl-hexine-diol hydrogenated, 566 imethyl-indol, 490, 633 imethyl-isobutyl-cyclohexane, 449 imethyl-ketazine isom., 196 imethyl-methylene-cyclopropane hydrog., 472 imethyl-octane, 415, 567 imethyl-octanol, 416, 567 imethyl-octadieneol, 416 imethyl-octatriene hydrogenated, 415 imethyl-octene-diol hydrogenated, 548 imethyl-pentane-thiol, 745 imethyl-phenols hydrogenated, 458 imethyl-propyl-carbinol, 587 imethyl-quinoline, 491 imethyl-toluidines, 684, 740 inaphthyl dehydrogenated, 685 inaphthyl-amine, 632 imtrobensenes, 269n, 512 imtro-compounds hydrogenated, 380 initro-toluenes hydrogenated, 380 ipentene depolymerized, 235 formed, 198 iphenols, ethers of syn., 904 reduced, 370 iphenyl formed, 907 in F.and C. syn., 896 hydrogenated, 452, 589 by hydrogenation, 403, 406 iphenyl-amine from aniline, 466 by dehydration, 642 hydrogenated, 469, 590 stabilizer, 13 sulphurized, 296 syn. of, 901 iphenyl-anthrone syn. of, 893 iphenyl-benzene formed, 907 iphenyl-butadione by F. and C. syn., iphenyl-butanes by hydrogenation, 520, 548 hydrogenated, 452 iphenyl-butadiene, 548 iphenyl-butenes hydrogenated, 415 iphenyl-butine-diol hydrogenated, 548 iphenyl-cyclopropane, 611 iphenyl-decadiene hydrogenated, 545

Elimination of ammonia, 631-633 Diphenyl-decane, 546 Diphenyl-diacetylene hydrogenated, 548 of aniline, 634 Diphenylene oxides, 787 of carbon, 613 Diphenyl-ethanes formed, 241, 728, 890 of carbon monoxide, 618-625 by hydrogenation, 391, 415, 721, 728 of halogens, 605 of hydrogen sulphide, 626-629 hydrogenated, 452 Diphenyl-ethylene formed, 890 of nitrogen, 606-610 hydrogenated, 415, 515 Ellis's apparatus, 962 Diphenyl-methane, 369, 389, 523, 538, Emulsine, 18, 327, 329 539, 590, 662, 720, 728, 806 Ensymes, 180e Diphenyl-pentanes hydrogenated, 452 Equilibrium in alcoholysis, 340n Diphenyl-pentens hydrogenated, 415 shifted, 180s Diphenyl-propane formed, 728 Erdmann's apparatus, 958 by hydrogenation, 389, 415 Erucic acid, 184 hydrogenated, 452 esters of, 937 Diphenyl-propenes hydrogenated, 415 Erythrol formed, 83 Diphenyl sulphide, 629 Esterification, 747–778 Diphenyl thio-urea, 630 by acid anhydrides, 761 Diphenyl-pyrazoline dec., 612 of bensoic acid, 758 by beryllia, 778 Diphenyl urea, 495 catalytic, 17, 747-778 Diphthalid formed, 107 Dipiperonal-acetone hydrogenated, 565 of formic acid, 773 Dipropionic nitrile, 231 in gas phase, 762-777 Dipropyl-amine, 427, 733 of glycerine, 760 Dipropyl-carbinol into amine, 735 limits, 21, 750-752, 767-770 Dipropyl ketone formed, 843 in liquid phase, 748-761 hydrogenated, Pt 567 mass law, 770 Dipropylene polym., 213 rates, 775 Divinyl polym., 213 theory of, 177, 752, 763 Dodecahydro-anthracene by hydrogenaby titania, 767 tion, 29, 363 velocity, 747, 774, 777 Dodecahydro-phenanthrene dehydrogen-Esters from aldehydes, 226-228 ated, 642, 646 with ammonia, 871 by hydrogenation, 484 as catalysts, 104 Doremol hydrogenated, Pt 570 condensed, 803 Doremone hydrogenated, Pt 570 decomposed, 180n, 858-874 Drying hydrogen for hydrogenation, formed, 75, 170, 175, 226-228 949 hydrogenated, 417 Drying oils, 266 hydrolysed, 83, 170, 313-316, 319, Dulcite, 588, 595 321, 337 Duodecene polym., 210 saponified, 175, 305, 337 Duratol, 967 Ethane, 423, 518, 526, 527, 546, 558, Durene hydrogenated, Pt 569 601, 605, 620, 631, 665, 709 from acetylene, 26, 914 Egg lecithine hydrogenated, Pd 555 decomp by heat, 911 ElaIdic acid formed, 82, 184 decomp. by Mg, 920 esters of, 937 from ethylene, 912 hydrogenated, 422 formed, 409 into ketone, 843 by hydrogenation, 26, 342, 377, 412, Electric heating, 349 413, 912, 914 Electrolytic dissociation, 175 Ether cond. with bensene, 817

decomp., 180m, 338 formed, 169, 180g, 690, 764, 872 in Grignard reagent, 6 oxidised, 254 process, 159, 691 Ethers, catalysts, 104 decomp., 180m, 321, 338, 494 formed, 169, 690, 764, 872 hydrogenated, 418 Ethoxy-cyclohexane, 464 Ethyl acetate from aldehyde, 228 catalyst, 304, 605 decomp., 180f, 180l, 180n, 858, 861, 861n. 871 formed, 228, 407, 749 hydrolysed, 313, 316, 819 neg. cat., 11, 303 Ethyl acetoacetate hydrogenated, 387 Ethyl-acetylene, 192 841 Ethyl alcohol, 680, 742 into acetal, 780, 783 into acetol, 783 into amines, 732 catalytic solvent, 38 decomposed, 650, 679 dehydrated, 688, 691, 694, 696, 700, 702, 709, 713, 716-719 864, 871 dehydrogenated, 538, 656, 667, CdO 674, MnO 672, Ni 664, Pt 668, SnO 673, Zn 670 esterified, 750, 770, 771, 778 hydrogenation agent, 538 oxidised, 150, 180b, 249, 254, 257, 260, 268 Ethyl adipate dec., 874 Ethyl-amine cat. prep., 732 con. agent, 804 dec. by Ni, 631 hydrochloride catalyst, 783 by hydrogenation, 377, 382, 386, 510 oxidised, 256, 268 Ethyl-aniline dec., 634 hydrogenated, 468 prep., 729 Ethyl-bensene, 451, 516, 520, 538, 539, 546, 548, 560, 641, 657, 728 decomp., 888, 930 681, 682 formed, 362, 369, 389, 415, 817, 888, 890 hydrogenated, 362, 448, 534 by hydrogenation, 362, 369, 389, 415

Ethyl bensoate, 744n, 749, 754, 755, 766 dec., 858, 864, 871 hydrolysed, 316, 319 Ethyl bromide formed, 104 Ethyl bromacetate reduced, 407 Ethyl-tert-butyl-bensene, 389 Ethyl-tert-butyl ether. 691 Ethyl butyrate dec., 858 by hydrogenation, 387 Ethyl caproate dec., 861, 862 Ethyl carbylamine hydrogenated, 430 Ethyl chloracetate red., 407 Ethyl chloride chlorinated, 282 by F. and C. reaction, 888 Ethyl cinnamate hydrogenated, 601 Ethyl evanide catalyst, 108, 605 hydrogenated, 427 Ethyl-cyclohexane dehydrogenated. Ni by hydrogenation, 362, 448, 451, 452, 455, 516, 520, 568, 569 Ethyl-cyclopropane, 193 Ethyl-diphenyl formed, 241 Ethylene, 423, 527, 548, 620, 626, 631, 634, 650, 670, 686, 689, 691, 696, 700, 708, 709, 713, 716, 726, 732, cond. with bensene, 241 cond. by sulphuric acid. 159 dec., 637, 912, 920, Co 912, Fe 912, Mg 920, Ni 413, Pt 912 formed, 78, 180g, 873, 914 hydrogenated, Co 500, Cu 515, Ni 413, 601, Pd 546, Pt 526, 558 manufacture, 180h, 689n, 717n oxidised, 180b polymerised, 211 preparation, 696n Ethylene bonds hydrogenated, 939 Ethylene compounds hydrated, 306 hydrogenated, 412-422, 587, Co 500, Cu 515, 594, Fe 506, Ni 601, Pd 546, 577, Pt 526, 558, 565 Ethylene chloride in F. and C. syn., 890 cyanide hydrogenated, 429 Ethylene hydrocarbons formed, 48, 86, in F. and C. syn., 90, 241 hydrogenated, Co 500, Cu 515, Fe 506 polymerized, 210 Ethylene oxides hydrogenated, 443

SUBJECT INDEX

isomerised, 200	by hydrogenation, 577
Ethylenic acids, 937	Ethyl valerate dec., 864
Ethylenic chlorides, 243	sapon., 316, 319
Ethyl ether, 694, 689, 713	Ethyl vanilline hydrogenated, 568
oat., 605 formed, 873	Eucalyptol dehydrogenated, 645
	Eucarvone hydrogenated, 552
hydrogenated, 494	Eudesmene hydrogenated, 570 Fuernal hydrogenated, N; 500, 602, Pd.
prepared, 691	Eugenol hydrogenated, Ni 590, 603, Pd
Ethyl formate, 773, 866	577, Pt 565, 569
dec., 866	isom., 191
hydrolysed, 316	Eugenol methyl ether hydrogenated, 590
Ethyl glutarate dec., 874	Primes 067m
Ethyl hexahydrobensoate, 471, 476	Fairco, 967n
Ethylidene chloride in F. and C. syn., 890	Farnesol hydrogenated, 570
Ethyl iodide cat., 299	Fats alcoholised, 341
in Grignard reaction, 302	hydrogenated, 542, 937–969
in syn., 605, 901	saponified, 314, 317
Ethyl-isoamyl-amine, 738	Fatty acids, effect on Ni, 948
Ethyl-isoamyl ether, 691	by hydrol., 314, 315, 318
Ethyl-isobutyl ether, 691	Fenchane, 611, 722
Ethyl isobutyrate, 316, 319	Fenchone, 611
Ethyl isovalerate, 417	Fenchyl alcohol dehydrated, 722
Ethyl malonate dec., 783	Ferments, soluble, 18
Ethyl mercaptan, 626, 744	Ferric chloride acetal cat., 781, 783
Ethyl-methyl-hexene hydrogenated, 414	catalyst, 687, 843, 849, 878
Ethyl-naphthalenes by hydrogenation,	chlorination cat., 285, 285n
390	cond. cat., 902
Ethyl naphthoates to nitriles, 871	in F. and C. syn., 899, 900
Ethyl nitrate cond., 819	Ferric oxide dehydrogenation cat., 677,
Ethyl nitrite as cat., 104, 207	686
hydrogenated, 382	ketone cat., 843, 849
Ethyl oleate hydrogenated, 565	mixed cat., 702
Ethyl ortho-formate, 783	Ferric sulphate cat., 725
Ethyl oxalate as cat., 104	Ferrous carbonate chlorination cat., 285
dec., 873	Ferrous chloride cat., 876, 954
Ethyl phenyl-acetate dec., 871	in F. and C. syn., 899
Ethyl-phenyl carbinol, 728	Ferrous oxide cat., 180j, 827
Ethyl-phenyl ether, 789	ketone cat., 843, 849
Ethyl propionate dec., 858	Ferulene hydrogenated, 570
Ethyl-propyl ether, 691	Fibrine cat., 110
Ethyl-pyridines syn., 901	Fish oils, effect on cat., 947
Ethyl-pyrrol, 742	hydrogenated, 939, 967n
Ethyl stearate dec., 858	Fittig syn., 11,
by hydrogenation, 565	Flake white, 967n
Ethyl succinate dec., 873, 874	Fluorides cats., 841
Ethyl sulphide, 626	Fluorene by dehydrogenation, 642
Ethyl terephthalate, 590	in F. and C. syn., 896
Ethyl tetrahydrobenzoate, 476	hydrogenated, 454
Ethyl toluate, 590	Formaldehyde, 73, 236, 562, 656, 664,
Ethyl-toluidines, 489	672, 674, 676, 678, 821, 825, 826,
Ethyl-trimethylene hydrogenated, 472	851, 870

SUBJECT INDEX

	7 4 000
catalyst, 269n	Furfurane, 620
dec., Cu 621, Fe ₂ O ₂ 677, Pd 623	Furfurane-dicarbonic acid, 727
into ester, 225, 228	Furfurane rings, 727
formed, 866, 871	Furfuroine formed, 220
hydrogenated, 432	Furfuryl alcohol, 371, 434
by oxidation, 249, 252–254, 256	Furfuryl-ethyl carbinol hydrogenated,
with phenols, 792	487
preparation, 249, 252-254	
into sugars, 221	Gaidic acid formed, 184
Formates, 851	Galactobiose, 18
Formic acid, 64, 621, 839, 851, 852, 855,	Galactose, 18, 186, 188
866	hydrogenated, Ni 588, Pd 595
decomp., 99, 143, 172, 180g, 624, 820-	by oxidation, 268
828	Galician petroleum, 927
esterified, 773	Galtose formed, 186
hydrogenating agent, 537, 539, 604	Gases condensed by metal powders, 135
by oxidation, 249	in porous bodies, 131, 132, 134
oxidised, 246	Gases from cracking, 909
syn. of, 574	Gasoline by cracking, 906, 932-936
toxic to Pt black, 117	Geometrical isomers, 182
Formic esters dec., 624, 866–870	Geraniol dehydrogenated, 658
Form of metals, 41, 53-55, 76-80	hydrogenated, Ni 416, 601, Pd 595,
Formyl chloride, 298	Pt 565
Fouling of catalysts, 118-120, 122, 932	Glass powder as cat., 811, 827, 828
Friedel and Crafts synthesis, 33, 87–89,	Gluconic acid, 187
157, 173, 174, 241, 241n, 297, 298,	Glucose hydrogenated, Ni 588, Pd 595
883 -9 00	by hydrolysis, 324–329
catalysts for, 899, 900	isomerized, 186
catalytic nature of, 898	multirotation of, 188
complications, 885	Glucosides dec., 18, 175
cyclic compounds, 896–898	hydrolyzed, 305, 327–330
with diphenyl, 889	syn., 15, 18, 793
with ethylene hydrocarb., 241	Glutaryl chloride in F. and C. syn., 893
isomerizations in, 888	Glyceric aldehyde cond., 237
of ketones, 891–894	formed, 236, 246, 268, 680
mechanism of, 898	by oxidation, 268
method of operating, 884, 892	Glycerides saturated, 939
with naphthalene, 889	Glycerine acetylated, 89
results of, 889	by alcoholysis, 340, 341
reversed, 887	dec. to formic acid, 855
Fructose, 186, 324	dehydrated, 725
Fumaric acid esterified, 756	dehydrogenated, 680
esters from maleic, 182	esterified, 757, 760, 761
hydrogenated, Pd 546	esters of, 340, 937
isom., 182	by hydrolysis, 314, 318
from maleic, 182	oxidised, 246, 249
Furfural cond., 686	Glycol dehydrated, 724
decom., Ni 620	oxidised, 249, 268
formed, 727	Glycolic acid esterified, 756
hydrogenated, 434	Glycolic aldehyde by oxid., 249, 268
oxidised, 268	Glyoxal by oxid., 249, 268

Gold, absorption of O₂ by, 137 catalyst, 66 colloid, 70, 72 dehydrogenation cat., 637 oxidation cat., 252, 254 Gold chloride, chlorination cat., 283 Goose fat, 938 Graphite catalyst, 702, 717, 911 formation, 180a Grease cat. poison, 1800 Greenwich gas works, 373 Grignard reaction, 44, 104, 300-302 Guaiacol hydrogenated, Ni 589 Gulose, 186 Gum arabic, 546, 561 Gunpowder dec., 8

Halides, 300 Halogenated alcohols dec., 876 Halogens eliminated by hydrogenation, 403-407, 545, 605 toxic to catalysts, 112-114 Hardened oils as foods, 967n, 969trade names of, 967 Hardening of fats, 577 of oils, 937-969 Heavy hydrocarbons cracked, 932 by cracking, 906, 936 Heavy oils by cracking, 906, 936 Helicine hydrolysed, 828 Helleborine, 330 Heptachlorpropane decom., 879, 357 formed, 242, 625, 903 Heptachlortoluene formed, 287 Hepta-isobutanal formed, 224 Heptaldehyde cond., 795 crotonised, 795 hydrogenated, 559 prepared, 853 Heptaldoxime hydrogenated, 383 Heptamethylene ring hydrogenated, 479 Heptane by cracking, 936 by hydrogenation, 519 Heptane-thiol, 745 Heptane hydrated, 306, 519 Heptine hydrogenated, 425, 519 Heptoic acid into aldehyde, 853 into ketone, 845 Heptoic aldehyde, 664 Heptyl alcohol dehydrogenated, 664 by hydrogenation, 559

Heterogeneous systems, 7, 34 Hexachlorbensene, 284 Hexachlorethane, 289 Hexachlorpropane, 242 Hexachlortoluene, 287 Hexadienal, 796, 801 Hexahydro-acetophenone, 476 Hexahydro-anisol, 589 Hexahydro-anthrone hydrogenated, 890 Hexahydro-bensoic acid, 471, 476, 551, 560, 569, 590 dehydrogenated, 649 esters of, 471, 649 Hexahydro-bensyl-amine, 470 Hexahydro-bensyl-aniline, 560 Hexahydro-carvacrol, 459 Hexahydro-cinchonine, 561 Hexahydro-cymene, 465, 478 Hexahydro-durene hydrogenated, 569 Hexahydro-guaiscol, 589 Hexahydro-indoline, 485 Hexahydro-naphthalid, 563 Hexahydro-phenanthrene, 484 dehydrogenated, 642 Hexahydro-phenylacetic acid, 476 Hexahydro-phthalic acid, 563, 590 Hexahydro-phthalid, 563 Hexahydro-phthalimide, 569 Hexahydro-terephthalic acid, 648 Hexahydro-toluene, 581 Hexahydro-toluic acids, 471, 563 Hexahydroxy-anthraguinone, 274 Hexahydroxy-bensene hydrogenated, 578 Hexa-isobutanal formed, 224 Hexamethyl-benzene decom., 887 formed, 212, 691 Hexamethylene hydrocarbons dehydrogenated, 649 hydrogenated, 475 Hexamethylene-tetramine cond., 792 hydrogenated, 496 Hexane from acetylene, 211 by cracking, 936 decom., 920 formed, 664, 665 by hydrogenation, 414 as solvent, 38 Hexaphenyl-cyclohexane, 880, 916 Hexene hydrated, 306 hydrogenated, 414, 515

Heptyl-amine by hydrogenation, 383

Waritan EOK	Hudanawania and hudanawanadad 949
Hexites, 595 Hexose from HCHO, 221	Hydrocyanic acid hydrogenated, 842, 528
Hexyl alcohol, 801	by hydrolysis, 329
Hexyl-bensene hydrogenated, 569	polym., 230
Hexyl cyanide, 814	stabilised, 11
High pressure in catalysis, 541	toxic to cats., 116
History of catalysis, 4	Hydrocyclic hydrocarb. dec., 921
Hofmann's reaction, 901	Hydrogen abs. by Co, 136, by Pd 165, by
Hog lard, 938	Pt 136
Homogeneous catalysis, 5, 144	elim. from hydrocarb., 905
Hydration, 305–312, 305–339	generator, 346
of acetylene comps., 308	for hydrogenation, 953
of ethylene comps., 306, 307	influence in dec. hydroc., 924
in gas phase, 337–339	from iron, 954
of imides, 311	manufacture, 953, 954
in liquid medium, 313–331	occluded by Co, 136, by Pd 165, by
mechanism of, 308	Pt 136, 166
of nitriles, 311	purification of, 346
Hydrasine compounds decom., 611	rate of production from Fe, 954
Hydrasobensene, 202, 531, 554	from water gas, 953
Hydraso compounds hydrogenated, 600	Hydrogenation, 15, 65, 111, 115, 121,
Hydrazones decom., 611	138, 165, 342-407, 408-497, 498-
Hydrindene, hydrogenated, 454	540, 541-583, 563n, 584-604, 721,
Hydrindone cond., 799	923, 931, 932, 937-969
Hydro-aromatic hydrocarbons, 424, 444	of acetylene, 501, 506
dehydrogenated, 640-649	of acetylene comps., 423 et seq., 518,
Hydrobensamide, 194	<i>5</i> 27, <i>5</i> 86, <i>5</i> 77
Hydrobromic acid elim., 901	of acid chlorides, 575
Hydrocarbons from acetylene, 925	of acids, 422, 471
from acids, 829–836, 839	of acridine, 491
from alcohol + aldehyde, 784	by alcohol vapors, 537, 538
condensed, 241, 905–936	of alcohols, 369, 416, 465
decom., 87, 493, 905–936	of aldehydes, 388, 419, 432, 433, 503,
decom. in H ₂ , 924	522, 532, 567, 588, 602
dehydrogenated, 639–649	of aliphatic aldehydes, 432, 532
hydrogenated, 413, 444-454, 472 et	of aliphatic amides, 386
seq. 481-485, 493, 500-502, 506,	of aliphatic ketones, 532
515-518, 526, 527, 534, 565, 566,	of aliphatic nitriles, 427
569, 570, 577, 601	of aliphatic nitro comps., 377
formed, 695–727, 784–815, 829–836,	of alkaloids, 555
839, 925	of amides, 386
oxidised, 254, 259	of amines, 466, 496
polym.,84	of anhydrides, 392
Hydrocarvols, 476	of anthracene nucleus, 483
Hydrochloric acid cat. acetale, 782	apparatus, 345–357, 543, 584–585, 597
cond. agt., 730, 782, 792, 799, 803-805	et seq., 957-964
dehydration cat., 687, 795	of aromatic acids, 471
in esterif., 748–750, 754–757	of aromat, ales., 369, 465
in hydration, 307	of aromat, alds., 388, 438
toxic to cats., 116 Hydrocina mic estern hydrocenated 471	of aromat. amines, 466 of aromat. diketones, 391
Hydrocinnamic esters hydrogenated, 471	or aromas, uncooling, 071

of aromat. halogen comps., 403 of aromat, hydrocarb, 466 et seg., 502 of aromat, ketones, 389, 455 of aromat, nitriles, 428 of aromat, nitro comps., 378 of aromat, nucleus, 444 et seq., 534, 569, 589 of benzene and homologs, 466 et seq., of carbasol, 490 of carbon 525 of carbonates to formates, 574 of carbon dioxide, 395, 504, 508 of carbon disulphide, 372, 492 of carbon monoxide, 393 by carbon monoxide and hydrogen. 537 by cobalt, 499-504 by colloidal Pd. 545-555 by colloidal Pt, 556 of complex rings, 571 by copper, 507, 523, 594 of cyclic comps., 578 with dehydration, 721, 722 of diago comps., 497 of dicvanides, 429 of diketones, 438 of esters, 417 of ethers, 418, 494 of ethyl acetoacetate, 387 of ethylene comps., 500, 506, 515, 526, 565, 577, 601 of ethylene hydrocarb., 500, 506, 515 of ethylene oxides, 443 by formic acid, 537, 539 of furfuryl alcohol, 371 furnace for, 347, 348 in gas system, 366-407 of halogen comps., 403 et seq. of heptamethylene ring, 479 of hexamethylene ring, 475 history of, 342-344, 542, 939 of hydrocarbons, 413, 493, 499 et seq. hydrocyanic acid, 528 of indol, 497 by iron, 505, 506, 593 of isocyanides, 431 of keto-acids, 437 of ketones, 389 et seq., 420, 435 et seq. 441, 455, 503, 522, 532, 567, 588, 602

of liquid fats, 937-969 in liquids, 350-352, 541 et seq., 584 et seq., 596-603 in manuf. of ill. gas, 397 et sea. methods, 343 et seq., 544, 562, 573, 584, 596, 597, 599, 604 of naphthalene nucleus, 481, 931 by nascent hydrogen, 537 of nitriles, 427, 428, 521 of nitro comps., 377, 378, 509, 529, 564, 576, 600 of nitrous esters, 513 of nitrous oxide, 368 of octomethylene ring, 480 of oxides of carbon, 504 by oxides of metals, 598, 943 of oxides of N, 374 of oximes, 283, 514 by palladium, 536, 544-555, 573-578 by palladium black, 573-578 of pentamethylene ring, 474 of phenanthrene nucleus, 484 of phenol ethers, 464, 494 of phenols, 370, 456 of phenyl isocyanate, 495 by platinum, 524-535, 556-571 by platinum black, 562-572 of polycyclic hydrocarbons, 452 of polymethylene rings, 535 of polyphenols, 370, 460 products, 355, 356, 965 of pyridine, 486 of pyromucic ald., 434 of pyrrol, 486 of quinoline, 488 of quinones, 442 removes odors, 939 results of, 355, 356, 965 of solids, 353 temperatures for, 599 of terpenes, 477, 570, 591 of tetramethylene ring, 473 theory of, 167, 365 of trimethylene ring, 472 by various metals, 580, 595 of various rings, 472 et seq., 571, 592, Hydrogen halides elim., 875-903 Hydrogen ions in hydrol, 82, 313, 324 Hydrogen peroxide decom., 2, 32, 38, 83,

160, 180a

mith charmin and 147	Timomata add 104
with chromic acid, 147	Hypogaelic acid, 184
oxidising agt., 268	esters of, 937
with permanganate, 148	Ultiminating and her herdengenation 207
stabilized, 11, 13	Illuminating gas by hydrogenation, 397–
Hydrogen persulphides, 83	402
Hydrogen selenide decom., 8	freed from CS ₁ , 372
Hydrogen sulphide, 686, 743, 791, 810,	manufacture, 397–402
924, 947	purification, 339, 372
with alcohols, 743	Imbibition of liquids by porous sub., 133
with aldehydes, 810	Imides, 305, 312
elim., 626–629	Indene polym., 217
isom. agt., 182	Indose, 186 Indian hydrogeneted 165, 602
toxic to cats., 1800, 598, 947	Indigo hydrogenated, 165, 603 Indigotine hydrogenated, 603
Hydroiodic acid cat., 82, 183 decom. limit, 15, 20	
	Indigo white, 165, 603 Indol, 684
formation, 342 isom. cat., 182	hydrogenated, 497, 571
Hydrolecithin, 555 Hydrolysis, 82, 175–178, 305, 313–336	Indols cond., 803
of acetals, 322	formed, 89, 91, 633 Induced catalysis, 149
by acids, 313	oxidations, 244
of amides, 331	Influence of solvents, 38–40
by bases, 318	Infra-red radiation as cat., 180j
of carbon disulphide, 339	Infusorial earth as carrier, 126, 587n,
of esters, 313–319, 337	598, 941
of ethers, 321, 338	Inosite, 578
in gas system, 337–339	Intermediate comps. in cat., 151-158,
of glucosides, 327	163–173, 179, 180, 752, 763, 859–
of halogen comps., 320	864, 866, 872, 878, 898, 916
of polysaccharides, 323	in esterif., 752, 763
Hydropivalic acid esterif., 227	in F. and C. syn., 898
Hydroquinine, 604	in oxidation, 258, 541, 677
Hydroquinone by hydrogenation, 442	Inversion of reactions, 14
reduced, 370, 461, 589	of sugar, 32, 324
Hydroxy-acids esterif., 756	Iodides cats., 84
Hydroxy-anthraquinones by oxidation,	Iodination, 294
274	Iodine absorbed, 938
Hydroxy-benzoic acid hydrogenated,	bromination cat., 291
569	catalyst, 6, 33, 43, 156, 278, 299, 632
Hydroxy-butyric aldehyde formed, 307	chlorination cat., 156, 278n, 287
Hydroxy-cyclohexanes dehydrated, 642	Iodine
Hydroxy-esters dehydrogenated, 663	dehydration cat., 699, 729, 790
Hydroxy-isoheptoic acid, 663	elim., 406, 605
Hydroxylamine by hydrol., 332	isom. cat., 182
Hydroxyl group elim., 465	sulphonation cat., 296, 815, 815n
introduced, 269	toxic to cats., 116, 359
Hydroxy-methylene comps. hydrogen-	Iodine numbers, 938, 955
ated, 550	of hardened oils, 967, 967,
Hydroxy-stearic acid formed, 306	in hydrogenation, 966
toxic to cats., 115	Iodine trichloride catalyst, 44, 85, 156
Hypochlorites as oxidising agents, 270	chlorination cat., 85, 278

Iodobensene reduced, 406	Iron retorts in cracking, 934
in syn., 904	Iron salts oxid. cat., 268, 271, 277, 320
Ionones formed, 198	reduced, Pd 165
hydrogenated, 554, 560	
Ions in hydrolysis, 305	Iron scale cat., 285 Iron sesquioxide chlorination cat., 285
Iridium black, 582	Iron sulphate chlorination cat., 285
catalyst, 64	oxidation est., 272, 275
colloidal oxid. cat., 251	Iron sulphide chlorination cat., 284
Iron, bromination cat., 298	Isoamyl acetate dec., 871
catalyst, 167, 180r-180u, 320, 344,	Isoamyl alcohol into amines, 733, 740,
505, 506, 540, 683	741
catalyst, prep. of, 58	dehydrated, 691, 696, 713, 715, 717,
chlorination cat., 278n, 285, 285n	719
cracking cat., 906, 910, 911, 932	dehydrogenated, 656, 664, 672
dec. C ₂ H ₃ , 913, 915, 920, 928	esterified, 771, 773
dec. alcohols, 667	oxidised, 254, 268
dec. aromat. hydrocarb., 921	Isoamyl amine from alcohol, 733
dec. CO, 615	catalyst, 836
dec. ethylene, 912	dehydrogenated, 681
dec. pinene, 922	by hydrogenation, 382
dehydrogenation cat., 637, 651, 652,	Isoamyl bensoate, 766
667	Isoamyl-carbinol hydrogenated, 570
in drying oils, 266	Isoamyl cyanide hydrogenated, 427
harmful in hydrogenation, 115	Isoamyl ether, 691
hydrogenation cat., 344, 505, 506, 593,	Isoamyl formate, 773
945	Isoamyl hexahydrobenzoate, 471
in hydrolysis of bensalchloride, 320	Isoamyl malonate dec., 873
influence on Pd, 946	Isoamyl mercaptan, 626, 744, 746
method for prep. of hydrogen, 953,	Isoamyl nitrite hydrogenated, 382
954	Isoamyl oxalate dec., 873
Iron benzoate, 320	Isoamyl-phenyl ether, 789
Iron borate cat., 265	Isoamyl-piperidine, 741
Iron bromide, bromination cat., 240, 293	Isoamyl succinate dec., 873
Iron chloride, bromination cat., 293	Isoamyl sulphide, 626
chlorination cat., 283	Isobutane, 472
cracking cat., 936	Isobutyl acetate dec., 861, 862
in F. and C. syn., 88	Isobutyl alcohol into acetal, 780
halogenation cat., 88	from aldehyde, 226
hydration cat., 310	dehydrated, 691, 696, 700, 713, 715-
polym. cat., 216	717
prep. acetals, 88	dehydrogenated, 656, 670
Iron compounds cats., 269n	esterified, 771, 776
Iron hydroxide oxid. cat., 150	oxidised, 249, 254, 268
Iron oxides cat., 6, 75, 100, 260, 285, 310, 320	Isobutyl-amine by hydrogenation, 382 Isobutyl benzoate, 766
dehydration cat., 702	Isobutyl bromide isom., 200
hydration cat., 310	Isobutyl chloride dec., 878, 881
ketone cat., 843, 849	in F. and C. syn., 900
oxidation cat., 257–259	Isobutyl cyanide, 681, 682
prep., 77	Isobutylene formed, 142, 713, 878
Iron powder cat., 320	hydrated, 306
	-J 100) 000

Technity other 601	from magas 206s
Isobutyl ether, 691 Isobutyl hexahydrobensoate, 471	from gases, 306n by hydrogenation, 391
	oxidised, 254n
Isobutyl-isoamyl-amine, 738 Isobutyl isobutyrate from ald., 226	preparation, 435
Isobutyl malonate dec., 873	Isopropyl amine from alc., 735
Isobutyl mercaptan, 744	by hydrogenation, 382
Isobutyl nitrite hydrogenated, 382	Isopropyl-bensene hydrogenated, 448
Isobutyl oxalate dec., 873	Isopropyl bensoate dec., 871
Isobutyl succinate dec., 873	formed, 766
Isobutyric acid from ald., 226	Isopropyl bromide by isom., 93, 199
dec., 839	Isopropyl chloride by isom., 199
esterif., 770, 771, 775, 776	Isopropyl-cyclohexane, 449, 452
into ketone, 840, 842–845	Isopropyl-cyclohexyl-amine, 739
Isobutyric aldehyde from alc., 670	Isopropyl-cyclopentanone, 546
cond., 808	Isopropyl-ethylene, 713
crotonized, 795	Isopropyl-guaiacol, 565
into ester, 226	Isopropyl iodide, 605
hydrogenated, 432, 588, 593	Isopropyl nitrite hydrog., 382
by oxidation, 249	Isopropylidene-cyclopentanone hydrog.,
phenylhydrazone dec., 635	546
polymerized, 224	Isosafrol hydrogenated, 418, 565, 590.
Isobutyryl chloride, 813	601
Isocamphane, 591, 722	Isosulphocyanic esters hydrol., 334
Isocrotonic acid hydrogenated, Pd 546	Isothujone formed, 198
Isocyanates from diazo, 610	hydrogenated, 552
hydrogenated, 431	Isovaleraldoxime, 814
Isocyanides hydrogenated, 431	Isovaleric acid into ald., 853
Isocyanic esters hydrol., 334	dec., 839
Isodulcite by hydrol., 328	esterif., 771
Isoeugenol formed, 191	into ketone, 842–844
hydrogenated, Ni 590	Isovaleric aldehyde cond., 808
oxidised, 249	formed, 664
Isoheptoic aldehyde, 635	hydrogenated, 432, 588
Isomerizations, 181-208	by oxidation, 268
of alkyl halides, 876	phenylhydrazone dec., 635
in F. and C. syn., 888	Isovaleric anhydride into ketone, 857
Iso-oleic soid hydrated, 306	Isovaleric esters dec., 871
Iso-oximes formed, 205	Isovalerone, 420
Isopentane, 681	Isovaleronitrile, 814
by hydrogenation, 414, 420, 472	Isovaleryl chloride, 813
Isopentene isom., 190	Isozingiberene hydrogenated, 570
Isoprene formed, 235, 723, 802, 909	Itaconic acid formed, 183
polym., 50, 106, 213, 214	hydrogenated, 558
Isopropyl-acetylene, 192	isom., 183
Isopropyl alcohol, 439, 503, 567, 588,	·
593, 594, 784	Jena glass cat., 827
into amine, 735	
dec. by C, 679	Kaolin carrier for Ni, 941
dehydrated, 700, 716, 719	dehydration cat., 99, 717, 723, 726,
dehydrogenated, 659, 665, 668	802
esterif., 757, 766, 775	oxidation cat., 267

Kayser's apparatus, 963 Ketimines formed, 809 Keto-acids esterif., 756 hydrogenated, 437 syn. of, 902 Keto-alcohols dehydrogenated, 663 Keto-esters, 663 Keto-hydrofurfurance formed, 195 Keto-isoheptoic esters, 663 Ketones from alcohols, 650, 659 into alcohols, 549 alicyclic hydrogenated, 436 aliphatic hydrogenated, 435 aromatic hydrogenated, 441, 455 condensed, 81, 238, 794-801, 803-810 condensed in gas phase, 801 crotonized, 794-800 crotonized in gas phase, 801 decom., Ni 620, Pt 532 dehydrated, 794-800, 802 from esters, 860, 861 formed, 31, 75, 208, 332, 701, 723, 764, 829, 830, 837-851, 857, 858, 865, 891-894 formed in liq. phase, 847 by hydration, 305, 308 hydrogenated, 420, 435, 436, 441, 455, Co 503, Cu 522, Fe 506, 593, Ni 588, 602, Pd 549, Pt 532, 567, 568 from oximes, 332 polym., 229 from second, alcs., 659 syn. by F. and C., 891-894 Ketoximes dehydrated, 814 hydrogenated, 383, 385, 514 Kieselguhr, carrier, 942 Kream Krisp, 967n

Lactones by hydrogenation, 392
Lactone, 323
Laevulinic acid esterif., 756
hydrogenated, 437
Laevulose formed, 221, 236, 237
hydrogenated, Ni 588, Pd 595
multirotation, 188
Lampblack cat., 811
Lard, 928
Lard oil hardened, 966
Lauric acid into ketone, 843, 850
Lead in drying oils, 266
influence on, Pd 946

toxic to cats., 115 Lead chamber process, 32, 158 Lead chloride cat., 876, not cat. 283 Lead hydroxide, isom, cat., 186 Lead nitrate, oxidation cat., 277 reduced with Pt. 166 Lead oxide cat., 676 Lead soaps toxic to cats., 115 Life of catalysts, 111, 708, 947 Ligroine as solvent in F. and C. syn., 892, 897 Lime catalyst, 540, 827 decom, methane, 911 dehydration cat., 795, 797 ketone cat., 840, 849 Limits of esterification, 750, 751, 767change with temperature, 768-770 Limits of reactions, 22, 313 Limonene dehydrogenated, 644 formed, 198 hydrogenated, Cu 517, Ni 477, 591, Pt 570 Linalool hydrogenated, Ni 416, 601, Pt 565 Linoleic acid constituent of fats, 937 hydrogen req. for sat., 955 hydrogenated, Pd 558 Linoleic esters, 937 Linolenic esters, 937 Linseed oil alcoholised, 341 hardened, 966 iodine number, 938 Liquid fats hydrogenated, 937-969 Lithium carbonate ketone cat., 846 Lyxonic acid, 187

Magnesia carrier, 127
catalyst, 540, 702, 828, 901, 906, 920
Magnesium carrier, Pd 946
cat., 51, 901
in cracking, 906
dec. C₂H₃, 920
powder cat., 901
Magnesium compounds cats., 269n
Magnesium sulphate dehydration cat., 101
Maleic acid cat., 196
hydrogenated, 558
isomer., 182

SUBJECT INDEX

by oxidation, 260n, 276	Menthene by dehydration, 714
oxidised, 268	dehydrogenated, 644
toxic to cats., 115	hydrogenated, 475
Malic acid esterif., 756	Menthol, 436, 567
Malichite green hydrogenated, 603	Menthone hydrogenated, Pt 56
Malonic acid cond., 804	by hydrogenation, 552, 591
Malonic anhydride, 873	isom., 189
Malonic ester cond. alds., 804	Menthone-oxime hydrogenated,
decom., 873	Mercaptans formed, 75, 170,
Malonyl chloride in F. and C. syn., 893	707n, 743–746
Maltose hydrol., 323, 325	Mercaptans, secondary, 628
	Mercaptides, 627
Manganese bromination cat., 52, 292	Mercuric bromide bromination
in drying oils, 266	
oxidation cat., 52, 254	hydration cat., 309
Manganese chloride cat., 283	Mercuric chloride with Al on
Manganese dioxide cat. H ₂ O ₂ , 75	299
Manganese oxides oxidation cat., 259	with Al in F. and C. syn., 886
Manganese salts cats., 100, 153, 264,	bromination cat., 293
269n	hydration cat., 92, 309
Manganous acetate, 268	isom. cat., 92
Manganous borate, 265	toxic to cats., 116
Manganous oxide cat., 259, 617, 702,	Mercuric nitrate nitration cat.,
828, 840, 845, 850, 853, 866	oxidation cat., 269
on alcohols, 142	Mercuric salts red. with Pd, 16
dehydrogenation cat., 651, 672	Mercuric sulphate hydration of
ketone cat., 840, 845, 850	309
Manganous salts oxidation cats., 100,	oxidation cat., 272–274
153, 264, 268	sulphonation cat., 6., 102, 810
Mannite esterif., 757, 761	Mercury dec. H ₂ O ₂ , 180a
by hydrogenation, 588, 595	Mercury oxide oxidation cat., 2
oxidised, 150	Mesaconic acid formed, 183
Mannite hexacetate, 761	Mesitylene with CO, 298
Mannonic acid formed, 187	hydrogenated, 447
Mannose isom., 186	by isom., 888
by oxidation, 150	by polymer., 212
Margaric esters, 937	Mesityl oxide, 697, 699, 797, 80
Mechanical shaking, 562	hydrogenated, 420, 549, Ni
Mechanism of amine formation, 731	546, 595, Pt 559, 567
of hydrogenation, 677	Meso-benso-dianthrone deh
of Grignard reaction, 300, 301	ated, 685
of hydration, 308	Meso-naphtho-dianthrone, 685
of mercaptan decomp., 627	Meta-aldehydes depolym., 234
	formed, 222
of oxidation, 258, 264, 276	Meta-butanal formed, 223
of poisoning, 180 <i>p</i> –180 <i>s</i>	
of promoters, 180s–180u	Meta-chloral formed, 224
Melissic acid into ketone, 843	Meta-heptaldehyde formed, 223
Melting points of hardened oils, 966,	Meta-isobutanal formed, 224
967n	Metal chlorides as cats., 876
Menthane, 369, 449, 465, 475, 477, 478,	Metal oxides as cats., 169, 675,
518, 570, 591, 722	dehydration cats., 686
Menthane-diol, 463	Metals, compounds formed, 299

ated, 644 ed. 475 567 drogenated, Pt 567 nation, 552, 591 ime hydrogenated, 385 formed, 75, 170, 626-628, 3-746 secondary, 628 627 mide bromination cat., 293 at., 309 oride with Al on alcohols, F. and C. syn., 886 n cat., 293 at., 92, 309 92 ts., 116 ate nitration cat., 269n at., 269 s red. with Pd, 165 phate hydration cat., 102, at., 272-274 on cat., 6., 102, 816 H₂O₂, 180a e oxidation cat., 269n id formed, 183 ith CO, 298 ed, 447 88 ., 212 , 697, 699, 797, 801 ed, 420, 549, Ni 587, Pd Pt 559, 567 lianthrone dehydrogeno-dianthrone, 685 les depolym., 234 formed, 223 formed, 224 lehyde formed, 223 nal formed, 224 es as cats., 876 as cats., 169, 675, 881 n cats., 686 ounds formed, 299, 300

cond. of gases on, 135 by hydrogenation, 377, 382, 510, 530 in cracking, 906, 932 oxidised, 256 decom. acetylene hydrocarb., 913-919 Methyl-amyl-acetylene, 308 Methyl-aniline dec., 634 dec. aromat. hydrocarb., 921 dec. esters.. 867 formed, 729, 740 dec. formic acid. 823 hydrogenated, 468 dec. formic esters, 867 Methyl-anthracene from oracking, 909 dehydration cats., 686, 687, 701 Methyl-anthraquinone nitrated, 269% ketone cats., 830. 847 Methyl bensoate alcoholized, 340n Meta-propional formed, 223 dec., 871 by esterif., 744n, 766 Meta-styrene, 657 Methane, 432, 495, 504, 536, 540, 593, hydrogenated, 471 620, 631, 634, 641, 645, 664, 672 into nitrile, 871 Methyl-butadiene, 802 from CO. by hydrogenation, 395-402 from CO, 393 Methyl-butane-diol dehydrated, 723 decom., Mg 920. Ni 911 Methyl-tert-butyl-amine, 430 Methyl-butyl ketone hydrogenated, 435 equilib. in formation, 409-411 Methyl-butyl-phenol, 459 formed, 362, 369, 370, 377, 393, 395-402, 409-411, 413, 525 Methyl-carbyl-amine hydrogenated, 430 Methyl-carvacryl ether, 789 formed from carbon, 586 Methyl-chlorcyclohexane, 569 by hydrogenation, 362, 369, 370, 377, 395-402 Methyl chloride in F. and C. syn., 884 oxidised, 253 Methyl cinnamate hydrogenated, 601 Methods of hydrogenation, 599 et sea. Methyl-p.cresyl ketone hydrogenated. Methoxy-cyclohexane, 464, 494 Methoxy-methyl-cyclohexanols, 464 Methyl-cyclohexane, 197, 447-450, 452, Methoxy-propylbensene, 590, 601 465, 467, 479, 560, 590, 641 Methoxy-propyl-cyclohexane, 590 dehydrogenated, 641 Methoxy-propyl-phenol, 590 by hydrogenation, 388 by isom., 197 Methyl acetate dec., 1807 hydrol., 313 Methyl-cyclohexyl-amine, 739 Methyl-acetyl-acetone hydrogenated, 439 Methyl-cyclohexanols into amines, 739 dehydrogenated, 660 Methylal cond. with phenols, 792 formed, 781 by hydrogenation, 457 by oxidation, 249 Methyl-cyclohexanones by dehydrogena-Methyl alcohol, 432, 538, 740, 771, 773, tion, 660 851 hydrogenated, 436, 567 Methyl-cyclohexanone-hydrasones. 611 into acetal, 781 dehydrated, 688, 690, 691, 693, 713, Methyl-cyclohexenes, 515, 660 Methyl-cyclohexyl-amine, 467, 737, 739 715, 716 dehydrogenated, 656, 676, CdO 674, Methyl-cyclohexyl-aniline, 467 Methyl-cyclopentane by hydrogenation, MnO 672, Ni 664, SnO 673, Zn 678 detection in EtOH, 656 390, 649 esterif., 771, 773 Methyl cyclopentane-carbonate, 649 from formic acid, 826 Methyl-cyclopentanone hydrogenated, oxidised, 249, 268 390, 436 with phenol, 789 Methyl-cyclopropene hydrogenated, 472 with Pt. 668 Methyl-diphenyl carbinol, 721 Methyl-allene, 784 Methylene chloride in F. and C. syn., Methyl-amine from HCN by hydrogenation 342, 528 Methylene-dithiol, 492

Methyl esters by alcoholysis, 341 dec., 860, 865, 871 Methyl ether, 688, 690, 691, 693, 718, 865, 871 Methyl-ethyl-acetylene, 192 Methyl-ethyl-acroleine hydrogenated. 595 Methyl-ethyl-amine, 430 Methyl-ethyl-bensene by hydrogenation, 389 from pinene, 922 Methyl-ethyl-butadiene formed, 192 Methyl-ethyl carbinol, 567 Methyl-ethyl-cyclohexane, 448, 449 Methyl-ethyl-cyclohexene hydrogenated, 475 Methyl-ethyl ether, 691 Methyl-ethyl-ethylene formed, 193 Methyl-ethyl ketone hydrogenated, 567 Methyl-ethyl ketone phenylhydrasone dec., 633 Methyl-ethyl-propenal hydrogenated 546, 559 Methyl-di-isopropyl-benzene, 930 Methyl formate from ald., 228 dec., 868 by esterif., 773 Methyl-furfurane hydrogenated, 487 by hydrogenation, 371 Methyl-heptanone by hydrogenation, 420 Methyl-heptenone hydrogenated, 420, Methyl hexahydrobensoate, 471 Methyl hexahydroterephthalate, 648 Methyl-hexanone, 420 Methyl-hexenone hydrogenated, 420 Methyl-hexyl carbinol dehydrogenated, 665 Methyl-hexyl ketone, 665 Methyl-indol, 489, 633, 684 Methyl-isobutyl-bensene by F. and C. syn., 900 Methyl-isobutyl carbinol, 549, 559, 568 Methyl-isobutyl ketone, 435, 545, 559, 567, 587, 595 Methyl-isopropyl-cyclohexane, 449, 475 Methyl-isopropyl ketone dehydrated, 802 hydrogenated, 435 Methyl mercaptan, 744

Methyl-naphthyl ketone hydrogenated, Methyl nitrite hydrogenated, 382, 513 Methyl-nonyl ketone hydrogenated, 435 Methyl-pentanol, 559, 595 Methyl-pentanone, 420 Methyl-pentamethylene, 444 Methyl-pentene hydrated, 306 Methyl-pentyl alcohol, 546 Methyl-phenyl-butine-ol hydrogenated, 548 Methyl-phenyl carbinol, 728 Methyl-propyl carbinol, 487 Methyl-propyl-octane by hydrogenation, 166 Methyl-propyl-octene hydrogenated, 414 Methyl-propyl ketone hydrogenated, 435 by hydrogenation, 487 Methyl-quinoline, 488 Methyl-salicylic ald. hydrogenated, 568 Methyl tetrahydroterephthalate dehydrogenated, 648 Methyl-toluidines, 684, 740 Methyl-valeric ald., 546 Methyl-vanilline hydrogenated, 568 Mexican petroleum cracked, 933 MFB, 967n Migration of atoms, 199 Migrations of double and triple bonds, 190 Mineral acids as cats.. 81 Mixed amines, 738 Mixed catalysts, 538, 651, 675, 702, 826, 827, 866 Mixed ethers formed, 170, 789 Mixed ketones, 75, 847-850 Mixed oxide-catalysts, 538 Mixed phenol-ethers, 788, 789 Moisture in oils, 949 Molybdenum chloride chlorination cat., 90, 283, 286 Molybdenum compounds cats., 269n Molybdenum oxide cat., 675, 676, 693, 702, 827 Molybdenum oxide, blue cat., 675, 746, 791 Molybdenum promoter, 180s, 180u Monobasic acids dec., 829-854 Morphine hydrogenated, 572 oxidised, 268

Mucic acid dehydrated, 727
esterif., 756
formed, 187
Multirotation of sugars, 188
Mustard oils hydrol., 333
Mutton tallow, 938
Myrcene formed, 214
Myristic acid into ketone, 850

Naphthalene cond., 806 from cracking, 908, 909 dec., 921, 931 dec., by AlCla, 931 by dehydrogenation, 642 formed, 908 in F. and C. syn., 889, 899 hydrogenated, 481, Ni 592, Pd 553, Pt 571 by hydrogenation, 379 oxidised, 273 Naphthalene hydrides from C₂H₂, 914 dehydrogenated, 642 Naphthalic acids hydrogenated, 594 Naphthalic anhydride hydrogenated, 563 Naphthane, 481 Naphthenes formed, 211 Naphthoic acid esterif., 756 Naphthols into amines, 790 hydrogenated, 481, 592 Naphthol ethers hydrogenated, 494 Naphthonitriles formed, 871 Naphthoyl chlorides in F. and C. syn., Naphthyl-amines, 512, 630, 632, 729 hydrogenated, 496 by hydrogenation, 379 Naphthyl ethers, 789 Naphthyl ketones by F. and C. syn., 899 Natural gas, 928 Negative catalysts, 9, 11 Neutral salt effect in hydrol., 317, 319 in inver. of sugar, 324 Nickel, α , β , and γ forms, 360 amount of required, 951 carrier for, Pd 946 on carrier, 126, 598, 939, 941, 942, 959, 960 cat., 15, 24, 53, 111-115, 122, 167, 180l-180n, 343, 344, 358, 539, 540,

563n, 584 et seq., 596-603, 614, 619,

620, 683, 721, 722 cat. preparation, 54-56, 598, 941 in cracking, 906, 910, 911 dec. C₂H₂, 913, 918-920, 925, 926 dec. alcohols, 180g, 664 dec. aldehydes, 619 dec. amines, 631, 634 dec. aromatic hydrocarb., 921 decomp. cat., 832, 834, 867, 910-913, 918-921, 923, 925, 926 dec. CO, 163 dec. chlorides, 882 dec. esters, 1805, 180f, 180j dec. formic esters, 867 dec. hydrocarb., 832, 834, 867, 910-913, 918-921, 923, 925, 926 dec. ketones, 620 dec. pinene, 923 dehydrogenation cat., 636, 637, 640-645, 647, 651, 664, 665, 681, 684, 701, 824 elim. NH₄, 631 in hardened oils, 969 hydrogen comps., 167 hydrogenation cat., 197, 801, 932, 939, 941-945, 947, 948, 950, 951, 969 hydrogenation cat. for fats, 939, 941-948, 950, 951 isom. cat., 208 from Ni(CO)₄, 163, 598, 616, 942, 953 preparation of, 53-56, 598, 941 on pumice, 126, 939, 941, 942 temp. for use, 952 Nickel acetate, 944 Nickel borate cat., 265, 944 Nickel carbonate, 941 Nickel carbonyl, 163, 598, 616, 942, 953 Nickel chloride cat., 283, 876, 880, 947 Nickeled asbestos, 959, 960 Nickeled pumice, 126, 939, 941, 942 Nickel formate, 944 Nickel lactate, 944 Nickel nitride, 375 Nickel oxide cat., 75, 80, 254, 258, 259, 722, 943 hydration cat., 310 hydrogenation cat., 584, 598 se. nickel, 584 theory, 258 Nickel peroxide, 180a Nickel sesquioxide, 589

SUBJECT INDEX

Nickel suboxide, 80, 598, 943 Nickel sulphate oxid. cat., 272 Nitranilines hydrogenated, Cu 513 Nitration catalysed, 269n Nitric acid from NHs, 150, 249 in hydration, 307 hydrogenated, 376 on metals, 8 oxidising agent, 269 Nitric oxide hydrogenated, 374, Cu 509, Pt 529 Nitriles, 305, 633, 635, 681, 682, 808 formed, 15, 631, 811, 812, 814 hydrated, 311 hydrogenated, 426-429, 521 polymerized, 230 Nitriles, aliphatic hydrogenated, 427 Nitriles, aromatic hydrogenated, 428 Nitroacetophenone hydrogenated, 545, Nitro-alcohols formed, 236 Nitrobensaldehyde cond., 798 by oxidation, 270 Nitrobensene formed, 819 hydrogenated, 378, 538, 545, Cu 511, Pd 536, 576, Pt 531, 537 oxidising agent, 277 solvent for F. and C. syn., 892 Nitrobenzophenone F. and C. syn., 893 Nitrobensoyl chlorides in F. and C. syn., 893 Nitro compounds cond., 803 from diaso, 609 hydrogenated, 377, 378, Cu 509, Fe 506, Ni 600, Pd 545, 576, Pt 529, 557, 564 Nitro compounds, aliphatic hydrogenated, 377 Nitro compounds, aromatic hydrogenated, 378 Nitro-ethane cond., 236 hydrogenated, 377, 510 Nitrogen eliminated, 606-612 Nitrogen dioxide hydrogenated, 529 Nitrogen oxides hydrogenated, 529 Nitrogen peroxide hydrogenated, 375, Nitromethane cond., 236, 803 hydrogenated, 377, Cu 510, Pd 536, Pt 580 Nitro-methanol-butanol, 236

Nitro-methylol-propane-diol, 236 Nitronaphthalene hydrogenated. Nitroparaffines cond., 236 Nitrophenols hydrogenated, 381, Cu 512, Pd 536 by oxidation, 269 Nitrophenyl-ethylene, 803 Nitropropane cond., 236 Nitropropanol, 236 Nitropropyl alcohol, 236 Nitrosamines, 108 Nitroso compounds as cats., 108 hydrogenated, 564 Nitroso-dimethyl-aniline in vulc., 104 Nitroso-naphthol hydrogenated, 564 Nitroso-phenol as cat., 108 Nitroso-terpenes hydrogenated, 564 Nitrostyrene hydrogenated, 565 Nitrotoluenes hydrogenated, 378, Cu 512, Pt 564 by hydrogenation, 378 Nitrous acid cat., 82, 184, 269n esters of hydrogenated, 382, 509, 513 Nitrous oxide hydrogenated, 368, 509 Nonane by hydrogenation, 414 Nonene hydrogenated, 414 Nonvlic acid into ald., 852, 853 into ketone, 845 Nonylic aldehyde, 852-854 Nonylic esters dec., 871

Occlusion of gases, 180 Ocimene hydrogenated, 415 Octadiene-diol hydrogenated, 566 Octadiene-diolic acid hydrogenated, 566 Octane by hydrogenation, 414, 601 Octane-diol, 566 Octene hydrogenated, 414, Cu 515, Ni 601 Octodecyl alcohol, 565 Octohydro-anthracene, 29, 363, 390, 483 Octohydro-indol, 571 536. Octohydro-phenanthrene, 484. 592 Octoic acid into ald., 853 Octoic ald., 853 Octomethylene ring hydrogenated, 480 Octo-trienal, 801 Octyl alcohol, 566 Octyl-bensene hydrogenated, 569

Odors of oils elim. by hydrogenation,	by permanganates, 275
939, 965	by persulphates, 276
Oenanthaldoxime, 814	by sulphur trioxide, 272
Oenanthylidene hydrogenated, 425	of oils, 265
Oenanthylidene-acetic acid hydrogen-	of phenois, 11
ated, 417	Oxides, carriers for, Pd 946
Oils hydrogenated in vapor, 939	catalysts, 73, 75, 784, 789, 807-809,
Oils oxidised, 265	813, 823, 837, 848, 858, 906, 921, 934
Oklahoma petroleum cracked, 935	in cracking, 906, 934
Oleic acid, amt. H ₂ required, 955	dec. hydrocarb., 906, 921, 934
in fats, 937	dehydrogenation cats., 638, 789
hydrogenated, 422, 562, 939, 955,	hydrogenation cats., 598
Cu 515, Ni 587, 601, Pd 546, 577,	ketone cats., 848
Pt 558, 565	prep. of, 76
isom., 82, 184	Oxides of carbon hydrogenated, 504
into ketone, 843	Oxides of nitrogen cats., 269n
Oleic esters in fats, 937	hydrogenated, 374
hydrogenated, 577, 601	Oximes hydrogenated, 383, 514
Oleic alcohol hydrogenated, 565	hydrolyzed, 332
Oleine, 937, 939, 955	Oxygen absorbed by C, 180b, by Au, Pt
amt. H ₂ required, 955	and Ag 137
into stearine, 939	Oxygen in catalysts, 165, 563, 563n,
Olive oil hardened, 966	943n
iodine number, 938	in Pt black, 563
Optical isomers, 186	Oxygenation of cat., $943n$, $947n$
Organic Mg compounds, 300–302	T 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Origin of petroleum, 925–928	Palladium absorbs hydrogen, 136, 150,
Osmium cat., 64, 251	165
Osmium black, 583	on alcohols, 669
Osmium oxide hydrogenation cat., 80,	on aldehydes, 623
583	amount of required, 951
oxidation cat., 262, 271	black, 251, 562, 573-579, 822
Oxal-acetic acid by oxid, 268	cat., 65, 126, 269n
Oxalic acid cat., 106	colloidal, 71, 141, 544-555, 604
dec., 12, 855	colloidal prep. of, 71
dec. formic, 822	dehydrogenation cat., 648, 649, 651,
esterif., 758	669, 824
by oxidation, 269	hydrogenation cat., 534, 536–595
oxidised, 246	in hydrogenation of fats, 946
retarder, 11	poisoned, 180o
Oxalic esters dec., 873	polymeris., 212
Oxamide, 105, 312	sponge, 604
Oxidation, 64, 150, 152, 244-277	temp. of use, 952
catalysts, 59, 60, 100, 152, 162, 245- 267	Palladium black, 251, 562, 573-579, 822 palladium hydride, 150
by chlorates, 271	Palladium sponge, 604
with gaseous oxygen, 244-267	Palladous chloride, 562
by hydrogen peroxide, 268	Palm oil bleached, 265
by hypochlorites, 270	Palmitic esters, 937
by nitric acid, 269n	Parabutanal formed, 223
by nitrobensene, 277	Paraldehyde cond., 801

crotonized, 795, 801 by dehydrogenation, 642 depolymerized, 234 ethers of formed, 75, 904, 785-789 formed. 82, 104, 222, 223, 724 formed, 150, 293, 404, 405, 843 Para-indene formed, 217 hydrogenated, 120, 444, 456, Ni 603, Parapropional, 223 Pt 569 Peanut oil. 938 by hydrogenation, 381 hardened, 966 by oxidation, 263, 268 Pelargonic acid into aldehyde, 852 into thiophenol, 791 esterif., 771 Phenol ethers formed, 785–789 Pennsylvania petroleum cracked, 911 hydrogenated, 494 nature of, 925 Phenols with aldehydes, 792 Pentachlorpropane, 242 condensed, 803 Pentachlorpropylene formed, 879 dehvdrated, 785, 789 Pentadecyl-benzene hydrogenated, 569 hydrogenated, 370, 456, 603 Pentamethylene ring hydrogenated, 474 nitrated, 269n Penta-isobutanal formed, 224 Phenolic glucosides, 793 Pentamethyl-bensene dec. F. and C.. Phenylactaldehyde by dehydrogenation. 887 857 Pentane from C₂H₂, 211 hydrogenated, Pd 549, Pt 560 dec. by Ni, 911 Phenyl acetate dec., 871 formed, 211, 558, 565, 931 Phenylacetic acid into ald., 853 Pentane-diol dehydrated, 726 dec., 830, 839 by hydrogenation, 595 esterif., 756-758 Pentane-thiol, 745 hydrogenated, 471 Pentol-one, 439 into ketone, 843-845, 850 Perchlorbenzene reduced, 404 Phenyl-acetylene hydrogenated. Perchlorethane prep., 289 Cu 520, Pd 548 Perchlormethyl mercaptan, 278n Phenyl-alkyl ethers formed, 789 Perhydroanthracene, 29, 363, 483, 592 hydrogenated, 464 Perkin's syn., 107 Phenylation of amines, 632 Phenyl-benzyl carbinol dehydrated, 714 Permanganates as oxidising agents, 275 Peroxides as intermediate comps., 150-Phenyl bromide in syn., 901, 904 153 Phenyl-butyl chloride in F. and C. syn., Persulphates oxid. agts., 276 Perylene, 685 Phenyl-carvacryl ether, 788 Petroleum cracked, 254n Phenyl chloride in syn., 904 by AlCla, 935 Phenyl-p.cresyl carbinol red., 369 formation, 506, 925-928 Phenyl-cresyl ethers, 788 Phellandrene, 198 Phenyl-p.cresyl-methane by hydrogena-Phenanthrene cond., 806 tion, 369 from cracking, 909 Phenyl-cyclohexane, 452, 475 hydrogenated, 484, 642, Ni 592, Pd Phenyl-cyclopentane by F. and C. syn., 536, 579, Pt 571 Phenanthridene oxidised, 270 Phenyl-cyclohexane formed, 889 by oxidation, 270 hydrogenated, 475 Phenetol, 464 Phenylene diamines by hydrogenation, Phenol from bensene, 150, 843 from bromphenols, 405 Phenylene-naphthalene oxides, 788 from chlorohenols, 404 Phenylene sulphide formed, 295 dehydrated, 16, 785 Phenyl esters dec., 871 by diaso, 606 Phenyl ether, 338, 785-787

hydrogenated, 494, 589	hydrogenated, 420, Pd 547, 549, Pt
formed, 59, 75, 786, 904	567
Phenyl ethers, 785–788	Phosgene formed, 134, 282, 282n, 284
Phenylethyl alcohol dehydrogenated, 657	Phosphine cat., 780
hydrogenated, 369	cat. poison, 180ø formed, 700
by hydrogenation, 560	
Phenylethyl chloride in F. and C. syn., 897	Phosphorus cat., 687, 689, 691, 696 Phosphorus cat., 46, 687
Phenyl-ethylene hydrogenated, 415, 451,	chlorination cat., 281
516	oxidised, 150
by hydrogenation, 520	toxic to cats., 115, 116
Phenyl-ethylene hydrocarbons hydro-	Phosphorus, red dehydration cat., 700
genated, 415	Phosphorus trichloride chlorination
Phenyl-ethyl ketone hydrogenated, 539	cat., 281
Phenyl-glycolic acid esterif., 756	Phthaleines, 90
Phenylhydrasine dec., 91, 611	Phthalic acid esterif., 756
from phenylhydrasones, 332	hydrogenated, 392, Ni 590, Pt 563, 569
hydrogenated, 497	by oxid., 273
negative cat., 11	Phthalic anhydride cond., 107
Phenylhydrasones dec., 633, 635	by oxid., 260n, 273
hydrol., 332	Phthalid by hydrogenation, 392
Phenyl-hydroxy-crotonic acid, 203	Phthalimide hydrogenated, 569
Phenyl iodide in syn., 904	Phthalophenone by F. and C., 893
Phenyl-isocrotonic acid hydrogenated,	Phthalyl-acetic acid, 107
417	Phthalyl chloride in F. and C. syn., 893
Phenyl isocyanate hydrogenated, 495	Physical cond. of cat., 41, 53-55, 76-80,
Phenyl-naphthyl-amine, 632	703
Phenyl-naphthyl ketone hydrogenated,	Physical theory of catalysis, 131 et seq.
685	Phytane, 565
Phenyl-nitrosamine formed, 206	Phytene hydrogenated, 565
Phenyl oxide by diaso, 59	Phytol hydrogenated, 565
formed, 75, 338, 785–787, 904	Picoline, 680
hydrogenated, 494, 589	Pinacoline, 724
hydrol., 16, 338	Pinacones, 195, 724, 726
Phenyl-naphthyl ethers, 788	Pinane, 552, 591, 594
Phenyl-pentyl chloride in F. and C.	Pinene cracked, 909
syn., 897	dec., 235, 909, 922, 923
Phenyl-propiolic acid hydrogenated, 548	dehydrogenated, 664
Phenyl-propionic acid, 417, 546, 560, 580,	hydrated, 307
581, 594, 601	hydrogenated, 477, Cu 594, Ni 591,
dec., 839	Pd 552, Pt 570
into ketone, 844	isom., 198
Phenyl-propyl alcohol, 560, 568	polym., 216
Phenyl-propylene by hydrogenation, 384	Piperidine, 486, 555, 561
Phenyl-propyl-pentane by hydrogena-	alkylated, 741
tion, 415	cat., 804, 836
	dehydrogenated, 647
Phenyl-propyl-pentene hydrogenated, 415	in vulc. of rubber, 104
Phenyl-pyridines, 807 Phenyl gulphide formed, 205	Piperonal hydrogenated, 568 Piperonal-agetona hydrogenated, 565
Phenyl sulphide formed, 295	Piperonal-acetone hydrogenated, 565 Piperonal-acetole acid hydrogenated, 601
Phorone by cond., 797	Piperonyl-acrylic acid hydrogenated, 601

Piperonyl-propionic acid, 601 Polycyclic hydrocarbons hydrogenated. Pipervlene by dehydration, 726, 784 polym., 213 Polymerization, 89, 209-233 Polymethylene hydrocarbons, 535, 926, Pittsburgh gas, 928 Platinum absorbs O₂, 137 asbestos, 247 Polymethylene rings hydrogenated. Pt catalyst, 61, 75, 126, 180c, 342, 539, 535 Polyphenois hydrogenated, 460 563n, 615, 829 in combustion anal., 250 Polyphenyl hydrocarbons hydrogenated, 452 in cracking, 906 dec. acetylene, 913, 914, 920 Polysaccharides hydrol., 323 Polyterpenes from cracking, 909 dec. alcohols, 668 dec. alds., 622 Polyvalerylene formed, 212 Poppyseed oil, 938, 966 dec. ethylene. 912 hardened, 966 dec. formic esters, 867 dehydrogenation cat., 636, 637, 643, Porous substances, 139 649, 651, 668 Potash as cat., 611, 795 Potassium cat. polym. hydrocarbons. hydrogenation cat., 524-535, 945 moss. 524 213, 232 Potassium acetate cat., 107 oxidation cat., 4, 15, 61, 154, 235, 245, Potassium bisulphate cat., 97, 687, 725 255, 249, 250, 255, 256 oxidation cat. for SO2, 4 cat. esterif., 759, 760, 783 poisoned, 180o Potassium chloride cat., 876 spiral, 829 Potassium evanide in aldolization, 95 wire, etc., 249 cat., 230 toxic to Pt. 117 Platinum black activity of, 63 cat., 235, 246, 247, 445, 562 Potassium copper cyanide cat., 95 Potassium ferricyanide reduced with Pt. dec. H₂O₂, 2 deoxidising, 14 heat weakens, 63 Potassium formate, 823 Potassium hydroxide cat., 799 hydrogenation cat., 344, 524, 563-572 Potassium iodide cat., 94 oxidation cat., 1, 14 Potassium soaps toxic to cats., 115 poisoned, 117, 947n Preparation of catalysts, 54-56, 58, 59, preparation, 61 Platinum chloride cat., 635 77, 78, 598, 606, 655, 704, 705, 941, 942 Platinum, colloidal, 69, 72, 141, 248, 544, 556-561 Pressure, effect of, 30 poisoned, 116 on dehydration, 711 on hydrogenation, 946, 956 Platinum moss, cat., 524 Platinum sponge cat., 193, 245, 342, 445, on hydrolysis, 317 524, 637, 824 on inversion of sugars, 324 Primary alcohols dehydrogenated, 650 Poisoning of catalysts, 112 et seq., 180o-Promoters, 180s-180u 180s, 946, 947n of Ni cat., 112, 598 Propane from ethyl acetate, 180j by hydrogenation, 414, 472, 912 of Pt cat., 116 Poly-alcohols dehydrogenated, 680, 723, Propane-thiol, 745 Propenol hydrogenated, 416 Propionamide hydrogenated, 386 Poly-aldehydes formed, 222 Poly-alkyl-bensenes dec., 887 Propionic acid dec., 838 Poly-ethyl-bensenes dec. F. and C., esterif., 751, 771 by hydrogenation, 417 888

into ketone, 840, 842-845 Propvi ether, 691, 694 Propionic aldehyde, 416, 419, 658, 664, Propvl iodide, 605 668. 680, 839 Propyl-isosmyl-amine, 738 cond., 795, 808 Propvi malonate dec., 873 crotonised, 795 Propyl mercaptan, 744 dec., Cu 621, Ni 620, Pd 623, Pt 622 Propyl-methoxy-cyclohexanol, 569 into ester, 228 Propyl-methoxy-phenol, 603 formed, 208, 249 Propyl nitrite hydrogenated, 382 hydrogenated, 432 Propyl oxalate dec., 873 by oxidation, 249 Propyl phenyl ether, 789 polym., 223 Propyl-piperidine, 741 Propionic aldehyde phenylhydrasone Propyl propionate from ald., 228 dec., 633 dec., 861 Propionic anhydride into ketone, 857 by esterif., 751 Propionic esters dec., 863, 871 Propyl succinate dec., 873 Propionitrile cat., 605 Protocatechuic ald., by oxid., 268 polym., 231 Pseudocumene hydrogenated, 447 Propionyl chloride cond., 902 isom., 888 into nitrile, 813 Pseudoionone, 800 Propiophenone-oxime hydrogenated, 384 Pseudomorphine by oxid.. 268 Propvl acetate dec., 861 Pulegomenthol, 436, 567 Propvl-acetylene formed, 192 Pulegomenthone, 421, 436 Propyl alcohol, 416, 419, 558, 680, 740. Pulegone, dehydrogenated, 645 741 hydrogenated, 421, Ni 591, Pd 552, into acetal, 780 Pt 567 into amine, 732 Pumice cat., 811, 828 dehydrated, 691, 694, 700, 713, 715carrier, 126, 598 Purification of oils for hydrogenation. 717, 719 dehydrogenated, 656, MnO 672, Ni 947-949 664. Pt 668 Pyridine cat., 187, 224, 836 esterif., 751, 771, 773, 775 by dehydrogenation, 647 in F. and C. syn., 893 by hydrogenation, 416 oxidised, 249, 254, 268 hydrogenated, 486, Pd 555, Pt 561 oxidised, 257 Propyl-amine from alc., 733 sulphonated, 816 by hydrogenation, 382 Propyl-bensene by hydrogenation, 384, in syn., 893, 901 448, 539, 560 Pyridine-carbonic acid hydrogenated, Propyl bensoate, 766 561 Propyl bromide isom., 199 Pyridine homologs hydrogenated, 561 Pyridyl-phenyl ketone by F. and C. syn., Propyl chloride dec., 877 893 isom., 199 Propyl cyanide as cat., 605 Pyrocatechol hydrogenated, 370, 461 by oxid., 268 Propyl-cyclohexane, 449, 590 Propvl formate, 773 Pyrogallol hydrogenated, 462 Propylene, 691, 694, 696, 700, 713, 716, oxidised, 150 Pyrogenetic equilibria, 905 735 from C₂H₂, 916 Pyrography, 249n Pyromucic ald. hydrogenated, 434 dec., Ni 912 formed, 877, 916 Pyrone formed, 835 hydrated, 306n Pyrrol, 686, 807 alkylated, 742 hydrogenated, 414, 515, 526

hydrogenated, 486, 571 Pyrrols cond., 803, 805 Pyrrolidine, 429, 485, 571

Quantity of cat., 32 Quercetine by hydrol., 328 Quinaldine hydrogenated, 488 Quinalizarine by oxid., 274 Quinidine hydrogenated, 555 Quinine as cat., 836 hydrogenated, 604 Quinine sulphate hydrogenated, 572 Quinite, 461, 589 dehydrated, 723 Quinisarine by oxid., 274 Quinoline as cat., 187, 793, 836 by dehydrogenation, 647 in F. and C. syn., 893 hydrogenated, 488, 489, Ni 592, Pd 555, Pt 561 Quinone hydrogenated, 442 by oxidation, 276 Quinones hydrogenated, 442

Radiation theory of catalysis, 180j Reaction tube for catalysts, 347 Reciprocal catalysis, 146 Regeneration of catalysts, 123-125, 563n, 932, 947n, 950 Regeneration of thoria, 708n Resinous substances by oxidation, 266 Resorcine cond., 806 hydrogenated, 370, 461 Reversible reactions, 19, 39 Rhodium cat., 64 Rhodium black cat., 822 in hydrogenation, 581 Ribonic acid formed, 187 Riché gas., 397 Ricinolele acid, 937 Ricinoleic esters. 937 Ring formation, 82, 194, 684, 685, 727, Rubber syn., 106, 213, 214, 784 vulcanis., 104 Ruberythric acid hydrogenated, 328 Russian petroleum cracked, 934, 936

Sabinene hydrogenated, 570

Ruthenium cat., 64

Ruthenium black, 580

Saccharic acid dehydrated, 727 Safrol hydrogenated, Ni 590, Pt 565 Salicine, 329 Salicylic acid esterif., 756-757 by hydrol., 328 Saligenine by hydrol., 329 Saliretine by hydrol., 329 Sand cat., 696, 811 Sandmeyer reaction, 91, 609 Santonin hydrogenated, 571 Saponification, 17, 305, 337 acid radical influence, 316-319 of esters, 337 of fats, 314, 319 neutral salt influence, 317 theory of, 176 Saturated hydrocarbons by hydrogenation, 412-415 Schlinck's apparatus, 960 Schwoerer's apparatus, 959 Sebacic acid into ketone, 843 Secondary alcohols, 420 dehydrogenated, 650, 659 esterif., 766, 775 prep., 435 Secondary amines by hydrogenation, 383 from naphthols, 790 prep., 427, 732 Selective absorption by cats., 180h Selenium hydride toxic to cats., 1800 Selex, 967n Separation of carbon, 613 Secame oil, 938 hardened, 966 Sesquiterpenes hydrogenated, 570 Side chains hydrogenated, Cu 594, Ni Siemens gas, 397 Silica cat., 75, 78, 540, 675, 676, 702, 811, 825, 911 dec. formic acid, 624 ketone cat., 847 prep., 705 Silica gel cat., 75n, 180c, 772n Silicates cat., 99, 267 ketone cat., 847 Silver absorbs O₂, 137 cat., 60 with CO, 615

dec. H₂O₂, 34

oxidation cat., 252, 254, 259	State of cat., 41, 53-55, 76-80
Silver chloride cat., 876	Stearic acid, 422, 515, 546, 558, 562, 577,
Silver colloidal, 70, 72	587, 601
Silver nitrate cat., 276	into ketone, 843, 847, 850
Silver oxide cat., 276	Stearic esters, 937
Silver salts in nitration, 269n	Stearine, 937
	Stearone, 847
Sise of grains, 35	Stearyl chloride hydrogenated, 575
Skatol, 647 Snow drift, 967n	Stibine dec., 8
Soda dehydration cat., 795	toxic to cats., 1800
Sodium isom. cat., 50	Stilbene by dehydration, 714 hydrogenated, 415, 515
polymer. cat., 50, 213, 231, 232	
Sodium acetate dehydration cat., 107,	by hydrogenation, 548
795	Stirring in hydrogenation, 587n, 601 Stoichiometric theory of catalysis, 180a
esterif. cat., 748, 761	
polym. alds., 224	Strontium carbonate cat., 838
Sodium alcoholate cat., 340n	Strychnine hydrogenated, 555
Sodium borate cat., 574	Styrene from acetylene, 914, 916
Sodium carbonate to neut. oils, 948	formed, 241, 520, 548, 657, 889
Sodium chloride cat., 876, 954	hydrogenated, 415, 451, Cu 516, Pd
Sodium formate cat., 822	546, Pt 569 Suberio said 566
Sodium hydroxide dehydration cat., 795,	Suberic acid, 566
798	into ketone, 843
Sodium methylate cat., 799	Succinic acid esterif., 756
Sodium nitrate effect on Ni, 947	by hydrogenation, 546
Sodium sulphide toxic to cats., 947	Succinic anhydride formed, 878, 874
Sodium thiosulphate in isom., 182	hydrogenated, 392
Solvent naphtha cracked, 909	Succinic esters decom., 873, 874
Solvents as cats., 36, 37, 40	Succinimide, 312
in hydrogenation, 599	Succincyl chloride in F. and C. syn., 893
influence of, 38–40, 180f	Sucrose hydrol., 323, 324
influence on equilibra, 39	Sugar oxid., 269
influence on inversion of sugars, 324	Sugars formed, 236
influence on reaction velocity, 38	by hydrol., 323
Sorbic acid hydrogenated, 558	inverted, 175
Sorbite, 588, 595	isom., 186
Sorbose formed, 186	multirotation of, 188
Specificity of cats., 142	Sulphates effect on, Ni 947
Spirocyclane hydrogenated, 535	Sulphides, 743, 744
Squibb's method, 161	Sulphocyanic esters, 333
Stabilisers, 13	Sulphonation, 815, 816
Stannic chloride acetylation cat., 240	aided by HgSO ₄ , 102
chlorination cat., 283, 288	Sulphur added, 295, 296
cond. cat., 243	catalyst, 6, 45, 630
in F. and C. syn., 899	chlorine cat., 280
Stannic oxide ketone cat., 849	eliminated from petroleum, 933
Stannous oxide cat., 288, 539, 676	toxic to cats., 115, 116, 947
dec. alcohols, 673	Sulphur compounds in hydrogenation,
dehydrogenation cat., 673, 824	569
Starch cat., 269n	toxic to cats., 946
hydrolyzed, 4, 323, 326	Sulphur dioxide added, 87, 297

cat., 74 oxidised with Pt 4, 247 polym. alds., 222 Sulphuric acid on alcohols, 159 catalyst, 687, 689, 691, 696, 713 cond. agt., 795, 803 in esterif., 748, 749, 751, 752, 756, 758 on formald., 822 in hydration, 306, 308 isom. terpenes, 198 manufacture, 32, 158 by oxidation, 258 Sulphuric acid furning as oxid. agt., 272-Sulphur trioxide cat., 12 manuf., 247 oxidising agt., 272 Sunlight in chlorination, 281n Surface, importance of, 35 Sylvestrene hydrogenated, 477 Synthetic tallow, 967

Tagatose formed, 186 Talgol, 967 Tallow, 938 hardened, 966 Talomucic acid, 187 Talose formed, 186 Tartaric acid esterif., 756 toxic to cats., 115 Tellurium oxid. cat., 45, 251 Tellurium hydride toxic to cats., 1800 Temperature coef. in dehydration, 709 coef. of reactions, 24, 25 effect on hydrocarbons, 905 of hydrogenation, 361, 952 of prep. of cats., 707n regulation, 348 Terephthalic acid, 648 Terpenes dec., 922 dehydrogenated, 643 hydrogenated, 477, Ni 591, Pt 570 isom., 198 Terpine, 307, 308 Terpinene formed, 198 Terpineol dehydrogenated, 645 hydrogenated, 478, 552 Terpinolene formed, 198 Tertiary alcohols esterif., 778

Tertiary butyl alcohol oxidised, 249

Tetra-acetyl-phenyl-glucoside, 793 Tetra-amylene formed, 211 Tetrabromethane in F. and C. syn., 897 Tetrachlorethane dec., 881 formed, 199 in syn., 903 Tetrachlorethylene formed, 879 Tetracosene formed, 210 Tetra-ethyl-ammonium iodide, 38 Tetrahydro-acenaphthene, 482 Tetrahydro-anthracene, 29, 363, 483, 592, 642 Tetrahydrobenzoic acid. 476 Tetrahydrocarvone, 552, 567 Tetrahydrocolchicine, 555 Tetrahydrodoremone, 570 Tetrahydrofurfuryl-ethyl carbinol, 487 Tetrahydroionones, 554 Tetrahydro-methyl-furfurane, 487 Tetrahydro - methyl - naphthalene - carbonic acid, 563 Tetrahydronaphthalene, 379, 481, 481n, 571, 592, 594 Tetrahydronaphthoic acid, 594 Tetrahydronaphthalid, 563 Tetrahydrophenanthrene, 484, 536, 579, 592, 642 Tetrahydrophenol, 723 Tetrahydropiperine, 555 Tetrahydroquinoline, 488, 561, 592 dehydrogenated, 647 Tetrahydrosantonine, 571 Tetrahydrostrychnine, 555 Tetrahydroterephthalic acid, 648 Tetrahydroxyanthracene oxid., 274 Tetrahydroxyflavanol by hydrol., 328 Tetra-isobutanal formed, 224 Tetralin, 481n Tetramethyl-bensene dec., F. and C., Tetramethyl-diamino-benshydrol cond., 730 Tetramethyl-leucaniline, 730 Tetramethylene-diamine, 429 Tetramethylene ring hydrogenated, 473 Tetraphenylethane, 538, 662, 720 hydrogenated, 453 Tetraphenylethylene, 736 Tetrolic acid hydrogenated, 546 Thalium cat., 47 Thalium chloride chlo. cat., 283

Theories of catalysis, 129 et seq., 131, chlorination cats., 283, 288 145, 180, 180a-180u cond. cat., 243 Theory of dehydration, 785 Tin oxides chlorination cats., 288 Theory of esterification, 752, 762, 763 dehydration cats., 702 Theory of ester decomp., 859-864, 866, Titania cats., 75, 143, 337, 624, 693, 702, 704, 709, 732 872 Theory of poisoning catalysts, 1800 aldehyde cat., 852 Thianthrene, 629 in cracking, 906, 934 Thiobensophenone syn., 894 dec., alcohols, 180g Thiodinaphthyl-amines formed, 296 dec. esters, 180b, 180f, 180j, 180n, Thiodiphenyl-amine formed, 296 861n, 863, 864, 868 Thio-indigo hydrogenated, 603 dehydration cat., 825 Thiophene formation, 686, 810 dehydrogenation cat., 686 toxic to cats., 112, 947n esterif. cat., 765, 767, 771, 772, 775 Thiophenol formed, 295 hydrolyt. cat., 686 Thiophenols dec., 629 ketone cat., 849 formed, 295 mixed cat., 675, 676 Thiophosgene in F. and C. syn., 894 prep. of, 704, 861n Thiols formed, 75, 743-746 Tolane, hydrogenated, 548 Thiourea isom., 207 Toluene, 465, 560, 590, 593, 641, 657, Thioureas, 630 681 Thoria catalyst, 16, 24, 75, 79, 143, 170, brominated, 292, 293 538, 676, 693, 700, 702, 707, 716, chlorinated, 278, 281, 285 from cracking, 908, 909 720, 731-738, 813 from cresoles, 370 aldehyde cat., 854 dec. chlorides, 881 from cymene, 930 dec. by F. and C., 887 dec. esters, 858, 861, 861n, 864-866, 872, 873, 180b, 180f, 180j, 180n by F. and C. syn., 884 in F. and C. syn., 899 dec. formates, 870 dehydration cat., 651, 743-746, 788, hydrogenated, 444, 447, Pt 534, 560, 789, 791, 801, 808, 809 569, Rh 581, Ru 580 dehydrogenation cat., 686 by hydrogenation, 369, 388 esterif. cat., 764-766, 772 oxidised, 257, 260n, 263 hydrolysis cat., 337, 338 from petroleum, 934 life of, 708 from pinene, 922 ketone cat., 840, 844, 848, 850, 857 sulphonated, 815n mercaptan cat., 746 from xylene, 930 Toluic acids dec., 830 mixed cat., 826 preparation of, 707n, 861n esterif., 758, 766 Thorium chloride cat., 90 into ketones, 848, 849 Thujane, 570 Toluic aldehydes dec., 623 Thujone hydrogenated, 478, 552, 570 syn., 298 isom., 198 Toluic esters dec., 864, 871 hydrogenated, 471 Thymol, ethers of, 789 Toluidines, 497, 564, 630-632, 683, 684. formed, 645 hydrogenated, 459 790 alkylated, 740 Thymoquinol hydrogenated, 463 hydrogenated, 467 Thymoguinone hydrogenated, 442 Tin chlorination cat., 47, 288 by hydrogenation, 380 dehydrogenation cat., 673, 824 manufacture of, 512 Tin chlorides acetylation cat., 240 oxidised, 256

Toluonitrile hydrogenated, 428	Triphenyl carbinol reduced, 369
Toluquinone hydrogenated, 442	Triphenylene, 646
Tolyl-dimethyl carbinol hydrogenated, 369, 465	Triphenyl-methane formed, 369, 728, 890
Toxic substances removed, 947-949	hydrogenated, 453
Toxicity of CO, 953	Tungsten as promoter, 180s
Toxicity scale, 116	filament, 180s, 180p
Trade names of hardened oils, 967	Tungsten, blue oxide, cat., 24, 693, 825
Trehalose hydrol., 323, 325	dehydration cat., 651, 791
Triacetin, 760	on formic acid, 624
Tri-anylene formed, 211	mercaptan cat., 746
Tribromphenol, 293	Tungstic oxide, 75
hydrogenated, 405	Turpentine oxid., 151
Tributene formed, 210	Types of hydrogenation apparatus, 957,
Trichlorbensene, 404	964
Trichlor-tert.butyl alcohol, 238	
Trichlorethylene cond., 242	Undecenal, 658
formed, 881	Undecenyl alcohol, 658
Trichlorethyl trichloracetate, 228	Undecylenic acid hydrogenated, 417
	Undecylic acid, 417
Trichlorphenol, 404	Unsaturated acids esterif., 756
Tricyclohexyl-methane, 453	
Triethyl-amine cond., 38	hydrogenated, 422
formed, 377, 427	isom., 203
Triethyl-amine hydrochloride cat., 783	Unsaturated alcohols hydrogenated, 416,
Triheptene, 519	418, 419
Tri-isoamyl-amine, 682	into sat. alds., 208
Trimethyl-amine cat., 224	into cat. ketones, 208
formed, 377, 496	Unsaturated chlorides dec., 876
Trimethyl-benzenes dec., 887	Unsaturated esters, 937
Trimethyl carbinol dehydrated, 713,	Unsaturated hydrocarbons, 743, 764,
719	802, 866
esterif., 776	dec., 912
into ether, 691	formed, 75, 142, 169, 695 et seq., 871
formed, 306	872, 876, 878
Trimethyl-cyclohexanes, 449	hydrated, 305
Trimethylene bromide, 605	Unsaturated ketones hydrogenated, 420
Trimethylene ring hydrogenated, 472	602
Trimethyl-ethylene dec., Ni, 912	Uranium in drying oils, 266
formed, 190	as promoter, 180s
hydrogenated, 414	Uranium chlorides cats., 90, 283
polym., 210	Uranium oxide cat., 75, 260, 702
Trimethyl-hydroxy-butyl-cyclohexane, 560	dec. alcohols, 142 ketone cat., 840, 849
Trimethyl-nonenone hydrogenated, 420	oxidation cat., 259
Trimethyl-pentane by hydrogenation,	Uranous oxide cat., 675, 676, 726
414	825
Trimethyl-pyrasoline formed, 196	dehydration cat., 791
Trioxymethylene into acetal, 781	mercaptan cat., 746
cond., 792, 806	Uranium soaps toxic to cats., 115
formed, 432	Urea acetylated, 87
in syn. rubber, 215	Urethane oxid., 259
	

Valeric acid from alcohol, 150 into ketone, 842, 845 Valerie estera dec., 863 Valerolactone, 437 Valerone, 547, 549 Valerviene polym., 212 Vanadium chloride chlorination cat., 283 Vanadium pentoxide cat., 269, 676, 693, 702, 828 oxidation cat., 260, 260n, 262n, 271 Vanadium sulphate in sulphonation, 816 Vanadous oxide cat., 675 Vanilline acetylated, 240 hydrogenated, 568 by oxidation, 191, 249 Vanilline triacetate, 240 Various rings hydrogenated, 592, 603 Velocity of catalytic reactions, 23 Vinvl bromide in F. and C. svn., 889 Vinyl-trimethylene hydrogenated, 577 Volume of hydrogen reg, by various oils. 955

Walls of vessel as cat., 244n
Water as cat., 73, 249
effect on dehydration, 710
effect on ethylene prep., 180h
effect on hydrogenation, 949
neg. cat., 12
Water gas, 398, 402
for hydrogenation, 953
reducing agent, 511
Wilbuschewitch's apparatus, 961
Williamson's reaction, 159, 169
Whale oil, 938
effect on cat., 947
Woltman's apparatus, 964
Wurts syn., 11

Xylenes brominated, 292 chlorinated, 278, 285 add CO, 298 from cracking, 908, 909 from cymene, 930 dec., by AlCl₃ 887, 930 by dehydrogenation, 641 hydrogenated, 444, 447, 534, 569 from pinene, 922 isomer., 888 Xylenols, ethers of, 786, 789 hydrogenated, 458 Xylonic acid formed, 187 Xylose, 188

Zinc cat. cond. cat. 52, 795 dec. alcohols, 670 dehydrogenation cat., 670, 678 in F. and C. syn., 899 hydrogenation cat., 595 oxidation cat., 269n polym, alds., 219 toxic to cats., 115, 946 Zine bromide bromination cat. 293 isom. cat., 200 Zinc carbonate cat., 824 Zinc chloride bromination cat., 293 cat., 6, 89, 234, 240, 283, 633, 635, 687, 689, 691, 695, 698 cond. cat., 795, 796, 803 esterif. cat., 748, 761, 795 in F. and C. syn., 899 hydrol, cat., 330 polym. cat., 211, 216, 222 Zinc hydroxide isom. sugars, 186 Zine organo-compounds, 304 Zinc oxide cat., 75, 143, 539, 675, 676 in cracking, 906, 934 dec. formates, 869 dehydration cat., 702 dehydrogenation cat., 824 hydration cat., 310 ketone cat., 841, 849 Zirconia, amine cat., 732 cat., 746, 791, 825, 840, 849 esterification cat., 772n dec. formic ac., 624 dehydration cat., 791 dehydrogenation cat., 676, 693 ketone cat., 840, 849 mercaptan cat., 746 mixed cat., 651, 675, 702