COMPRESSIBLE LAGRANGIAN HYDRODYNAMICS WITHOUT LAGRANGIAN CELLS

FREE-LAGRANGIAN METHODS CONFERENCE
Science Applications International Conference
134 Holiday Court, Suite 318
Annapolis, Maryland 21401

DISCLAIMER

This report was prepared as an account of work sponsored by an agency of the United States Government. Neither the United States Government nor any agency thereof, nor any of their employees, makes any warranty, express or implied, or assumes any liability or responsibility for the accuracy, completeness, or usefulness of any information, apparatus, product, or process disclosed, or represents that its use would not infringe privately owned rights. Reference herein to any specific commercial product, process, or service by trade name, trademark, manufacturer, or otherwise does not necessarily constitute or imply its endorsement, recommendation, or favoring by the United States Government or any agency thereof. The views and opinions of authors expressed herein do not necessarily state or reflect those of the United States Government or any agency thereof.

By acceptance of this article, the publisher recognizes that the U.S. Government retains a nonexclusive, royalty-free license to publish or reproduce the published form of this contribution, or to allow others to do so, for U.S. Government purposes.

The Los Alamos National Laboratory requests that the publisher identify this article as work performed under the auspices of the U.S. Department of Energy.
COMPRESSIBLE_LAGRANGIAN_HYDRODYNAMICS_WITHOUT_LAGRANGIAN.Cells

Robert A. Clark
Computational Physics
Group X-7, MS B257
Los Alamos National Laboratory
Los Alamos, New Mexico 87545

I. INTRODUCTION

The formulation normally used to calculate compressible Lagrangian hydrodynamics in two dimensions is the following. First define a two-dimensional mesh containing a set of Lagrangian cells. Assign each cell a fixed mass. Compute the acceleration of the mesh points and move the points. The volume of the cell changes with the motion of the points. The changes in cell density, energy, and pressure are computed from the changes in volume. Difficulties occur when there are large distortions in the flow that cause similar large distortions in the Lagrangian cells. The usual solution is to somehow adjust the mesh as the calculation proceeds. This involves either moving individual mesh points or actually re-connecting the mesh. In either case, it becomes necessary to re-map the mass from the old cells to the new. This necessarily produces some amount of undesirable numerical diffusion. When and how to adjust the mesh and how to accurately re-map the mass and other variables so as to minimize numerical diffusion are the problems.

One way to eliminate these problems is to abandon the idea of the Lagrangian cell since it is the distortion of the Lagrangian cell that is the cause of all the other problems. In the next section we will discuss how the conservation equations can be solved directly without resorting to Lagrangian cells. Next we will give some examples of calculations using this method. Finally, we will give details of the calculational method presently being used.

II. SOLVING THE CONSERVATION EQUATIONS

The equations we are trying to solve can be written

\[ \frac{D}{Dt} \rho - \rho \nabla \cdot v = 0 \]  [1.1]
\[
\frac{D}{Dt} \vec{u} = -\frac{1}{\rho} \vec{\nabla} P \tag{2.2}
\]
\[
\frac{D\rho}{Dt} = -\frac{P}{\rho} \vec{\nabla} \cdot \vec{u} \tag{2.3}
\]
\[
P = P(\rho, e) \tag{2.4}
\]

where \(\vec{u}\) represents the vector velocity, \(\rho\) the density, \(e\) the specific internal energy and \(P\) the pressure of the fluid. Equation [2.1] expresses conservation of mass, [2.2] conservation of momentum and [2.3] conservation of energy. The Lagrangian time derivative, i.e., the derivative following the fluid, is indicated by \(\frac{\partial}{D t}\).

In a standard Lagrangian calculation only Eq. [2.2], the momentum equation is solved directly. The procedure is to integrate [2.2] over some region of space to arrive at the acceleration of each mesh point. The mesh points are then moved and the new cell volumes along with the fixed cell mass determine the new density, hence, indirectly solving Eq. [2.1]. The associated \(PdV\) work term updates the cell energy and indirectly solves Eq. [2.3] and the new pressure is obtained from the equation of state [2.4].

We propose the following: Instead of Lagrangian cells, we think of a set of Lagrangian points which are embedded in and move with the fluid. There is no mass associated with these points. They are just moving tracer points at which we will attempt to keep track of the velocity, density, energy, and pressure of the fluid. In our later example calculations we will show point positions at various times in the calculation. At each of these points, we know the density, energy and velocity of the fluid, but we do not associate any particular mass with the point.

Looking now at Eq. [2.1], we note that to approximate the time integral of the density change from time \(t\) to time \(t + \delta t\) we need an approximation to \(\vec{\nabla} \cdot \vec{u}\) at that point. To solve Eq. [2.2], we need an approximation for \(\vec{\nabla} P\) and for [2.3] we again need \(\vec{\nabla} \cdot \vec{u}\). To obtain these, we select a set of "representative" neighbors. We then make a
finite difference approximation to $\hat{\nabla}P$ and $\hat{\nabla} \cdot \vec{u}$, using these neighbors, and update $p$, $\vec{u}$, and $e$ at each point. Each point is then moved the distance $\vec{u} \Delta t$ and one time step is completed.

At the next time step the selection of a set of "representative" neighbors may change, but this does not require any sort of re-mapping of variables. It only means that a different set of points will be used in the next finite difference approximation to $\hat{\nabla} \cdot \vec{u}$ and $\hat{\nabla}P$. Large distortions in the flow will produce frequent changes in neighbor selection, but since there are no cells to distort and no remapping to be done the calculation proceeds from cycle to cycle with no difficulty.

III. SOME EXAMPLE CALCULATIONS

3.1 Here we will give three examples of calculations performed by the code HOBO using the free Lagrangian method described herein. The first test problem is the implosion of a gaseous sphere. The initial condition is a sphere of perfect gas with a gamma of $5/3$. The gas is divided into four regions as seen in Fig. 3.1. Pressures are in megabars, density in gm/cc and dimensions in cm.

![Fig. 3.1](image)

The high pressure in region IV will drive a spherical implosion which will greatly compress region III, II, and particularly I. There are two challenges to this problem, the first is to maintain a spherical ball while running the calculation in cylindrical $(r, z)$ geometry. Six snapshots of region II are shown in Fig. 4.2. Region I is interior to region II. The minimum volume of region I occurs in the fifth snapshot after which region I begins to expand. We can hope...
cycles with 73 points in the radial direction and 64 points covering 180° of angle. The left half of the snapshot is a reflection of the right half which was calculated.

Fig. 3.2

The second challenge is the accuracy of the solution. For comparison purposes we ran a standard one-dimensional Lagrangian code using 800 zones, 200 zones in each region. In Figs. 3.3.a, b, c, and d. We have plotted the average density and average specific internal energy in regions I and II as calculated by HOBO with 73 points in the radial direction and the one-dimensional Lagrangian calculation with 500 points. We feel the agreement to be quite good. One notable difference is the time at which minimum volume is reached. HOBO is
slow by about .0075 \textmu sec or $\frac{1}{2}$ of the problem time at that point. Since average density and energy are integral quantities we have plotted one of the variables as a function of radius in Fig. 3.4. We chose radial velocity, but the agreement in all other variables is very similar. The plots are from slightly different times to compensate for the time shift just mentioned. The 1D Lagrange plot is at 2.125 \textmu sec and the HOBO plot is from 2.25 \textmu sec. Apart from the inability of the more coarsely zoned HOBO to resolve the shock front at the radius 1.2 cm we feel the agreement is excellent. The time chosen for the plot is late in the calculation when region II has expanded almost back to its original volume.

![Graph showing radial velocity vs radius](image)

Fig. 3.4

3.7. For our second test problem we have chosen a Meshkov instability\cite{1} based on the geometry used in one of Meshkov's experiments. The initial conditions are shown in Fig. 3.5. A piston driven shock is driven through a region of air and then helium. The

![Table of initial conditions](image)

Fig. 4.5
air to helium density ratio is just over 7. There is an initial perturbation in the air -- He interface which grows with time after the shock passes through the interface. In Fig. 3.6 we plot several snapshots of the Lagrangian point positions in the air (the He is not plotted). For comparison purposes we ran the same problem on a two-dimensional Eulerian code with the cell size similar to the point

Fig. 3.6

separation used in HOSC. In Figs. 3.7a and b, we compare the size of the perturbation as it grows in time. In 3.7.a the initial perturbation, \( \delta \), is .2 cm and in 3.7.b it is .4 cm in width. The agreement between the two codes is excellent.

Fig. 3.7.a, b

3.3. Our third test problem is the penetration of a concrete plate by a steel rod moving at an initial velocity of .7144 cm/sec. The rod is 9.064 cm in diameter and 45 cm in length. The concrete is 50 cm thick. In Fig. 3.8 we show six snapshots of the rod penetrating the concrete. Incompressible theory\(^2\) predicts a constant time rate of
change in the length of the steel rod. The sound speed in the rod is \(0.4545 \text{ cm/\mu sec} \) and \((v/c)^2 = 0.22\), so this problem should not be too far from the incompressible solution. As is shown in Fig. 3.9, the rod length as a function of time matches the incompressible theory very well. Calculations with a two-dimensional Eulerian code produced an almost identical result.
IV. THE FINITE DIFFERENCE SCHEME

4.1 The pressure gradient

We want to approximate $\hat{P}$ at the point $k$ whose neighbors are the points $k_1, k_2, \ldots, k_{n_{\text{max}}}$. Our neighbor selection guarantees at least three neighbors for each point, the average is six and there is no maximum number. Clearly there are many methods that could be used to approximate $\hat{P}$. The following was arrived at through much trial and error and appears to work very well.

Consider point $k$ in Fig. 4.1 that has five neighbors. We construct a polygon with vertices midway between the point and each of its neighbors. The position of the $n$th vertex is $\hat{x}_n = 1/2 (\hat{x}(k) + \hat{x}(k_n))$ and the vector from $\hat{x}_k$ to $\hat{x}_n$ is denoted by $\delta \hat{x}_n = \hat{x}_n - \hat{x}_k$. The pressure at the $n$th vertex, $P_n$, is a weighted average of $P(k)$ and $P(k_n)$ (to be described in Section 4.3). We assume a linear pressure distribution along each edge of the polygon and integrate the pressure over the surface to get a force $\mathbf{F}$. We assume a constant density $\rho_k$ over the polygon to calculate a mass $M$. Then we have $\frac{D}{Dt} \mathbf{U} = -\mathbf{F}$. Now let $\hat{x}_n' = \hat{x}_k + \delta \hat{x}_n$ and the pressure at the new vertex is $P_n' = P_k' + r(P_n - P_k)$. Now $r$ and $M$ are functions of $t$ and we compute

$$\lim_{t \to 0} \mathbf{F}(t) \mathbf{M}(t)$$
The resulting expression for the pressure gradient is

\[ \mathbf{\hat{P}}_k = \frac{x \sum P_n (\delta y_{n+1} - \delta y_n) + y \sum P_n (\delta x_{n+1} - \delta x_n)}{\sum (\delta x_{n+1} \delta y_n - \delta y_{n+1} \delta x_n)} \]  \hspace{1cm} [4.1] \]

where \( \mathbf{\hat{x}} \) and \( \mathbf{\hat{y}} \) are respectively the unit vectors in the x and y directions and \( \delta \mathbf{\hat{x}}_n = \delta x_n \mathbf{\hat{x}} + \delta y_n \mathbf{\hat{y}} \).

If the preceding is done in cylindrical geometry, the result is identical for \( \mathbf{\hat{P}} \) with x and y replaced by r and z. It is of interest to note that if the \( \frac{\partial}{\partial n} \) is not taken, the result does not give a spherically symmetric pressure gradient in a spherically symmetric problem using cylindrical coordinates.

There is an easier way to arrive at Eq. [4.1] although the method just described is how we originally derived it. Since it takes only three points to describe a plane surface, each consecutive pair of neighbors along with the point k defines a pressure plane to first order. If we assign a weight to each of these approximations we have an approximation for \( \mathbf{\hat{P}} \). If the weighting function is the area of the triangle formed by the three points, the result is the same as Eq. [4.1]. We have tried other weighting functions, \( \Theta \) and \( \sin \Theta \) where \( \Theta \) is the angle between \( \delta \mathbf{\hat{x}}_n \) and \( \delta \mathbf{\hat{x}}_{n+1} \) both work fairly well, but area weighting appears to be best at this time.

### 4.2 The divergence of the velocity field

In cartesian coordinates we represent the velocity at the point k by \( \mathbf{\hat{U}}_k = u_k \mathbf{\hat{x}} + v_k \mathbf{\hat{y}} \). The divergence of the velocity field can be expressed as \( \nabla \cdot \mathbf{\hat{U}} = \frac{1}{V} \frac{\partial V}{\partial t} \) where \( V \) is the specific volume of the fluid. Referring back to Fig. 4.1 the specific volume of the constructed polygon is proportion to the area of the polygon given by

\[ A = 1/2 \sum_n (x_{n+1} + x_n) (y_{n+1} - y_n) \]
Hence we can write

\[
\Phi \cdot \bar{U} = \frac{1}{A} \sum \frac{u_{n+1} + u_n}{\Delta t} (y_{n+1} - y_n) + \frac{(x_{n+1} + x_n)}{\Delta x} (v_{n+1} - v_n) - \sum \frac{x_{n+1} y_n - y_{n+1} x_n}{4A}
\]

Equation 4.2 can be derived directly from Eq. 4.1 by noting that 4.1 implies a definition for the operators \( \frac{\partial}{\partial x} \) and \( \frac{\partial}{\partial y} \) and when these are applied to \( \Phi \cdot \bar{U} = \frac{\partial u}{\partial x} + \frac{\partial v}{\partial y} \) Eq. [4.2] is obtained. Thus, we have in effect three ways of deriving the same finite difference approximation to the operators \( \frac{\partial}{\partial x} \) and \( \frac{\partial}{\partial y} \). In cylindrical coordinates we express the divergence of the velocity field as

\[
\Phi \cdot \bar{U} = \frac{1}{r} \frac{\partial}{\partial r} (ru) + \frac{\partial v}{\partial z} = \frac{u + \frac{\partial u}{\partial r} + \frac{\partial v}{\partial z}}{r}
\]

where \( \frac{\partial u}{\partial r} + \frac{\partial v}{\partial z} \) is calculated by Eq. [4.2] with \( x, y \) replaced by \( r, z \).

4.3 The midpoint pressure and velocity

In 4.1 we use a pressure \( P_n \) which is midway between points \( k \) and \( k_n \). This is not a numerical average. Consider the one-dimensional problem depicted in Fig. 4.2.a.

What pressure should we use for \( P_{i+1}^+ = P_{i+1}^- \)? If we use the average, \( \frac{1}{2} (P_i + P_{i+1}) \) the acceleration at \( i+1 \) will be much greater than at \( i \). However, we know that the velocity should be continuous across the discontinuity. Given equal zoning the boundary pressure which gives equal accelerations to points \( i \) and \( i+1 \) is

\[
P_{i+1}^+ = \frac{P_i + P_{i+1}^*}{2}
\]

It can be shown that the resulting finite difference approximation \( P_{i+1}^+ - P_{i+1}^- \) is second order accurate when the density is continuous.

Now consider the problem depicted in 4.2.b. Here we have a heavy material on the left moving into a very light material on the right.
What should we use for $U^+_1 = U^-_{i+1}$? If we use the average, $1/2 \left( U^+_1 + U^-_{i+1} \right)$, there will be a very large rate of compression in region 2 which is incorrect because region 1 is moving into a near vacuum. The quantity that should be continuous is pressure. The velocity which causes equal pressure increases at points $i$ and $i+1$ is

$$U^* = \frac{\left( p c^2 \right)_i u_i + \left( p c^2 \right)_{i+1} U^-_{i+1}}{\left( p c^2 \right)_i + \left( p c^2 \right)_{i+1}}.$$ 

This assumes the sound speed $c$ is a constant. Again it can be shown that the resultant finite difference approximation to $U_x$ is second order accurate if $p c^2$ is continuous.

The midpoint pressure used in Eq. [4.1] are inverse density weighted and the midpoint velocities in Eq. [4.2] are $p c^2$ weighted.

4.4. The artificial viscosity

An artificial viscosity, $q$, is added to the midpoint pressure in Eq. [4.1]. It is quadratic in $f$. Let $U_c$ be the closing rate between points $k$ and $n$, i.e.

$$U_c = (\hat{U}_k - \hat{U}_n) \cdot \frac{(\hat{x}_k - \hat{x}_n)}{\left| \hat{x}_k - \hat{x}_n \right|}$$
Then let \( q_k = a^2 \rho_k U_c^2 \) and \( q_{kn} = a^2 \rho_{kn} U_c^2 \). In the spirit of paragraph 4.3, we inverse density weight the two to get our expression for the midpoint \( q \), i.e.,

\[
q_n = 2a^2 U_c^2 / \left( 1/\rho_k + 1/\rho_{kn} \right)
\]  

[4.3]

In all of our example calculations in section 2 we used \( a^2 = 5.76 \). Now we must fold \( q \) into the internal energy equation in which we need to evaluate \( (P - q) \hat{v} \cdot \hat{U} \). Our approximation for \( \hat{v} \cdot \hat{U} \) is given by Eq. [4.2]. The \( q \) term is brought inside the summation so that

\[
(P + q) \hat{v} \cdot \hat{U} = \sum_n (p_k + q_n) u_n (y_{n-1} - y_{n+1}) + \sum_n (p_k + q_n) v_n (y_{n+1} - x_{n-1})
\]

\[
\sum_n x_{n+1} y_n - y_{n+1} x_n
\]

[4.4]

4.5. **Prevention of density striations**

The method so far described has one remaining difficulty. By having all of the variables centered in space it becomes impossible to detect a sawtooth type wave as depicted in one dimension in Fig. 4.3.
scheme. To correct for this, we define an artificial velocity \( u' \) as depicted in Fig. 4.3. We use our calculated \( \hat{\mathbf{v}} \mathbf{p} \) to extrapolate from point \( k \) to point \( k_n \) giving \( P_{\text{ext}} = P_k + (\hat{\mathbf{x}}_{k_n} - \hat{\mathbf{x}}_k) \cdot \hat{\mathbf{v}} P_k \). If the pressure field is linear then \( P_{\text{ext}} = P_k \). If they are not equal, there is a second derivative in the pressure field which we attempt to reduce. Physically what should happen is a velocity would be produced at the midpoint as indicated in 4.3, which would decompress point \( i+1 \) and compress point \( i-1 \). This velocity must be proportioned to \( \delta P = P_{\text{ext}} - P_k \). We chose to use \( u' = b^2 \delta P / p_c \). We then use \( p_c^2 \) weighting between points \( k \) and \( k_n \) to arrive at

\[
\frac{b^2 \delta P (c_k + c_{k_n})}{p_k c_k + p_{k_n} c_{k_n}^2}
\]

\[\text{[4.4]}\]

\( u' \) is added to \( u_n \) in calculating \( \hat{\mathbf{v}} \mathbf{v} \).

In our present calculations \( b^2 = 1.44 \). We further limit \( |u'_{n}| \) to be less than 20% of the maximum of \( (C_k, C_{k_n}) \). In practice, \( u' \) is a very small term, but an absolutely necessary one. For example, in test problem 1, density striations of around 50% will occur without using \( u' \). We note also that \( \delta P \) is proportional to \( \delta x^2 p_{xx} \) and thus is quadratic in nature. The similarity between \( q \) and \( u' \) is striking. The \( q \) is an artificial pressure which smooths the velocity field while \( u' \) is an artificial velocity which smooths the pressure field.

4.6 Neighbor selection

The method requires a good selection of representative neighbors at each point in time. We have found out that the neighbors whose bisectors form the Voronoi polygon\(^3\) around the point \( k \) are an excellent choice. The \( k_{th} \) Voronoi polygon is defined as that region of space which is nearer point \( k \) than any other point.

V. SUMMARY

The partial differential Eqs (2.1, 2.2, and 2.3), along with the equation of state 2.4, which describe the time evolution of compressible fluid flow can be solved without the use of a Lagrangian mesh. The
Method follows embedded fluid points and uses finite difference approximations to \( \nabla P \) and \( \mathbf{v} \cdot \mathbf{u} \) to update \( p, \mathbf{u} \) and \( e \). We have demonstrated that the method can accurately calculate highly distorted flows without difficulty. The finite difference approximations are not unique, improvements may be found in the near future. The neighbor selection is not unique, but the one being used at present appears to do an excellent job. The method could be directly extended to three dimensions. One drawback to the method is the failure to explicitly conserve mass, momentum and energy. In fact, at any given time, the mass is not defined. We must perform an auxiliary calculation by integrating the density field over space to obtain mass, energy and momentum. However, in all cases where we have done this, we have found the drift in these quantities to be no more than a few percent.

References


3. G. Vornoi, J. Keine Angew, Math., 134, 198 (1904)