Feedback from Freeze-out in Hydrodynamics∗

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Abstract

Most hydrodynamical calculations used in heavy-ion physics ignore the effect of freeze-out matter carrying energy and momentum away from the expanding fluid. In a simple one-dimensional model we compare calculated energy density and velocity profiles, with and without interaction between fluid-like and freeze-out parts of the system, in order to estimate the importance of this effect.

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I. INTRODUCTION

With new data from CERN’s Superconducting Proton Synchrotron (SPS) and with the expected completion of the Relativistic Heavy Ion Collider (RHIC) in Brookhaven in 1999, nuclear physicists will be able to look at higher temperatures in larger space-time volumes than ever before. Of particular interest is the identification of signatures of the transition to the high-energy phase of nuclear matter, commonly referred to as the quark-gluon plasma (QGP). It should be easier to measure the properties of such a transition with heavier ions because of the larger times and volumes occupied by the resulting hot and dense matter. With larger amounts of matter produced, we can hope that a description in terms of smoothly-varying locally equilibrated quantities such as temperature and fluid-dynamical velocity is reasonable, making hydrodynamical models especially valuable in this scenario. The point of view where the physics is described in terms of such average properties rather than in terms of individual particle motions has obvious advantages. The savings in computer simulation time allow research into a larger parameter space and a wider variety of initial conditions can be explored. Particle production calculations are routinely carried out in the framework of such models. However, these models make a drastic approximation when assuming a sudden change from fluid-dynamical behavior to free-streaming (the usual effective definition of “freeze-out”). It should be kept in mind that, since our particle detectors register free-streaming particles far from the interaction zone, a transition in the description from continuous matter variables to free particles is unavoidable if the model is to have any predictive power or relevance to measured data. The underlying question of better interfacing the hydrodynamical and kinetic descriptions at freeze-out is studied by several groups [1–4].

In traditional hydrodynamical models, all the matter in the system of interest is assumed to be a thermalized fluid which expands according to idealized hydrodynamics. Within the fluid lies an isothermal surface which is at the “freeze-out temperature”, $T_{\text{fo}}$. While the freeze-out surface plays no role in influencing the hydrodynamic evolution, one assumes, for purposes of calculating particle spectra, that matter outside the freeze-out surface is no longer in thermal equilibrium, i.e. particle interactions cease, and particles that cross the freeze-out surface are free to move unimpeded to the detectors. If, for example, lepton pairs are measured, the decay of the freeze-out particles can contribute a significant portion of the observed leptons, and their spectra have interesting properties which are sensitive to the hydrodynamic evolution [5].

The trajectory of the freeze-out surface is typically found by assuming that all the fluid is thermalized, even at temperatures below $T_{\text{fo}}$. On the other hand, calculations of particle spectra assume that particles near the freeze-out surface leave the fluid. The energy and momentum carried away by these freeze-out particles can be sizable, and should properly be taken into account in a self-consistent calculation. The fact that some freeze-out particles may return to the fluid through the moving freeze-out surface should also be accounted for. There has been some recent analytical investigation of how to do this by Bugaev [6], who derived equations which describe the trajectory of the freeze-out surface. Here we take the point of view of evolving a system from a given point in time within the context of a computer simulation which includes the feedback from this dynamical freeze out. A Godunov-type calculation [7] appears well-suited for this endeavor. In this paper we show
how to extend the Godunov method to handle the evolution of both thermalized fluid and freeze-out matter, and we compare to calculations of expanding matter assuming all thermalized fluid (traditional hydrodynamics). We demonstrate the effect on a one-dimensional example using the simplest possible relativistic equation of state (EOS) for the thermalized fluid, that of massless pions. These simplifying assumptions can be relaxed in the future. Note that at the highest energies presently in the focus of attention, where a very large number of pions is created, even this most simplified study may be relevant to the interpretation of data.

The paper is organized as follows. In Section II, we describe the essence of Godunov-type methods as typically used today and prepare the way for the inclusion of dynamical freeze-out. The details of how this is done are given in Section III with the technicalities relegated to the Appendix. Section IV summarizes the generalized algorithm implemented by our computer code. Section V contains the results, and we conclude in Section VI.

II. GODUNOV METHOD

One successful approach to solving hydrodynamics on a computer is to use a so-called “Godunov-type” method [6]. In such a method, the space-time continuum is discretized into cells, i.e. the algorithm calculates the changes in the average properties of spatial cells as it makes finite steps in time. Because conservation laws tell us how quantities (such as energy, momentum, and particle densities) in a cell change due to the flows of those quantities through the cell “walls” during a timestep, it only remains to determine what the flows should be to give the correct hydrodynamical behavior.

There are two common approaches within the framework of Godunov-type methods to calculating the flows between thermalized cells, an attempt to exactly calculate the intercell flow shape, and the approach of approximating the flows by uniform regions between cells. Both methods assume that the cells can be treated as semi-infinite and uniform at the beginning of a given timestep. This is reasonable insofar as the timestep is kept small enough that the developing flow patterns do not extend past the middle of the cell, i.e. \( \delta t < \frac{\delta x}{2} \), where \( \delta t \) and \( \delta x \) are, respectively, the timestep size and cell width\(^1\). Once this is assumed, it follows that neither a length nor a time scale exists for the development of a flow pattern between two cells. Therefore a similarity pattern, i.e. a pattern that depends only on the dimensionless variable \( \chi \equiv x/t \), develops. This is also a reasonable approximation in more complex geometries as long as the similarity pattern is much smaller than the size of the system as a whole.

In the most ambitious approach, one attempts to find the actual shape of the flow pattern, e.g. the energy density \( e(\chi) \) and the fluid velocity \( v(\chi) \) across the cell boundary. These “Riemann solutions” can be derived from the continuity equations for the relevant quantities (in this case, energy and momentum) and the equation of state (EOS). In the case of a simple EOS, analytical solutions are possible [7,8]. For a realistic EOS, the shape of the flow pattern is found by numerical integration. The alternative approach proposed by Harten, Lax, and van Leer, and completed by Einfeldt [9], is commonly known as the “HLLE” method [10]. Here the flows between thermalized cells are approximated by uniform regions of length proportional to the signal velocities (sound speed relativistically added to
fluid velocity). HLLE saves considerable CPU time because the task of calculating the actual flow patterns by numerical integration is replaced by choosing approximate signal velocities, which by conservation laws determine the conserved quantities between the two receding discontinuities. Discussion of the options for and choice of the signal velocities can be found in Ref. [10]. In the present work we follow the HLLE algorithm as described there. We emphasize that in any Godunov-type method the calculation of the flows between cells is at the heart of the fluid-dynamical problem.

For our purposes, a Godunov-type method is especially well suited, because the Godunov equations (see eq. (2) below) used to evolve a cell in time are derived directly from conservation laws. Thus the validity of these equations is not restricted to ideal fluids, but they are true in general. Energy-momentum conservation is expressed in terms of the continuity equation for the energy-momentum tensor,

\[ \partial_{\mu} T^{\mu\nu} = 0 . \]  

(1)

The Godunov equations are obtained by averaging the time derivatives over a cell, and for a system that varies in only one spatial dimension, e.g. \( x \), take the form

\[ \frac{d}{dt} \left( \int_{\delta x} dx \, T^{00}(x) \right) = T_{-}^{0x} - T_{+}^{0x} , \]  

(2)

\[ \frac{d}{dt} \left( \int_{\delta x} dx \, T^{0x}(x) \right) = T_{-}^{xx} - T_{+}^{xx} , \]

where \( \delta x \) refers to the spatial extent of the cell, and the subscripts – and + refer respectively to the lower and upper spatial bounds of the cell. The variable \( t \) refers to the time as measured in the frame of the cells, which can be considered to be fixed in space (the “cell frame”), with matter moving between them in the course of a simulation. From inspection of eqs. (1) and (2) it is clear that \( T^{0x} \) represents the flow of \( T^{00} \) and \( T^{xx} \) is the flow of \( T^{0x} \).

The energy-momentum tensor \( T^{\mu\nu} \) is defined, in the rest frame of thermalized fluid, to be

\[ T^{\mu\nu} = \begin{bmatrix} e & 0 & 0 & 0 \\ 0 & p & 0 & 0 \\ 0 & 0 & p & 0 \\ 0 & 0 & 0 & p \end{bmatrix} , \]  

(3)

i.e. \( T^{00} \) is the energy density \( e \), \( T^{0x} \) is the momentum density, and \( T^{xx} \) is the pressure \( p \) (flow of momentum). When the fluid is given a boost \( v \) in the \( x \)-direction, we have

\[ E \equiv T^{00} = (e + p)\gamma^2 - p , \]  

(4)

\[ M \equiv T^{0x} = (e + p)v\gamma^2 , \]

\[ P \equiv T^{xx} = (e + p)v^2\gamma^2 + p , \]

where \( \gamma^2 = 1/(1 - v^2) \), and we have given mnemonic names (reflecting the physical quantities to which these tensor elements reduce in the rest frame) \( E \) (energy density), \( M \) (momentum density), and \( P \) (pressure or flow of momentum density \( M \)) to the three tensor elements of interest. (\( E \) and \( M \) are used in Ref. [10], \( P \) is introduced in this work.) For concise
treatment, we introduce the notation $U_i : E, M, P$ for $i = 1, 2, 3$. We also use $q_i : e, p$ ($i = 1, 2$) for the energy density and pressure in the rest frame of thermalized fluid. The notation $F(U_i)$ for the flow of quantity $U_i$ will also be used occasionally.

Recall that in a non-curved geometry the flows are similarity patterns, and thus the values of $M(\chi = 0)$ and $P(\chi = 0)$ remain constant throughout the timestep. It then follows that

$$\Delta \langle E \rangle = \frac{\delta t}{\delta x} (M_- - M_+),$$

$$\Delta \langle M \rangle = \frac{\delta t}{\delta x} (P_- - P_+),$$

where $\langle T^{\mu\nu} \rangle$ signifies the average value of $T^{\mu\nu}$ in the cell. This is the form that the computer code actually uses to evolve a cell from one timestep to the next. In the examples presented in this work we use a simple ultra-relativistic equation of state, so particle number conservation is ignored and energy-momentum conservation as presented above is sufficient.

**III. GENERALIZED GODUNOV METHOD INCLUDING FREEZE-OUT**

We wish to extend the Godunov method to handle freeze-out as well as thermalized fluid. Strictly speaking, we are not extending the Godunov method per se, but simply using it in a way that exploits its generality. The general problem as we have mapped it out consists of solving for flows in three cases: between two cells of thermalized fluid, between two cells of non-thermalized (freeze-out) matter, and between one cell of each kind. For this last case, the freeze-out surface lies somewhere in one of the two cells, and for simplicity, we approximate the flow with the flow due to thermalized matter (approximation with non-thermalized matter is an equally valid choice). This can be done because we expect the density gradient to be well-behaved in the region connecting thermalized fluid with freeze-out matter, as justified a posteriori by our results (Sect. 5).

The physics of freeze-out can be described in terms of matter that was originally thermalized and treated as a fluid expanding to such a low density that large mean-free paths make fluid behavior an invalid approximation. In fluid-dynamical models of nuclear collisions, such as ours, this is assumed to happen at a well-defined “freeze-out temperature” below which the particles which comprise the matter lose thermal contact. In relativistic nuclear collisions $T_{fo} \approx 100 - 150$ MeV, and the freeze-out particles are expected to reach the detectors without any further interaction. In freeze-out matter it no longer makes sense to calculate flow patterns based on fluid behavior, or even to assume that local quantities such as energy density ($e$) and pressure ($p$) are related by an equation of state. However, since the Godunov equations do nothing more than express the conservation of energy and momentum in a discretized system, they can still be applied, as long as the correct values for the flows, $F(E) = M$ and $F(M) = P$, (and in our case, $F(P)$, the flow of $P$) are used.

It should be pointed out that there is no relativistically covariant continuity equation for $P$, meaning that pressure is not conserved like energy and momentum. However, in weakly interacting matter, such as freeze-out, one can still calculate the contribution to the pressure
due to a particle of given momentum, so one can calculate the "flow of $P$" by considering the rate at which particles cross the cell wall. Thus, in the case of freeze-out matter, we assume $\Delta \langle P \rangle = (\delta t / \delta x) [F(P)_- - F(P)_+]$ (the calculation of $F(P)$ is described in the Appendix), and for thermalized fluid we calculate $P$ from $E$, $M$, and the EOS.

In order to calculate flows of freeze-out matter, it is necessary to keep track of more than $E$ and $M$. This is because for non-thermalized matter, which does not follow an equation of state, $P$ does not have an implicit relationship to $E$ and $M$. In principle, one should keep track of a number of degrees of freedom of the order of the number of particles in the matter. However, including just tensor element $P$ (in addition to the usual $E$ and $M$) in the description captures the essence of freeze-out behavior, which is that the free-streaming particles tend to move at a single velocity, making $P_0$ (the value of $P$ in the zero-momentum frame, where $M=0$, referred to as the "matter rest frame"") smaller than would be expected for thermalized matter with the same $E_0 = e$. Thus our calculation can be thought of as a first order approximation of the coupling effect.

If the potential energy is negligible compared to the kinetic and rest mass energies, e.g. in a non-interacting pion gas, it is natural to express the elements of the energy-momentum tensor in terms of the distributions of particles [11]:

$$T^{\mu \nu} = \int \frac{d^3p}{p^0} p^\mu p^\nu f\left(\frac{p^0}{T}\right),$$

where $f\left(\frac{p^0}{T}\right)$ describes the distribution of the particles. For thermalized particles viewed in the matter rest frame, as defined above, the function $f\left(\frac{p^0}{T}\right)$ is the Bose-Einstein or Fermi-Dirac distribution (Bose-Einstein for the pion gas considered here). The momentum distribution of particles immediately inside the freeze-out surface is known, because we have made the assumption of thermal equilibrium everywhere inside the freeze-out surface. Because we assume a sharp freeze-out surface, the particles crossing the surface will retain the momenta they had while still in the thermalized fluid. We will use (7) for the calculation of the elements of the energy-momentum tensor in non-thermalized matter. The only remaining freedom is in specifying the distribution function, which is described in detail in the Appendix.

We assume massless pions for the freeze-out particles. This is because pions are the lightest, and therefore most abundantly produced hadrons, so our model is closely linked to actual experimental data at the highest beam energies available. Massive particles could be used at a price of some complication of the EOS for thermalized fluid and of the calculation of velocity distributions. Here we limit ourselves to an extremely simple case to illustrate the method, which we expect to find more general applications (e.g. the nonrelativistic scenario of evaporation from a fluid).

IV. STRUCTURE OF THE ALGORITHM

Here we summarize the tasks performed by the generalized Godunov algorithm for the above simplified scenario at each timestep. Before evolving the system in time, the cells are first initialized with values for $v$, the cell frame matter velocities, and $q_i$, the rest frame thermodynamic quantities ($e$ and $p$). After all the cells are initialized, the cell frame tensor
elements $U_i$ ($E$, $M$ and $P$) are calculated. The piecewise-constant distribution is replaced by a piecewise-linear distribution, the slopes being calculated from the values of $U_i$ in neighboring cells. Thus two distinct values of $U_i$ that correspond to two adjacent cells are associated with each boundary (see [10] for details). The difference between these two $U_i$ determines a flow at a given boundary. These flows give rise to second order accuracy in time by allowing the calculation of half-timestep quantities, including the signal velocities and flows used in the HLLE scheme. The new second-order flows determine the evolution of $U_i$ inside each cell (throughout the code, if the $U_i$ are changed, the corresponding $q_i$ and $v$ are updated immediately afterward). Finally, $e$ and $v$ for each cell are output. At this point one time step is completed.

For cells with freeze-out matter, we depart from the standard HLLE scheme, as described in [10]. If a cell is determined to be below the freeze-out temperature (based on the rest-frame energy density $e$), the flow is calculated assuming free-streaming particles. To do this, we first need to parametrize the velocity distribution in terms of the tensor elements $U_i$, as described in the Appendix. To make a calculation with the freeze-out coupling turned off, one simply uses the standard HLLE to calculate all the flows regardless of energy density.

V. RESULTS

We ran most of our simulations with $\delta t=0.03$ fm for the timestep and $\delta x=0.1$ fm for the cell size. The dependence on the timestep was tested, and we found that decreasing $\delta t$ to 0.01 fm did not alter the results appreciably. We use the equation of state of a thermalized massless boson gas (pions). The initial temperature was $T_0 = 200$ MeV for $-2.7 \text{ fm} \leq x \leq 2.7 \text{ fm}$, and vacuum everywhere else. Initial velocities were all set to zero. Because of the symmetry about $x = 0$, only one side of the system ($x \geq 0$) is shown in the figures. The freeze-out temperature was chosen to be $T_{fo} = 140$ MeV, close to the mass of a pion. (This appears to be a realistic estimate of the freeze-out temperature [5].) The corresponding isotherm is called the freeze-out isotherm and defines the freeze-out surface. In a calculation of particle production, particles that cross the freeze-out surface are assumed to lose thermal contact and move freely to the detectors. Therefore, the trajectory of the freeze-out surface is important, and is our primary concern in this work. The results shown were obtained by a second order calculation in time, as described in Section 4. For our simple problem, with a “well-behaved” EOS, we get nearly identical results for first and second order simulations; for equations of state with a phase transition, or other systems where macroscopic discontinuities may develop, a second order treatment is necessary to get accurate results.

Figures 1 and 2 are contour plots of $e$ in $x$-$t$ space for the evolution of a finite one-dimensional slab expanding into the vacuum in both directions; only one side, $x \geq 0$, is shown, without and with feedback from freeze-out, respectively. The isotherms, starting from the right side of the diagram, represent energy densities from 5 to 70 MeV/fm$^3$, in increments of 5 MeV/fm$^3$. The thick line is the $e=49$ MeV/fm$^3$ isotherm, which corresponds to the freeze-out temperature ($T_{fo} = 140$ MeV), or freeze-out surface. We see that it has a distinctly different slope in $x$-$t$ space when we include the coupling between thermalized fluid and freeze-out. The increased inward velocity of the freeze-out surface would result
in more freeze-out particles being measured, because the number is sensitive to the relative
velocity between the matter and the freeze-out surface.

The contour plot for $e$ also demonstrates that there are large regions of small density
gradient in the freeze-out matter in the case of freeze-out coupling. Nearly the entire region
outside the freeze-out surface is at around half the freeze-out energy density. This is in
contrast to the case without freeze-out coupling, which displays a much smoother change of
the energy-density profile. In the case with feedback from freeze-out, there are also much
larger fluctuations in energy density, notably a region of lower density (a “bubble”) that
seems to have formed in the freeze-out matter around $t = 5.0$ fm, but disappears at around $t = 7.0$ fm. Whether this is indicative of new physics or merely a result of the uneven flow
cau sed by the crudeness of our choice of velocity distributions, or the limited number (three)
of moments of the momentum distribution we use, needs further investigation.

Figures 3 and 4, the contour plots for $v$ in $x$-$t$ space, corroborate our expectations and the results displayed in the $e$-plots. In the case of freeze-out coupling (Figure 4), there are large regions outside the freeze-out
surface that have little variation in velocity, indicating that the particles “free-stream” with
no resistance. The contours are, from $x = 0$ outward, $v = 0.01, 0.1, 0.2, \cdots 0.8, 0.9$. The
group of lines in the lower right corner is an artifact of the contour plotting due to the fact
that fast rarefied matter abuts the vacuum, whose velocity is taken to be zero. Most of the
freeze-out matter moves outward at a speed around $0.65c$ in the case when the feedback
from the freeze-out is taken into account. The inhomogeneities that appear seem to be due
to disturbances that originate at or near the freeze-out surface, where particles with a range
of velocities cross into the freeze-out region.
FIG. 1. Lines of constant energy density, for the case without coupling between the freeze-out and thermalized fluid. The thick line ($e=49$ MeV/fm$^3$) is the freeze-out isotherm.

FIG. 2. Lines of constant energy density, for the case with coupling between the freeze-out and thermalized fluid. The thick line ($e=49$ MeV/fm$^3$) is the freeze-out isotherm.
VI. CONCLUSIONS

We have generalized the conventional Godunov algorithm to consistently include freeze-out matter in fluid-dynamical simulations. In particular, the energy and momentum carried away by the freeze-out particles and the feedback from those particles is accounted for in our generalized treatment. In the present paper we illustrated this dynamical freeze-out in a simple one-dimensional example. The net result of the dynamical treatment is a rapid eating into the thermalized fluid as shown by the trajectory of the freeze-out surface, as compared to the treatment without dynamical freeze-out. Also, there is a more sudden change across the freeze-out surface to freeze-out matter that is moving more or less uniformly, as expected.
from free-streaming particles. In addition, the treatment with freeze-out coupling displays more fluctuations, partly due to our simplifying assumptions.

For a more realistic simulation we intend to use the generalized Godunov algorithm in more complicated geometries and with more complicated equations of state. We expect the method to find many applications. One of these may be to combine the improved freeze-out modeling with three-fluid hydrodynamics. Such a combination would be especially useful for calculating production of particles that are sensitive to freeze-out (e.g. dileptons). One suspects that the effect of the freeze-out/fluid coupling on the trajectory of the freeze-out hypersurface will influence pion rapidity distributions at least as much as differences between isothermal and isochronous freeze-out. Photon spectra are also expected to be modified, albeit to a lesser extent.

We believe that fluid-dynamical calculations with an improved treatment of freeze-out will be very competitive with transport-theoretical modeling. This is partly because of great savings in CPU time and partly due to the physics insight afforded by a treatment dealing with average quantities, as opposed to running computer simulations with a very large number of chance events.

VII. APPENDIX

In our approach each cell is characterized by the three tensor elements $E$, $M$ and $P$. These are Lorentz-transformed into the frame where $M$, the momentum density, is zero. This is referred to as the zero-momentum frame, or “matter rest frame”, and we will denote the tensor elements in this frame by $E_0$, $M_0 = 0$ and $P_0$. The velocity of this frame relative to the frame of the cells is the “matter velocity” $v$. In the matter rest frame, the cells are characterized by $E_0$ and $P_0$. For weakly interacting matter in general, $E_0 = e$, the energy density, and $P_0 = p$, the pressure. For the simple (ultra-relativistic) equation of state of our thermalized fluid, $P_0 = E_0/3$. For freeze-out matter there is no relationship between $E_0$ and $P_0$ except for constraints imposed by causality.

For thermalized massless particles, the momentum distribution is isotropic, and $p^r = p^0 \xi$, where we use the notation $-1 \leq \xi \leq 1$ for the dimensionless velocity (in terms of the speed of light) along the $x$-axis, and $p^0$ is the time-like component of the 4-momentum. Therefore we can factor each integral in (7) into an integral over total energy and an integral over velocity in the $x$-direction, e.g.

$$E_0 = \int \frac{d^3p}{p^0} p^0 p^0 f \left( \frac{p^0}{T} \right)$$

$$= \int_{-1}^{1} d\xi \ 2\pi \int_{0}^{\infty} dp^0 \ (p^0)^{3} \ f \left( \frac{p^0}{T} \right)$$

$$\equiv \int_{-1}^{1} d\xi \ h(\xi), \quad (8)$$

where $h(\xi)$, the energy distribution with respect to velocity, is obviously a constant $E_0/2$ due to the fact that the integral over $p^0$ is independent of $\xi$ for thermalized massless particles. (Note that here the subscript on $E_0$ refers to the matter frame, while the superscripts on
p refer to components of the 4-momentum.) Due to this factorization for massless particles we can recast the momentum distribution as a velocity distribution along the x-axis. For other equations of state, \( h(\xi) \) would be defined, but not necessarily a constant. In a similar manner, we obtain:

\[
M_0 = \int_{-1}^{+1} d\xi \, \xi h(\xi) = 0, \quad (9)
\]

\[
P_0 = \int_{-1}^{+1} d\xi \, \xi^2 h(\xi).
\]

For non-thermalized matter, we parametrize \( h(\xi) \) subject to the conditions that it be (1) even with respect to \( \xi = 0 \), which guarantees \( M_0 = 0 \), and (2) non-negative. For cells near thermal equilibrium, i.e. \( P_0 \approx E_0/3 \), we want the simplest function that differs gently from a constant and still satisfies the above criteria; the choice \( h(\xi) = a\xi^2 + b \) (\( a \) and \( b \) constant) works well, where \( a = (45P_0 - 15E_0)/8 \) and \( b = (9E_0 - 15P_0)/8 \).

For much of the freeze-out matter, \( P_0 \) is too small (namely, \( P_0 < E_S \equiv E_0/5 \)) to satisfy \( h(\xi) = a\xi^2 + b \geq 0 \) for all \(-1 \leq \xi \leq +1\). In this case we interpolate between a function of the form \( a\xi^2 + b \) and a delta function, \( (E_0/2)\delta(\xi) \). The use of the delta function reflects our assumption that the particles in freeze-out matter tend to move along together at nearly the same velocity. Occasionally it is necessary, for large \( P_0 \) (> \( E_L \equiv 3E_0/5 \)), to use a double delta function of the form \( h(\xi) = (E_0/2)[\delta(-d) + \delta(d)] \), where \( d \) is a velocity. In this case the matter is described better by a superposition of movement at a couple velocities than by homogeneous particle movement.

To calculate the flows of matter over the right-hand (left-hand) cell boundary, we calculate the integrals (8) and (9) for particles with cell-frame velocities \( v(\xi) \) greater than (less than) zero, where \( v(\xi) = (\xi + v)/(1 + \xi v) \). For example, the flow of energy \( F(E) \) (due only to matter in the cell being considered) over the right-hand side of the cell is

\[
F(E) = \int_{-v}^{+v} d\xi \frac{dM_0}{d\xi} + \int_{-v}^{+v} d\xi \frac{dP_0}{d\xi} = \int_{-v}^{+v} d\xi \gamma^2 \frac{d}{d\xi} \left( vE_0 + (v^2 + 1)M_0 + vP_0 \right),
\]

where \( M_0' \) is the momentum density, viewed in the cell frame, due to particles with velocity between 0 and 1. The derivatives of the rest frame quantities are simply

\[
\frac{dE_0}{d\xi} = h(\xi),
\]

\[
\frac{dM_0}{d\xi} = \xi h(\xi),
\]

\[
\frac{dP_0}{d\xi} = \xi^2 h(\xi).
\]

The evaluation of the flow of \( P \) requires the numerical integration of

\[
F(P) = \int_{-v}^{+v} d\xi \, v(\xi) \frac{dP}{d\xi} = \int_{-v}^{+v} d\xi \, v(\xi) \gamma^2 \frac{d}{d\xi} \left( v^2 E_0 + 2vM_0 + P_0 \right).
\]
where \( \gamma^2 = 1/(1 - v^2) \) is the gamma factor due to the relative velocity \( v \) between the rest frame and the cell frame.

Flows from two neighboring cells are added together to determine the total flow across the shared interface. These flows are then used by the code to update \( U_i \) at the next full timestep.

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**NOTES**

1. We use the convention \( \hbar = c = 1 \).
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