Simulation of two- and three-dimensional dense-fluid shear flows via nonequilibrium molecular dynamics: Comparison of time-and-space-averaged shear stresses from homogeneous Doll’s and Sllod shear algorithms with those from boundary-driven shear

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Homogeneous shear flows (with constant strainrate \( dv_x/dy \)) are generated with the Doll’s and Sllod algorithms and compared to corresponding inhomogeneous boundary-driven flows. We use one-, two-, and three-dimensional smooth-particle weight functions for computing instantaneous spatial averages. The nonlinear normal-stress differences are small, but significant, in both two and three space dimensions. In homogeneous systems the sign and magnitude of the shearplane stress difference, \( P_{xx} - P_{yy} \), depend on both the thermostat type and the chosen shearflow algorithm. The Doll’s and Sllod algorithms predict opposite signs for this normal-stress difference, with the Sllod approach definitely wrong, but somewhat closer to the boundary-driven truth. Neither of the homogeneous shear algorithms predicts the correct ordering of the kinetic temperatures: \( T_{xx} > T_{zz} > T_{yy} \).

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I. INTRODUCTION

In the present work, we use nonequilibrium molecular dynamics [1] to study microscopic simulations of “simple shear flow” (also called “plane Couette flow”),

\[
v_x \approx y \rightarrow P_{xy} = P_{yx} = -\eta \left( \partial v_x / \partial y \right) = - \eta \left( \partial v_y / \partial x \right) = - \eta \dot{\epsilon}.
\]

The flow is in the \( x \) direction so that the tensor \( \nabla v \) has only one nonzero element, \( \left( \nabla v \right)_y = dv_y / dy = \dot{\epsilon} \). We use the symbol \( P \) for the (symmetric second-rank Cauchy) pressure tensor (positive in compression and the negative of the stress tensor, \( P = -\sigma \)); \( v \) for the (time- and space-dependent) hydrodynamic flow velocity; \( \eta \) for the shear viscosity; and \( \dot{\epsilon} \) for the magnitude of the imposed (or measured) shear strain rate. The various simulation types we consider here were designed to clarify the relationships between periodic homogeneous methods, thermostated everywhere, and flows with moving thermostated boundaries. All of the methods we use are consistent with the Green and Kubo linear response theory at small rates of shear [2]. We are specially interested in characterizing and understanding the nonlinear shear plane stress difference: \( P_{xx} - P_{yy} = \sigma_{yy} - \sigma_{xx} \) which arises in sufficiently small systems at sufficiently large shear rates. In carrying out microscopic simulations both the boundary conditions and the thermostats or ergostats which control the flow need to be carefully considered [1].

Though simple shear flow is “stationary,” fluctuations in local properties necessitate averaging, both in time and in space. Here we reduce the importance of these fluctuations by using spatial averaging techniques borrowed from smooth-particle continuum simulation methods [3]. We measure instantaneous spatially averaged flow velocity, temperature, and pressure-tensor components.

The two best-known homogeneous microscopic methods, the Doll’s tensor [4] and Sllod algorithms [5], treat fluid rotation differently, leading to qualitatively different predictions for the nonlinear normal-stress difference. Boundary-driven flows can help to resolve this disagreement [6–9]. Useful flows need to satisfy four conditions: The spatial scale \( L \) of these flows needs to be large enough (relative to the particle size), but not too large (to avoid turbulence), with flow velocities \( v \) large enough (to emerge above fluctuations), but not too large (again, to avoid turbulence), in order to provide useful information. The relative greater importance of fluctuations in two dimensions is responsible for the reduced utility of viscosity there, as shown in Fig. 1. Additionally, the frictional boundary-fluid interaction needs to be sufficiently strong to prevent excessive boundary slip.

In the present work, we carry out both two- and three-dimensional simulations using all three approaches (Doll’s, Sllod, and boundary-driven) for two simple, and rather similar, pairwise-additive repulsive potentials. We focus here on

FIG. 1. Limits imposed on system size \( L \) and boundary velocity \( v \) by atomistic size, thermal fluctuations, turbulence, and shock waves. Viscosity is a useful concept for flows in the enclosed areas. The figure here is constructed for a fluid at unit mass, number density, temperature, viscosity, and heat conductivity. \( L \) is measured in units of the microscopic particle size and \( v \) is measured in units of the thermal velocity. Modeled after Ref. [6].
the normal-stress difference, $P_{xx} - P_{yy} = \sigma_x - \sigma_y$, in the shear plane. We will see that the Doll’s and Silod algorithms typically predict different signs for the simple monatomic dense fluids considered here. Both these homogeneous algorithms yield clearcut results, insensitive to system size.

In the early days of such shear-flow simulations [10] it was thought that the two-dimensional viscosity might depend logarithmically on system size [11]. More recently, careful studies [12–15] of the corresponding Green-Kubo stress-strain correlation function

$$\eta = (V/kT) \int_0^\infty \langle P_{xy}(0)P_{xy}(t) \rangle_{eq} dt,$$

indicate no such dependence in dense fluids. The work we carry out here is consistent with this lack of size dependence. Here we find that the coefficient of the hypothetical logarithmic viscosity contribution can be no larger than $10^{-4}$ in the natural reduced units of atomic size, mass, and velocity. The more realistic boundary-driven flows necessarily entail larger fluctuations and considerable size dependence. We are, nevertheless, able to determine the sign and the size of the normal-stress difference for such flows so as to characterize the errors inherent in the homogeneous algorithms.

This paper is organized as follows: in Sec. II we lay out the macroscopic description of the problem; in Sec. III the microscopic description; In Sec. IV we describe the two homogeneous algorithms emphasizing their similarities and differences; in Sec. V we describe the boundary-driven algorithm used in the present work; Sec. VI describes the smooth-particle spatial averaging method used in analyzing results from simulations; Sec. VII describes numerical results for two similar (smooth repulsive) force-law models in two very different density regimes and in two space dimensions; Sec. VIII describes corresponding three-dimensional results for one of these force-law models; Sec. IX lists the conclusions we have drawn from this work.

II. MACROSCOPIC DESCRIPTION OF SIMPLE SHEAR FLOW

Classical fluid flow can be modeled and understood from either the microscopic or the macroscopic standpoint. In the microscopic description individual particles obey the ordinary differential equations of motion of nonequilibrium molecular dynamics [1]:

$$\{m\ddot{r}_i = m\dot{v}_i = F_A + F_B + F_C + F_D\},$$

and everything follows from the functional forms of the assumed atomistic, boundary, constraint, and driving forces [1,16]. In molecular dynamics, just as in continuum hydrodynamics, the pressure tensor is (defined to be) the comoving momentum flux, comoving relative to the flow velocity. In a many-body system with forces derived from the pairwise-additive pair potential $\phi(r)$, the pressure tensor is made up of $\{i,j\}$ pair contributions as well as individual particle convective contributions [1]:

$$P_{xy} = \sum_{i<j} \left[ \frac{x_{ij} y_{ij}}{r_{ij}^2} \right] [F \cdot r]_{ij} + \sum_i (p_i p_j / m)_i,$$

$$x_{ij} = x_i - x_j, \quad y_{ij} = y_i - y_j, \quad r_{ij} = |r_i - r_j|,$$

$$F_{ij} = -\nabla_i \phi(r_{ij}).$$

It is evident that the pair-potential pressure tensor is symmetric, with $P_{xy}$ and $P_{yx}$ equal.

In the macroscopic description the continuum field variables [such as the mass density $\rho(r,t)$, the velocity $\mathbf{v}(r,t)$, and the energy per unit mass $e(r,t)$] obey partial differential equations

$$\rho = -\rho \nabla \cdot \mathbf{v},$$
$$\rho \ddot{\mathbf{v}} = -\nabla_e \mathbf{P} = \nabla \cdot \mathbf{\sigma},$$
$$\rho \dot{e} = -\mathbf{Q} \cdot \nabla \mathbf{v}.$$

In these general continuum field equations the constitutive relations for the stress tensor $\mathbf{\sigma} = -\mathbf{P}$ and the heat flux vector $\mathbf{Q}$ distinguish one material from another. In both approaches a computational algorithm for solving the equations is needed. Its implementation gives $\{\mathbf{v}(t),\mathbf{v}(t)\}$ in the microscopic case and $\{\rho(r,t),\mathbf{v}(r,t),e(r,t)\}$ in the macroscopic case. In either case, a well-posed problem also requires boundary and initial conditions, constraints, and driving forces.

First consider the simplest model system illustrating stationary simple shear: Imagine an incompressible Newtonian fluid, with constant shear viscosity $\eta$, and which also follows Fourier’s linear heat transport law with a constant heat conductivity $\kappa$:

$$P_{xy} = -\eta (dv_x / dy) + (dv_y / dx),$$
$$\rho \dot{e} = \kappa \nabla^2 T - P \nabla \mathbf{v}.$$

We ignore thermal expansion, so that the mass density $\rho$ is constant. We denote the local thermodynamic variables in the conventional way: Temperature $T(r)$, pressure tensor $\mathbf{P}(r)$, and internal energy per unit mass $e(r)$. We adopt the colon convention in the tensor product $A : B$ to indicate a sum over all four $A_{ij}B_{ij}$ terms in two dimensions, and all nine such terms in three dimensions. In some texts the alternative sum (immaterial for the symmetric tensors considered here) $A_{ij}B_{ji}$ is used.

Simple shear flow for this bare bones textbook model is perhaps the simplest imaginable nonequilibrium flow problem. It gives a linear variation of velocity in space along with a quadratic variation of temperature. Simple shear flow can be driven by two moving parallel boundaries, both of them at temperature $T_B$, and able to absorb heat and to impose their boundary velocities, $v_z = \pm v$, on a two-dimensional strip or three-dimensional slab of model fluid of thickness $L$,

$$v_z(y = \pm L/2) = \dot{e} y.$$

The stationary macroscopic description of such a flow, for the model Newtonian fluid with Fourier heat conduction, has a constant stress tensor, a linear velocity profile and (because $\nabla^2 T$ is constant) a quadratic temperature profile,

$$(dv_x / dy) = \dot{e} = 2\nu / L = -P_{xy}/\eta.$$
The maximum temperature difference, relative to the boundaries’ temperature $T_B$,

$$\Delta T(y) = \Delta T_{\text{max}} = \Delta T(0) = \eta \bar{v}^2 L^2/8 \kappa = \eta \bar{u}^2/2 \kappa,$$

occurs at the midplane $y=0$. See Fig. 2 for a schematic illustration of this prototypical simple shear flow.

This stationary solution satisfies energy balance, with the rate at which heat is generated throughout the volume $V$ (necessarily the same as the rate at which external work is done, $-W$), equal to the rate at which heat is transferred through the two boundary walls of area (length in two dimensions) $A$, at $y = \pm L/2$,

$$-W = -P_{xy} V \dot{\epsilon} = \eta V \dot{\epsilon}^2 = \kappa A \left[ (dT/dy)_{-L/2} - (dT/dy)_{L/2} \right].$$

This macroscopic description of shear flow guides our interpretation of microscopic simulations. We focus on the complications caused by fluctuations and nonlinearities in what follows.

III. MICROSCOPIC DESCRIPTION OF SIMPLE SHEAR FLOW

Molecular dynamics was developed over 50 years ago [17–19] and soon gave rise to successful interpretations of equilibrium properties based on hard-sphere perturbation-theory [20] analogous to Enskog’s hard-sphere understanding of dense-fluid transport properties [21]. Field-driven diffusion [22], shear and bulk viscous flows [23], and heat conducting flows [24,25] all came to be simulated with a variety of algorithms. Special boundary conditions and computational thermostats were developed to model these flows. Throughout this development, the limiting case of the Green and Kubo linear theory of transport served as a guide [26].

In the past 30 years simulation has come a long way. Billions of atoms can be simulated now [27]. Nonequilibrium simple shear flows have been developed for diatomic and polyatomic molecules [28,29], not just simple fluids. More recently, alternative shear flow algorithms have been developed for periodic irrotational “elongational” flows [30–32], flows with a steady stretching in one direction and a simultaneous shrinking in a perpendicular direction. It might be thought (as it once was [33]) that such a flow could not be followed forever, but a clever choice of periodic boundaries makes it possible to study steady thermostated elongational flows. The conceptual difficulties involved in deriving these algorithms have led to a spirited literature [34–36] as to the “correctness” of the various algorithms. Such discussions can easily lead outsiders to the impression that it is hard to distinguish “correct” from “incorrect” algorithms.

The controversial aspects of these shear flow algorithms [28–36] led us to reconsider the problem. It seemed to us that a fresh look at the basic algorithms for monatomic simple-shear flows would help to develop a perspective clarifying this situation. It is evident that the nonlinear aspects of the computer algorithms are to some extent arbitrary, as the only true guidelines for correctness are consistency with the well-known and well-accepted linear flow theory described by Newtonian viscosity and Fourier heat conduction.

In viscous shear flow, which we consider here, any reasonable algorithm needs to satisfy the requirement that the shear viscosity for small strain rates agrees with the Green and Kubo linear response relation [2,26] linking the viscosity to equilibrium fluctuations in the shear stress $\sigma_{xy} = -P_{xy}$,

$$\eta = \langle V/kT \rangle \int_0^\infty \langle P_{xy}(0) P_{xy}(t) \rangle_{eq} dt,$$

$$\eta = -\frac{P_{xy}}{\dot{\epsilon}}, \quad \dot{\epsilon} = \frac{\partial \bar{v}}{\partial y} + \frac{\partial \bar{v}}{\partial x}.$$

For simplicity we consider systems with pairwise-additive forces $\{F_{ij}\}$ derived from a potential function $\Phi = \Sigma \phi$,

$$\{F_{ij} = -\nabla_i \phi(|r_i - r_j|) = -F_{ji}\}.$$

In such a system both the energy $E$,

$$E = \Phi + K = \sum_{i<j} \phi_{ij} + \sum_i p_i^2/2m,$$

and the microscopic pressure tensor $P$,

$$PV = \sum_{i<j} \langle F r_{ij} \rangle + \sum_i \langle pp/m \rangle,$$

are sums of two-particle potential and single-particle kinetic contributions.

There are two interesting ambiguities in the definition of pressure in a particulate system. The contributions of the potential pair interaction terms $\{F_{ij}\}$ to $PV$ need to be allocated spatially. Either $\delta$-function contributions at $r_i$ and $r_j$, or at $(r_i + r_j)/2$, or smoothed distributions centered at these locations can be used. The old “Irving-Kirkwood” preference for $\delta$ functions is motivated more by analytic convenience than by any physical considerations. A smoothed approach is certainly preferable in computational work.

The kinetic part of $PV$ is ambiguous too. How is the “co-moving” velocity to be defined in a system with transient velocity fluctuations? Either an instantaneous or a time-averaged hydrodynamic velocity can be chosen. Only the kinetic part of $P_{xy} V$ is affected by this choice. The simplest versions of the two choices are as follows:

$$P_{xy}(t)V = \sum_i m [v'_i - v_i(t)] [v'_i - v_i(t)] \text{ (instantaneous)}$$

or
forces schemes, all of which are based on time-reversible constraint needs to be removed. This can be done with any one of many finite spatial range, and emphasizes the contributions of the instantaneous kinetic temperature for

The kinetic part of the pressure tensor is the usual definition in three space dimensions the usual definitions of individual particle velocities.

In a system composed of moving particles, is to use a

must be estimated numerically. The simplest way to do this must be estimated numerically. The simplest way to do this in a system composed of moving particles, is to use a weighted (smoothed-particle) average of nearby particle velocities, where the time-independent scalar weight function $w_{ir}$ has a finite spatial range, and emphasizes the contributions of those nearby particles to the location $r$ where $v(t)$ is to be computed. The details of this spatial average are described in Sec. VII. Fortunately, the main conclusions of this work are independent of the necessarily arbitrary choice between using instantaneous and time-averaged flow velocities. By choosing to study stationary simple shear we avoid the additional complexities associated with time averaging flows which have an explicit time dependence in their boundary conditions.

The atomistic simulation approach is also complicated by heat and by fluctuations. Because viscosity is dissipative, a viscous shear flow necessarily generates heat. Particle motions and collisions in systems tractable with molecular dynamics necessarily exhibit fluctuations in all their local properties [38]. Despite the time reversibility of the underlying equations of motion, the overall long-time-averaged character of these flows is necessarily dissipative, in accord with macroscopic hydrodynamics and the second law of thermodynamics [38–41].

To simulate a steady viscous flow, the heat generated needs to be removed. This can be done with any one of many schemes, all of which are based on time-reversible constraint forces [1,16]. Such constraint forces can play the role of a feedback-based thermostat or ergostat,

\[ \{F_{\text{Constraint}} = -\zeta p\} \rightarrow \dot{K} = 0 \quad \text{or} \quad \dot{E} = 0. \]

Typical constraint-force choices keep the kinetic energy $K$ or the total energy $E$ fixed, or allow these energies to fluctuate about a specified mean value in a way consistent with Gibbs’ statistical mechanics at equilibrium [42,43]. For simplicity, we restrict ourselves here to “Gaussian thermostats” (so named after Gauss’ principle of least constraint) [44], which fix the kinetic energy $K$ or the total energy $E$ of a particular set of degrees of freedom by imposing feedback-based constraint forces, \( \{F_{\text{Constraint}} = -\zeta p\} \). The Gaussian “friction coefficient” $\zeta$ constrains the momenta ($p$) contributing to $K$ or $E$.

The instantaneous hydrodynamic velocity \( [v_x(t), v_y(t)] \) must be estimated numerically. The simplest way to do this in a system composed of moving particles, is to use a weighted (smoothed-particle) average of nearby particle velocities,

\[
v(t) = \sum_i v_{ir} w_{ir} \bigg/ \sum_i w_{ir}, \quad w_{ir} = w(|r - r_i|),
\]

where $i$ is the index of all the particles, \( v_{ir} \) is the velocity of the particle at position $r$ in the direction of $r_i$.

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We next describe the two best-known computational algorithms for simulating simple shear.

IV. DOLL’S AND SLLOD ALGORITHMS FOR SIMPLE SHEAR FLOW

Shear flows were the first application of homogeneous nonequilibrium molecular dynamics [6]. Steady inhomogeneous shock waves [45] and boundary-driven shear flows [6–9] had both been simulated earlier. Special boundary conditions, or external forces, as well as thermostats had to be developed for the homogeneous shear flows. The numerical applications of these ideas, in the early 1970s, preceded their formal theoretical development by decades. The Doll’s tensor Hamiltonian for viscous flows was discovered in 1985 [4]. Dettmann and Morriss’ Hamiltonians (for the Nosé-Hoover and Gauss thermostats) were discovered in 1996 [46,47]; and a proper formulation of elongational flows first appeared in 1998 [30].

Simple homogeneous shear flow, with the $x$ velocity proportional to the $y$ coordinate,

\[
\dot{\epsilon} = d\nu_x / dy \leftrightarrow \nabla \nu = \begin{bmatrix} \partial \nu_x / \partial x & \partial \nu_x / \partial y \\ \partial \nu_y / \partial x & \partial \nu_y / \partial y \end{bmatrix} = \begin{bmatrix} 0 & 0 \\ \epsilon & 0 \end{bmatrix},
\]

can be implemented with periodic “Lees-Edwards” boundary conditions [48] [developed independently by Ashurst [7]]. See page 26 of Ref. [6] for a brief description of Ashurst’s algorithm. The corresponding flow is illustrated in Fig. 3.

This simple shear flow, with $\nu_x$ nonzero, is “rotational,” in the sense that the clockwise rotation rate, $-\omega$, is nonzero, and equal to one-half the strain rate,
In our specific special case $\nabla v$ can be written as a sum of irrotational and rotational contributions

$$\nabla v = \begin{bmatrix} 0 & 0 \\ 0 & 0 \end{bmatrix} + \begin{bmatrix} 0 & +\varepsilon/2 \\ +\varepsilon/2 & 0 \end{bmatrix} = \begin{bmatrix} 0 & +\varepsilon/2 \\ +\varepsilon/2 & 0 \end{bmatrix}.$$ 

See Fig. 3 for an illustration of the periodic boundary conditions consistent with such a flow. A stationary nonequilibrium state can result if the irreversible heating rate, $-W = \eta \dot{\varepsilon}\dot{V}$ is appropriately compensated by momentum-dependent thermostat forces.

The Doll’s and Sllod algorithms are straightforward possibilities for simulating such a flow, and the differences between them are small (of order $\dot{\varepsilon}^2$) at moderate strain rates. Ashurst’s periodic shear algorithm preceded the formal derivations of differential evolution equations for the shear flow and its associated thermostats. His finite-difference algorithm was equivalent, for small time steps, to a thermostated version of what is now called the Sllod algorithm:

$$\{ \dot{x} = (p_x/m) + \dot{\varepsilon}y; \dot{y} = (p_y/m); \dot{p}_x = F_x - \dot{\varepsilon}p_y - \dot{\xi}p_x; \dot{p}_y = F_y - \dot{\xi}p_x \}.$$

In the absence of the thermostat forces $\{ \dot{\xi} \}$, these motion equations give an energy change exactly consistent with the rate at which thermodynamic work is performed by the instantaneous shear stress $-F_{xy}$.

In the alternative Doll’s-tensor approach [4], a coordinate-momentum $(q,p)$ sum (giving rise to the “Kewpie-Doll” name) is added to the usual Hamiltonian. The added term includes the strain rate $\dot{\varepsilon}$.

$$\mathcal{H}_{Doll} = \mathcal{H}_{Usual} + \varepsilon \sum_i (yp_i).$$

The resulting Hamiltonian equations of motion,

$$\left\{ \begin{array}{l} \dot{q} = + \frac{\partial \mathcal{H}}{\partial p} \quad \dot{p} = - \frac{\partial \mathcal{H}}{\partial q} \end{array} \right\},$$

are only slightly different to the closely related “Sllod” equations. The Doll’s-tensor motion equations are

$$\{ \dot{x} = (p_x/m) + \dot{\varepsilon}y; \dot{y} = (p_y/m); \dot{p}_x = F_x - \dot{\xi}p_y - \dot{\xi}p_x; \dot{p}_y = F_y - \dot{\xi}p_x \}.$$

In the absence of heat-extracting thermostats, both approaches, Sllod and Doll’s, provide an energy change exactly consistent with thermodynamics

$$\mathcal{H}_{Doll}(q,p,\nabla v) = 0 \rightarrow \mathcal{H}_{Usual}(q,p) = - (d/dt) \sum q \dot{p} \nabla v = -\varepsilon P_{xy} V.$$

The extra momentum-dependent forces, $\{ \dot{\xi}p_x \}$ for Sllod and $\{ \dot{\xi}p_y \}$ for Doll’s, model rotation. In a rotational flow in the $xy$ plane the rotation rate $\omega$ is given by

$$\omega = \frac{1}{2} \left( \frac{\partial v_y}{\partial x} - \frac{\partial v_x}{\partial y} \right),$$

and corotating momenta would include rotational contributions

$$\dot{p}_x = -\omega p_y, \quad \dot{p}_y = +\omega p_x.$$
FIG. 4. Four-chamber $L \times 4L$ periodic flow with $4 \times 400$ particles. Chamber 1 (filled circles at the bottom) moves to the right at speed $+\varepsilon L/2$; chamber 3 (filled circles, third chamber from the bottom) moves to the left at speed $-\varepsilon L/2$. This imposes the nominal strain rates $\pm \varepsilon$ on the two Newtonian chambers 2 and 4 (open circles). Periodic boundaries apply in both the $x$ and the $y$ directions. Note the ordering effect of the moving tethers in chambers 1 and 3.

The multichamber shear flows are induced by tethering the boundary particles to steadily moving lattice sites of all the space dimensions. Normal-stress differences from the two-chamber flows are shown in Figs. 10 and 11. They are not not significantly different from their four-chamber analogs.

VI. SMOOTH-PARTICLE SPATIAL AVERAGES

The spatial averaging algorithms our simulations require are borrowed from a continuum technique, “SPAM” [3]. Smooth particle applied mechanics (“SPAM”) is a technique for solving the partial differential continuum equations of continuity, motion, and energy for the evolution of the density, velocity, and energy. It makes use of a normalized weight function with a maximum range $h$, $w(|r| < h)$. The weight function describes the spatial extent of a representative particle of mass $m$. The weight function is formulated with at least two continuous derivatives everywhere in order that the first and second continuum derivatives of smooth particle sums [corresponding to instantaneous local quantities such as $\nabla \rho(r,t)$, $\nabla u(r,t)$, and $\nabla^2 f(r,t)$], are everywhere continuous in both space and time. This continuity and differentiability facilitates a smooth transition between particle and continuum analyses.

Consider the two simplest cases, mass and momentum sums. The density $\rho(r)$, and the momentum density $\rho(r)u(r)$ at the location $r$ are defined by summing up the contributions of all particles ($j$) within the maximum range $h$ of that location,

$$\rho(r) = m \sum_j w(|r - r_j|) = m \sum w_{rj},$$

$$\rho(r)u(r) = m \sum_j v_j w(|r - r_j|) = m \sum w_{rj}v_j.$$  

An advantage of this formulation is that the continuum continuity equation,

$$\frac{\partial \rho}{\partial t} = -\nabla \cdot (\rho v),$$

is satisfied exactly by the interpolated smooth particle fields,

$$\frac{\partial \rho(r)}{\partial t} = \sum_j \left( \frac{\partial w_{rj}}{\partial t} (r - r_j) \right)_{r_j}$$

$$= m \sum_j \left( \nabla w_{rj} \cdot \frac{\partial r - r_j}{\partial t} \right)_{r_j}$$

$$= + m \sum_j (\nabla w_{rj}) \cdot v_j$$

$$= - m \nabla \cdot \sum_j w_{rj}v_j = - \nabla \cdot (\rho v).$$
TABLE I. Relatively high-density Lucy viscosity in two dimensions. The mass and number densities are unity: \(\rho = m N / V = N / V = 1\). The range of the Lucy potential is \(h = 3\), so that each particle interacts simultaneously with approximately 30 neighbors. Space-time-averaged pressure tensors are given here for homogeneous SiIod and Doll’s algorithm with a square periodic \(20 \times 20 = 400\)-particle cell with a strain rate of \(\dot{\varepsilon} = d\varepsilon / d t = 0.05\). The kinetic temperature is fixed, \(kT = 0.07\), using a single control variable \(\varepsilon\) (in the two runs S1 and D1) and two separate control variables in the \(x\) and \(y\) directions, \(\xi\) and \(\eta\) (in the two runs S2 and D2). The total energy is fixed at 0.500 for the runs SE and DE. The pressure-tensor components are given in the order \(x, y, x\) with the kinetic, potential, and total terms indicated. The boundary conditions are periodic, with a total run time of 200 \(\times 10000\) time steps. The fourth-order Runge-Kutta time step is 0.005. The average potential energies per particle for the first four runs are all in the range 0.4272 < \(\Phi / N < 0.4274\) so that the total energy in these runs is 0.497.

<table>
<thead>
<tr>
<th>Run type</th>
<th>(p_{xx}^k)</th>
<th>(p_{xx}^b)</th>
<th>(p_{xx}^e)</th>
<th>(p_{xy}^k)</th>
<th>(p_{xy}^b)</th>
<th>(p_{xy}^e)</th>
<th>(p_{xy}^e)</th>
<th>(p_{xy}^b)</th>
<th>(p_{xy}^e)</th>
</tr>
</thead>
<tbody>
<tr>
<td>400S1</td>
<td>0.078</td>
<td>0.484</td>
<td>0.562</td>
<td>0.062</td>
<td>0.485</td>
<td>0.547</td>
<td>-0.022</td>
<td>+0.002</td>
<td>-0.021</td>
</tr>
<tr>
<td>400D1</td>
<td>0.060</td>
<td>0.486</td>
<td>0.546</td>
<td>0.079</td>
<td>0.483</td>
<td>0.563</td>
<td>-0.021</td>
<td>+0.001</td>
<td>-0.020</td>
</tr>
<tr>
<td>400S2</td>
<td>0.070</td>
<td>0.485</td>
<td>0.555</td>
<td>0.070</td>
<td>0.484</td>
<td>0.554</td>
<td>-0.025</td>
<td>+0.002</td>
<td>-0.023</td>
</tr>
<tr>
<td>400D2</td>
<td>0.070</td>
<td>0.485</td>
<td>0.555</td>
<td>0.070</td>
<td>0.484</td>
<td>0.554</td>
<td>-0.025</td>
<td>+0.002</td>
<td>-0.023</td>
</tr>
<tr>
<td>400SE</td>
<td>0.081</td>
<td>0.484</td>
<td>0.564</td>
<td>0.064</td>
<td>0.485</td>
<td>0.549</td>
<td>-0.023</td>
<td>+0.002</td>
<td>-0.022</td>
</tr>
<tr>
<td>400DE</td>
<td>0.062</td>
<td>0.486</td>
<td>0.548</td>
<td>0.083</td>
<td>0.483</td>
<td>0.566</td>
<td>-0.022</td>
<td>+0.001</td>
<td>-0.021</td>
</tr>
</tbody>
</table>

Notice that the spatial gradient operator, \(\nabla\), affects only \(w(r-r_j)\) and not the individual point-particle properties \(\{v_i\}\).

By choosing properly normalized weight functions spatial averages can be computed in one, two, or three spatial dimensions. The pressure tensor in the vicinity of a particle or at a grid point in three dimensions would be computed by using the three-dimensional weight function,

\[
w_{3D}(r) = \frac{105/16\pi h^3}{4\pi r^2} \left[1 - 6(r/h)^2 + 8(r/h)^3 - 3(r/h)^4\right] \int_0^h 4\pi r^2 w_{3D}(r) dr = 1.
\]

This particular weight function was discovered and used by Lucy in 1977 [53]. It is the simplest normalized polynomial with (1) a maximum at \(r=0\); (2) a finite range \(r < h\), and (3) two continuous derivatives vanishing at \(r=h\). The corresponding one- and two-dimensional weight functions (for averages in thin strips or slabs, and for averages at particles or grid points in two-dimensional problems, are

\[
w_{2D}(r) = \frac{5/\pi h^2}{2\pi r} \left[1 - 6(r/h)^2 + 8(r/h)^3 - 3(r/h)^4\right] \int_0^h 2\pi r w_{2D}(r) dr = 1.
\]

\[
w_{1D}(r) = \frac{5/4h}{2\pi} \left[1 - 6(r/h)^2 + 8(r/h)^3 - 3(r/h)^4\right] \int_0^h 2w_{1D}(r) dr = 1.
\]

The smooth-particle equations of motion for the time development of the individual particle velocities \(\{v_i\}\) are generally formulated so as to conserve linear momentum exactly. The failure of this approach to conserve angular momentum is a relatively subtle point worthy of more research investigation [54].

VII. NUMERICAL RESULTS: TWO DIMENSIONS

A. Homogeneous simulation results with Lucy’s potential

To simplify the numerical work, reduce numerical integration errors, and to make contact with existing data [55–57], we initially chose to use Lucy’s pair potential in two space dimensions. This pairwise-additive potential is identical to the weight function \(w_{2D}(r)\) of Sec. VI,

\[
\phi(r) = \frac{5}{\pi h^2} \left[1 - 6x^2 + 8x^3 - 3x^4\right], \quad x = \frac{r}{h} < 1.
\]

We can make a rough estimate of the potential contribution to the energy and pressure for this potential by assuming a random distribution of particles. Viewed as a statistical-mechanical potential function for molecular dynamics in two dimensions, Lucy’s potential (due to the normalization of the Lucy weight function) then corresponds to the simple equation of state for a two-dimensional ideal gas,

\[
\Phi = \sum_{i<j} \phi_{ij} = \frac{N^2}{2V} \leftrightarrow p^V = \frac{N^2}{2V}.
\]

The energy expression follows from the normalization condition, while the pressure expression follows from the virial theorem:

\[
p^V = \frac{1}{2} \sum_i \sum_{j \neq i} r_{ij} f_{ij} = -\frac{N^2}{4V} \int_0^h 2\pi r^2 \phi'(r) dr = \frac{N^2}{2V}.
\]

At unit mass and number density and at a strain rate of \(\dot{\varepsilon} = 0.05\) the shear viscosity and normal stresses (for SiIod) were measured precisely in 1995 [55, 56]. Because the results are insensitive to the number of particles used, we list in Table I below only a single representative set of simulations for \(N = 400\). The insensitivity of the viscosity to the algorithm type is also evident in Daivis recent work [58].

We chose \(h = 3\), for which the SiIod algorithm shear viscosity (with a single thermostat variable) has previously been computed. At unit density, with \(L = L = N\), the average number of pair interactions is approximately 14N. We also used exactly the same Lucy’s function as a weight function [3] for
carrying out spatial averages. Table I lists the pressure-tensor components using both the Doll’s and Sllod algorithms. The first four simulations are “isothermal” with both the average temperature, \((T_x + T_y)/2\) thermostated (S1 and D1), and with both temperatures separately thermostated (S2 and D2), by using two control variables, \(\zeta_x\) and \(\zeta_y\). The last two simulations (SE and DE) constrain the internal energy rather than the temperature. Notice that in all these simulations the average pressure is fairly close to the simple virial-theorem estimate, \(P^0 V/N = 0.5\).

The normal-stress difference is sensitive to the single-thermostat type

\[
P_{xx} - P_{yy} = +0.015, \quad P_{xx} - P_{yy} = -0.017, \quad \text{(Doll’s) s},
\]

and is almost entirely kinetic. At the high effective density of these simulations the potential portion of the stress distribution is nearly isotropic. These results are essentially unchanged if energy, rather than average temperature, is controlled. On the other hand, this striking normal-stress effect disappears completely (because it is kinetic in nature) if the two temperatures are separately thermostated. We conclude from these simulations that a definitive determination of the normal-stress difference requires a more realistic boundary-driven flow. One can have no confidence in nonlinear kinetic effects when the evolving kinetic energy itself can be dominated by the choice of homogeneous thermostats, one or two.

**B. Boundary-driven simulation results with Lucy’s potential**

Because the homogeneous investigations with the Lucy potential were sensitive to thermostat type, we next carried out several four-chamber simulations with two boundary chambers moving oppositely, as shown in Fig. 4. The density and strain rate were chosen to match the data in Table I. These simulations produced no useful normal-stress results. This failure reflects the nearly negligible coupling between the two moving chambers and the two Newtonian chambers. See the typical velocity profile in Fig. 5. Particles in the two Newtonian chambers simply rest quietly between the two rapidly moving walls. Evidently, this very dense repulsive fluid with relatively weak collisional forces has insufficient friction for boundary driving to reach strain rates with significant nonlinear stress differences. This same difficulty was found by Liem, Brown, and Clarke in their three-dimensional Lennard-Jones simulations [9].

On the other hand, in Ashurst’s thesis work his “fluid-wall” boundary driving regions (velocity constraints, but no tethers, were used) produced good linear velocity profiles for both Lennard-Jones and soft-sphere potentials [6,7]. Because those simulations correspond to a much lower density (with about three interactions per Lucy particle rather than 30) and much more violent collisions than those of Table I, we abandoned the high-density Lucy simulations and took up instead soft-disk and soft-sphere simulations at conditions more closely resembling those of Ashurst. The force law change was motivated by the desire to check our results with those from previous simulations [57]. The new simulations are discussed in the following two sections.

**C. Homogeneous simulation results with a soft-disk potential**

Here we consider the short-ranged smooth soft-disk repulsive potential [57],

\[
\phi(r < 1) = 100(1-r^2)^4, \quad r^2 = x^2 + y^2,
\]

very similar in general shape to Lucy’s, but with three vanishing derivatives rather than just two, at its maximum range of unity. The two potentials are compared in Fig. 6. In comparison simulations we found that the two potentials provide quite similar normal stresses at corresponding temperatures and densities. Here we choose unit density and energy per particle. We will see that boundary-driven simulations with this potential choice provide good velocity profiles, as did Ashurst’s similar “fluid-walls” in 1972, and also allow comparisons with previous viscosity simulations in the same thermodynamic state. First we consider again Sllod and Doll’s simulations with homogeneous shear.

Just as in the Lucy simulations, the soft-disk normal stresses are quite different for the Doll’s and Sllod algorithms and are even less sensitive to system size. Now the short-ranged potential contribution to shear stress is actually dominant. The results listed in Table II, all using a single friction coefficient, fixing the energy per particle, \(E/N=1\), show that now the potential contribution to the stress difference is the same order as the kinetic contribution. The Sllod

---

FIG. 5. Comparison of Lucy’s potential and the short-ranged soft-sphere repulsive potential \(\phi\). The ordinate and the abscissa are divided by their maximum values, so that both scales vary from 0 to 1.
TABLE II. Moderate-density soft-disk viscosities. The range of the potential is unity so that each particle interacts, on the average, with only three neighbors at unit density. Space-and-time-averaged pressure tensors for homogeneous Slod and Doll’s algorithms using the soft-disk pair potential illustrated in Fig. 6,  \( \phi = 100(1-r^2)^2 \). The energy and density are fixed, and equal to unity, and the strain rate, \( \dot{e} = dv_x/dy \) is 0.50. The pressure-tensor components, are again given in the order \( xx, yy, xy \) with the kinetic, potential, and total terms indicated. The boundary conditions are periodic, with a total run time of \( 200 \times 10^6 \) time steps for \( N = 64, 256, 1024; 200 \times 2000 \) time steps for \( N = 4096 \), and \( 200 \times 500 \) time steps for \( N = 16384 \). The fourth-order Runge-Kutta time step is 0.005 and a single isoenergetic friction coefficient is used, as explained in Sec. II of the text.

<table>
<thead>
<tr>
<th>N type</th>
<th>( \rho_{xx}^k )</th>
<th>( \rho_{yy}^k )</th>
<th>( \rho_{xy}^k )</th>
<th>( \rho_{xx}^\phi )</th>
<th>( \rho_{yy}^\phi )</th>
<th>( \rho_{xy}^\phi )</th>
<th>( \rho_{xx}^\Sigma )</th>
<th>( \rho_{yy}^\Sigma )</th>
<th>( \rho_{xy}^\Sigma )</th>
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</thead>
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<td>3.217</td>
<td>3.907</td>
<td>0.681</td>
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<td>3.904</td>
<td>-0.081</td>
<td>-0.534</td>
<td>-0.615</td>
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<td>0.692</td>
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<td>-0.617</td>
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</tr>
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<td>-0.623</td>
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<tr>
<td>4096D</td>
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<td>3.193</td>
<td>3.872</td>
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<tr>
<td>16384S</td>
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<td>3.211</td>
<td>3.903</td>
<td>0.681</td>
<td>3.214</td>
<td>3.895</td>
<td>-0.085</td>
<td>-0.537</td>
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</tr>
<tr>
<td>16384D</td>
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<td>3.872</td>
<td>0.694</td>
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<td>3.926</td>
<td>-0.085</td>
<td>-0.537</td>
<td>-0.622</td>
</tr>
</tbody>
</table>

pressure difference is considerably smaller in magnitude,
\[ P_{xx} - P_{yy} = +0.008 \quad \text{(Slod)}, \]
\[ P_{xx} - P_{yy} = -0.054 \quad \text{(Doll’s)}, \]
Because the rotated normal-stress difference is equivalent to a shear stress,
\[ \frac{(P_{xx} - P_{yy})^{45}}{2} = P_{xy}, \]
an alternative phenomenological description of the normal-stress effect corresponds to a rotation of the principal stress direction (the direction of maximum tension in the traceless stress tensor). The Doll’s algorithm gives a clockwise shear-stress rotation while the Slod rotation is counterclockwise.

The Slod program used to generate the data in Table II reproduced earlier Slod results [57] for strain rates of 0.10 and 0.25 very well. Gass’ Enskog-theory prediction [59] of the (linear) viscosity for the conditions of Table II, \( \eta_{\text{Enskog}} = 1.5 \), is just slightly higher than the value \( \eta = 1.25 \) given by the data in this table. Just as in the work of Refs. [10–12], there is no indication of any logarithmic \( N \) dependence in these results. Let us now compare these homogeneous results to those from boundary-driven simulations.

D. Boundary-driven results with the soft-disk potential

With the soft-disk potential \( \phi(r < 1) = 100(1 - r^2)^2 \) a useful boundary-driven flow does result if the strain rate is moderate and the system is not too large. The boundary-driven flows are sensitive to system size. This is because the heat generated in a shear flow, of order \( L^0 \) for \( L \) large eventually overwhelms the capacity of the boundary, of order \( L^{D-1} \), to absorb it. The maximum strain rate that can be reached by boundary-driven flows is therefore limited by the heat conductivity as well as the efficiency of the frictional thermal contact at the reservoir walls. In addition to varying the size and stiffness of the tethering potential, we also varied the lattice structure of the tether sites, but settled on the simple square and cubic lattices when the pressure-tensor results proved to be insensitive to lattice type.

Systems with Newtonian strips 20 atoms wide were already large enough that no systematic deviation from linear-stress behavior (with vanishing normal-stress difference, \( \sigma_{xx} = \sigma_{yy} = 0 \)), resulted. Accordingly, we use a narrower system width here, 10, for an analysis of boundary driven flows. The system size is \( N = 4 \times (10 \times 10) = 400 \) particles.

Time-averaged boundary-driven velocity and normal-stress-profiles are shown in the figures. The time-reversible frictional forces within the two steadily moving “boundary” chambers were chosen to maintain a kinetic temperature in those chambers (relative to the tether velocity) of 0.70. This approximately reproduces the conditions of the Slod and Doll’s states in Table II. Figure 7 shows the average velocity \( \langle v_x(y) \rangle \), computed using the one-dimensional weight function of Sec. VII. For each of 400 equally spaced grid points the instantaneous values of the laboratory-frame velocity components \( \{ v_i \} \) were spatially averaged at the grid points

\[ v_i(y_G, t) = \frac{\sum_i w_{1D}[|y_G - y_i(t)|]}{\sum_i w_{1D}[|y_G - y_i(t)|]} \cdot \]

These instantaneous grid-point averages were then themselves averaged over time and are plotted in Fig. 7.

Pressure-tensor components at each particle were calculated in two different ways. Using the assumed analytic form, a linear velocity profile,
\[ 0 < y < 10 \rightarrow \langle v_x \rangle_{\text{analytic}} = +5, \]
\[ 10 < y < 20 \rightarrow \langle v_x \rangle_{\text{analytic}} = 15 - y, \]
\[ 20 < y < 30 \rightarrow \langle v_x \rangle_{\text{analytic}} = -5, \]
the pressure tensors for particles in the vicinity of each grid point were computed as averages, using the two-dimensional weight function $w_{2D}(r<h=3)$. These individual pressure-tensor components were then averaged, with $w_{1D}(r<h=3)$ as were the velocities of Fig. 7. Pressure-tensor components were also computed and averaged using the smooth-particle weights $w_{2D}(r<3)$ to calculate the instantaneous flow velocity $\langle v(r_j) \rangle$ at each particle $j$ rather than using the assumed linear profile. The differences are relatively small, as can be seen in Fig. 8, where the two approaches are compared. Using the instantaneous velocity at each particle is analogous to, but smoother than, the “unbiased” procedure discussed by Evans and Morriss [16].

The normal-stress differences following either approach are considerably larger than the Sllod results (Table II), and have the opposite sign to the Doll’s homogeneous results (Table II).

$$P_{xx} - P_{yy} = 0.03 \quad \text{(boundary driven).}$$

The Sllod algorithm predicts a smaller effect (smaller by an order of magnitude) while the Doll’s algorithm predicts the wrong sign. We conclude that the two homogeneous algorithms provide no more than an order of magnitude estimate of the normal-stress effects and further that these effects can be otherwise measured reliably, but with some difficulty.

Figure 8 shows stresses for the two portions of the four-chamber system with 13 < $y$ < 17 and 33 < $y$ < 37. These portions have “typical” bulk fluid averages, without any influence from the two driving boundary regions. This is a consequence of the smooth-particle weight functions’ range, $h = 3$.

![Figure 7](image)

**FIG. 7.** Velocity profile for a 400-particle system with the aspect ratio illustrated in Fig. 4. The tethering potential’s force constant, $k = 100$ provides an efficient coupling between the driving chambers at $0 < y < 10$ and $20 < y < 30$ and the driven Newtonian chambers at $10 < y < 20$ and $30 < y < 40$. The locations of the two Newtonian chambers are emphasized in the velocity plot. The averaged velocity profile was calculated with the smooth-particle weighting function $w_{1D}(r<3)$. The measured strain rate in the Newtonian regions is about $±0.48$.

$$30 < y < 40 \rightarrow \langle v_y \rangle_{\text{analytic}} = y - 35,$$

where the pressure tensors for particles in the vicinity of each grid point were computed as averages, using the two-dimensional weight function $w_{2D}(r<h=3)$. These individual pressure-tensor components were then averaged, with $w_{1D}(r<h=3)$ as were the velocities of Fig. 7. Pressure-tensor components were also computed and averaged using the smooth-particle weights $w_{2D}(r<3)$ to calculate the instantaneous flow velocity $\langle v(r_j) \rangle$ at each particle $j$ rather than using the assumed linear profile. The differences are relatively small, as can be seen in Fig. 8, where the two approaches are compared. Using the instantaneous velocity at each particle is analogous to, but smoother than, the “unbiased” procedure discussed by Evans and Morriss [16].

The normal-stress differences following either approach are considerably larger than the Sllod results (Table II), and have the opposite sign to the Doll’s homogeneous results (Table II).

$$\frac{P_{xx} - P_{yy}}{2} = 0.03 \quad \text{(boundary driven).}$$

The Sllod algorithm predicts a smaller effect (smaller by an order of magnitude) while the Doll’s algorithm predicts the wrong sign. We conclude that the two homogeneous algorithms provide no more than an order of magnitude estimate of the normal-stress effects and further that these effects can be otherwise measured reliably, but with some difficulty.

Figure 8 shows stresses for the two portions of the four-chamber system with 13 < $y$ < 17 and 33 < $y$ < 37. These portions have “typical” bulk fluid averages, without any influence from the two driving boundary regions. This is a consequence of the smooth-particle weight functions’ range, $h = 3$.

![Figure 8](image)

**FIG. 8.** Boundary-driven two-dimensional flow using the “soft-disk” potential. Average normal-stress differences are shown. At the left the stresses are calculated relative to the instantaneous velocity profile. There, the spatially averaged instantaneous velocity and stress at each particle are computed with $w_{2D}$, then averaged to get instantaneous profiles using $w_{1D}$, and finally time averaged. At the right the stresses are calculated relative to an assumed linear velocity profile. In both cases the stress difference is shown for the same regions 13 < $y$ < 17 and 33 < $y$ < 37 free of boundary influences and hence typical of bulk fluid. The run length was 5000.

**VIII. NUMERICAL RESULTS: THREE DIMENSIONS**

**A. Periodic shear**

Three-dimensional boundary-driven simulations require only the addition of $z$ coordinates, with periodic boundary conditions in the $z$ direction. For comparison purposes, we first generated series of isoenergetic Sllod and Doll’s periodic shears. These results, shown in Table III, are for periodic shearing of $L\times L\times L$ cubes of soft-sphere fluid at unit density and energy

$$\phi = 100(1 - r^2)^4, \quad r^2 = x^2 + y^2 + z^2, \quad N/V = Nm/V = E = K + \Phi = 1.$$ 

The constant-energy ergostat forces keep the total energy of the $N=L\times L\times L$ particles fixed. The kinetic energy $K$ is a sum in which each particle’s contribution is measured relative to the local velocity. Choosing the cube center as the coordinate origin, the systematic velocity in the $x$ direction is taken to be proportional to $y$,

$$\{p^2/2m = (p_x^2 + p_y^2 + p_z^2)/2m; (p_z/m) = v_x - \bar{v}_y\}.$$ 

The pressure-tensor results, given in Table III, are very insensitive to system size. Notice that the average kinetic temperature for all these isoenergetic simulations is approximately 0.5. The Sllod algorithm gives

$$T_{xx} > T_{yy} > T_{zz} \quad \text{(Sllod)},$$

while the Doll’s tensor algorithm gives instead

$$T_{yy} > T_{xx} > T_{zz} \quad \text{(Doll’s)}.$$ 

The (correct) “Boundary-driven” results, described next, show instead the ordering

$$T_{xx} > T_{zz} > T_{yy} \quad \text{(boundary driven)}.$$ 

The boundary-driven results also show qualitative differences from the Sllod and Doll’s results in the normal-stress differences ($P_{xx} - P_{yy}$) and ($P_{xx} - P_{zz}$).
TABLE III. Soft-sphere viscosities in three dimensions with periodic boundary conditions. Space-and-time-averaged pressure tensors for homogeneous Sllod and Doll’s algorithms using the soft-sphere pair potential illustrated in Fig. 6, \( \phi = 100(1 - r^2)^4 \). The energy and density are equal to unity and the strain rate, \( \epsilon = \frac{dv_y}{dy} \), is 0.50. The pressure-tensor components, are given in the order \( xx, yy, zz, xy \) with the kinetic, potential, and total terms indicated. The boundary conditions are periodic, with a total run time of \( 200 \times 10000 \) time steps for \( N=10 \times 10 \times 10=1000 \) with a fourth-order Runge-Kutta time step of 0.005.

<table>
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<tr>
<th>(N)</th>
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<th>(p_{xx}^p)</th>
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<th>(p_{xy}^p)</th>
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<th>(p_{yy}^p)</th>
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<td>-0.344</td>
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</table>

B. Boundary-driven shear

We next implemented 4000=4 \times (10 \times 10 \times 10)-particle boundary-driven shear flows similar to the two-dimensional flows of Sec. VII B, but with periodic boundaries in the \( z \) direction and with a fixed isothermal boundary temperature 0.5, chosen to match the homogeneous periodic results. For comparison with these Sllod and Doll’s results the same nominal strain rate, \( dv_y/dy=0.5 \), was used. Thus, the two thermostated chambers move with velocities \( (v_z, \pm 2.5) \).

The time and spatially averaged velocity profile computed with the weight function \( w_{1D}(|\delta y|<3) \) is shown in Fig. 9. The measured strain rates in the straight-line portions of the profile are about \( \pm 0.47 \). The energy dissipation rate for 6000 thermostated degrees of freedom in the two moving 1000-particle reservoirs was 6000\(kT\left \langle \frac{\delta y}{\delta t} \right \rangle \approx 6000 \times 0.5 \times 0.112=336 \), giving an estimate for the viscosity \( \eta = TS_{\text{external}}/(V\epsilon^2) = 336/(2000 \times 0.5^2) = 0.672 \), within 2\% of the periodic result, 0.688 from Table III. The Newtonian shear stresses from the 4000=4 \times (10 \times 10 \times 10)-particle simulation are \( \pm 0.37 \) in the bulk Newtonian regions, corresponding to

\[
\eta = -P_{yy}/(dv_y/dy) = 0.74.
\]

The time-averaged spatially smoothed normal-stress differences,

\[
(P_{xx} - P_{yy})/2 = 0.01, \quad (P_{xx} - P_{zz})/2 = 0.01,
\]

are only different with marginal significance, and are shown in Figs. 10 and 11. We considered three different system sizes, to ensure that these results are insensitive to small geometrical changes. Data for \( N=4 \times 10^3 \) (run length 1000), \( N=4 \times 10^3 \) (two runs of length 500; results from only one of them are shown here as the difference between the two was insignificant), and \( N=4 \times 14^3 \) (run length 400) are included.

![FIG. 9. Time-averaged velocity profile for a 4000-particle three-dimensional system with the aspect ratio illustrated in Fig. 4. The tethering potential’s force constant, \( \kappa = 100 \), provides an efficient coupling between the driving chambers at 0 < \( y < 10 \) and 20 < \( y < 30 \) and the driven Newtonian chambers at 10 < \( y < 20 \) and 30 < \( y < 40 \). The locations of the two Newtonian chambers are emphasized in the velocity plot. The averaged velocity profile was calculated with the smooth-particle weighting function \( w_{1D}(r < 3) \). The measured strain rate in the Newtonian regions is about \( \pm 0.47 \). The run length was 1000.](image1)

![FIG. 10. Normal-stress differences for boundary-driven three-dimensional flows using the “soft-sphere” potential. The normal-stress difference \( (P_{xx} - P_{yy})V/2N \) is shown here for three different system sizes, with stress calculated relative to the instantaneous velocity profile. The spatially-averaged instantaneous particle values of velocity and stress are computed with \( w_{1D} \). Then, the particle values are averaged to get instantaneous profiles using \( w_{1D} \). The figure shows time averages of those instantaneous profiles. Only data from the two distinct regions free of boundary averaging influences are shown here. The large filled circles are from three analogous two-chamber flows with 800 < \( t < 4000 \).](image2)
in the figures. Just as before, the Newtonian regions for which the data are plotted are those free of any boundary influences in the stress averaging. The corresponding Sllod and Doll’s values for the stress differences, +0.003 and −0.016, respectively, are quite different, just as in two dimensions. The disparity shows that neither homogeneous algorithm is even close to “correct.” The actual difference between $P_{yy}$ and $P_{zz}$ is apparently quite small, while both the Doll’s and the Sllod algorithms indicate a relatively large difference of order ±0.04. The statistical fluctuations in the boundary-driven simulations are not quite so large as to mask the ordering of the two normal-stress differences,

$$P_{xx} - P_{yy} > P_{xx} - P_{zz}.$$ 

Somewhat faster (larger) computers could make this conclusion more convincing.

As the system size is increased, with the strain rate fixed, the Newtonian temperature increases also, in rough accord

with the linear model treatment of Sec. II. That is, the central temperature increase is proportional to $L^2$. Figure 12 shows temperature profiles for the three system sizes considered here. We consider the kinetic temperature here, because of its relative conceptual simplicity and its physical conceptual basis [37]. In the boundary-driven shear flows the ordering of the kinetic temperatures is

$$T_{xx} > T_{zz} > T_{yy},$$

with the difference between $T_{xx}$ and $T_{zz}$ being 2 or 3 times larger than that between $T_{zz}$ and $T_{yy}$.

IX. SUMMARY, CONCLUSIONS, AND REMARKS

We were able to characterize the tensor temperature and the nonlinear stresses for both homogeneous and boundary-driven versions of simple shear. Neither Sllod nor Doll’s gives the correct ordering of the kinetic temperatures $T_{ii}$. Generally, the “Sllod” algorithm gives a somewhat “better” approximation to the normal-stress differences $\{P_{ij}-P_{ji}\}$, though Sllod is certainly far from “correct.” Despite its evident failures, there is a fairly widespread faith in the Sllod approach [60]. It is clear (as emphasized to us by Lutsko; see also Ref. [61]) that the algorithms’ extra rotational terms in the motion equations, $-\epsilon p_{i}$ for Sllod and $-\epsilon p_{i}$ for Doll’s, when left to their own devices, would eventually cause $p_{zz}^2$ and $T_{zz}$ to diverge for the Sllod algorithm, and $p_{zz}^2$ and $T_{yy}$ to diverge for Doll’s. This provides a clear explanation of the qualitative difference between the two algorithms’ predictions and the corresponding opposite directions for the rotation of the principal axis of the stress.

Lutsko [62] has reviewed the hard-sphere-based Enskog theory for nonlinear stress (“the only viable theory”) and his finding that $P_{xx} > P_{yy}$ in simple shear is quite consistent with our results. On the other hand, some theoretical models [63] and some computer simulations [16] find $P_{xx} < P_{yy}$, even for relatively simple fluids, so it is clear that more investigations are required. Evidently, the temperature tensor and the nonlinear stresses are not given accurately by the Sllod algorithm. The more realistic boundary-driven flows need to be used whenever confidence in the results is required.

We wish to address here the concern that our thermostated-boundary flows might not be “realistic” or “typical.” Our work adopts and applies the usual assumption that dense fluids can be described by constitutive relations when the fluid is sufficiently far from any boundaries—this is analogous to Saint Venant’s assumption in elasticity. Here the constitutive relation gives the normal stresses in terms of the local stream velocity. We obtained very similar results with the linear model treatment of Sec. II. That is, the central temperature increase is proportional to $L^2$. Figure 12 shows temperature profiles for the three system sizes considered here. We consider the kinetic temperature here, because of its relative conceptual simplicity and its physical conceptual basis [37]. In the boundary-driven shear flows the ordering of the kinetic temperatures is

$$T_{xx} > T_{zz} > T_{yy},$$

with the difference between $T_{xx}$ and $T_{zz}$ being 2 or 3 times larger than that between $T_{zz}$ and $T_{yy}$.

FIG. 11. Normal-stress differences for boundary-driven three-dimensional flows using the “soft-sphere” potential. The normal-stress difference $(P_{xx}-P_{zz})V/2N$ is shown here for three different system sizes, with stress calculated relative to the instantaneous velocity profile. The spatially-averaged instantaneous particle values of velocity and stress are computed with $w_{3D}$. Then the particle values are averaged to get instantaneous profiles using $w_{1D}$. The figure shows time averages of those instantaneous profiles. Only data from the two distinct regions free of boundary averaging influences are shown here. The large filled circles are from three analogous two-chamber flows with $800 < t < 4000$.

FIG. 12. Tensor temperatures for boundary-driven three-dimensional flows using the “soft-sphere” potential. The time-averaged kinetic temperatures, relative to the instantaneous velocity profile, computed with $w_{3D}$ and $w_{1D}$, are shown here for three system sizes, all with boundary temperatures and strain rates equal to 0.5. The systems contain four chambers with $10^6$ (run length 1000), $12^3$ (run length 500), and $14^3$ (run length 400) particles per chamber in the three cases shown. Although data from the two distinct regions free of boundary averaging influences are shown here the differences between $T_{xx}$, $T_{yy}$, and $T_{zz}$ are of order ±0.02 and are too small to see on the scale of the figure.
pressure) indicate that the normal stresses can indeed be described by a constitutive relation so that the exact details of the boundaries are not crucial.

Boundary-driven flows are actually extremely complex, even for this simplest possible model of shear. The flows we can study are dominated by fluctuations which can be tamed by averaging, in one, two, or three dimensions, but the time-averaged flows describe the time-dependent situation no better than they would for a physical waterfall or a turbulent stream.

It is fortunate that a hydrodynamic description of flows is feasible on a very small scale (just a few particle diameters), as was apparent from the earliest shock-wave simulations, which showed shock widths of only a few particle diameters. It still remains a puzzle that shock waves indicate an enhanced nonlinear viscosity while the homogeneous shear algorithms considered here predict a reduction rather than an enhancement [45].

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