Efficient mesoscale hydrodynamics: multiparticle collision dynamics with massively parallel GPU acceleration

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Abstract

We present an efficient open-source implementation of the multiparticle collision dynamics (MPCD) algorithm that scales to run on hundreds of graphics processing units (GPUs). We especially focus on optimizations for modern GPU architectures and communication patterns between multiple GPUs. We show that a mixed-precision computing model can improve performance compared to a fully double-precision model while still providing good numerical accuracy. We report weak and strong scaling benchmarks of a reference MPCD solvent and a benchmark of a polymer solution with research-relevant interactions and system size. Our MPCD software enables simulations of mesoscale hydrodynamics at length and time scales that would be otherwise challenging or impossible to access.

Keywords: multiparticle collision dynamics, mesoscale hydrodynamics, molecular dynamics, hybrid simulations, GPU, MPI

1. Introduction

Complex fluids and soft matter, including colloidal suspensions, polymer solutions, and biological materials, are characterized by structure on disparate length scales, dynamics that occur over a range of time scales, and interactions on the scale of thermal energy \cite{1,2}. This combination of properties can give rise to many complex behaviors, including non-Newtonian rheology \cite{3,4,5} and the ability to self-assemble into organized structures \cite{6,7}. Computer simulations have emerged as useful tools for studying complex fluids and soft matter because they offer detailed, simultaneous resolution of structure and dynamics, and can be used to predict and systematically engineer a material’s properties.

Many complex fluids and soft materials consist of a mesoscopic solute, such as polymer chains or spherical colloids, dispersed in a molecular solvent. The solute can be orders of magnitude larger than the solvent and have corresponding slower dynamics \cite{8}. For example, colloidal particles typically have diameters ranging from a few nanometers to micrometers, but are dispersed in solvents such as water that have molecular

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diameters less than a nanometer. Classical molecular dynamics (MD) simulations \[9,10\] are often unsuitable for such problems because explicitly resolving the solvent with the solute quickly becomes intractable, both in terms of the number of particles required for the model and the number of simulation time steps that are necessary to study the time scales of interest. On the other hand, continuum constitutive models employed in computational fluid dynamics \[11,12\] neglect details of fluctuating microscopic structures, which may be important for accurately describing complex liquids such as polymer solutions when the rheological properties are influenced by the local structure. A simulation approach bridging these regimes to retain a detailed model of the solute while resolving only the essential effects of the solvent is necessary. Various mesoscale methods have been developed for this purpose, including lattice-based models such as the family of Lattice-Boltzmann methods \[13,15\]; implicit-solvent models including Brownian dynamics \[9\], Stokesian dynamics \[16\], and fast lubrication dynamics \[17\]; and particle-based methods such as dissipative particle dynamics \[18,20\], direct simulation Monte Carlo \[21,22\], and multiparticle collision dynamics (MPCD) \[23,24\]. The reader is referred to \[25\] and the references therein for an excellent discussion on the various advantages and disadvantages of these methods, particularly as applied to colloidal suspensions. This article focuses on the MPCD method, first developed by Malevanets and Kapral \[23\], and its massively parallel implementation for graphics processing units (GPUs).

In MPCD, the solvent is represented as a set of off-lattice point particles that undergo ballistic streaming steps followed by cell-based, stochastic multiparticle collisions \[23,24,26\]. The nature and frequency of the collisions determines the transport coefficients of the solvent \[27,29\]. The MPCD algorithm is unconditionally numerically stable, has an H-theorem, and naturally includes the effects of thermal fluctuations \[23\]. Polymers \[30,31\], colloids \[25,32,34\], and fluid-solid boundaries \[35,37\] can be coupled to the solvent, making MPCD an excellent tool for studying soft matter \[24,48,45\]. The MPCD equations of motion are less computationally demanding to integrate than those in MD because MPCD particles do not usually have interactions with each other (e.g., dispersion forces), which typically require some of the most time-consuming calculations in the MD algorithm \[9,10\]. On the other hand, MPCD can require a high particle density in order to achieve a sufficiently liquid-like solvent \[29\], which still poses significant demands on the memory required to represent the particle configuration and the number of calculations required to propagate the solvent particles.

MPCD is particularly well-suited for parallelization because of its particle-based nature and spatially localized collisions. Yet, to our knowledge, there are few publicly available massively parallel implementations of the MPCD algorithm. RMPCDMD \[46\] is a multithreaded program for performing MPCD simulations, but lacks support for decomposing the simulation onto multiple CPUs, limiting the simulation sizes that can be reasonably accessed. The SRD package \[47\] of the massively parallel MD code LAMMPS \[48\] implements a limited set of MPCD features and supports spatial decomposition onto multiple CPU processes using the Message-Passing Interface (MPI). Recently, Westphal et al. \[49\] described an implementation of the
MPCD algorithm for GPUs. GPUs have become indispensable for performing MD simulations at large scale because of their massively parallel architectures, and essentially all major MD software packages support GPU acceleration to some extent. Westphal et al. demonstrated a 1-2 order of magnitude acceleration of their MPCD code on a GPU compared to single-threaded CPU code [49]. However, their implementation, which to our knowledge is not publicly available, was designed to use a single GPU as an accelerator, and so the GPU memory capacity effectively limits the size of the simulations that can be performed. It is critical to be able to utilize multiple GPUs through MPI in order to efficiently perform large-scale MPCD simulations.

In this article, we present a high-performance, open-source implementation of the MPCD algorithm within the HOOMD-blue simulation package [50, 51] that scales to run on hundreds of NVIDIA GPUs. We first give an overview of the MPCD algorithm (Section 2). We then describe the implementation of this algorithm with a focus on optimization for the GPU and for supporting multiple GPUs with MPI (Section 3). We assess the accuracy and performance of a mixed-precision computing model for the MPCD solvent. We report weak and strong scaling benchmarks of a reference MPCD solvent and a benchmark of a polymer solution (Section 4). We demonstrate that massively parallel GPU acceleration can enable simulations of mesoscale hydrodynamics at length and time scales that would be otherwise challenging or impossible to access.

2. Algorithm

In MPCD [23, 24, 26], the solvent is modeled by point particles of mass $m$ with continuous positions $\mathbf{r}_i$ and velocities $\mathbf{v}_i$. MPCD particles are propagated through alternating streaming and collision steps. During the streaming step, the particles follow Newton’s equations of motion. In the absence of any external forces, the positions are simply updated during an interval of time $\Delta t$ according to:

$$\mathbf{r}_i(t + \Delta t) = \mathbf{r}_i(t) + \mathbf{v}_i(t) \Delta t.$$  \hspace{1cm} (1)

The time step $\Delta t$ effectively sets the mean free path for the particles, $\lambda = \Delta t \sqrt{k_B T/m}$, where $k_B$ is Boltzmann’s constant and $T$ is the temperature. Particles are then binned into cubic cells of edge length $a$, which sets the length scale over which hydrodynamics are resolved [52]. For values of $\lambda$ much smaller than $a$, the MPCD algorithm violates Galilean invariance, which can be restored by applying a random shift to the particle positions of $\pm a/2$ before binning [53, 54]. All particles in a cell undergo a stochastic collision that updates their velocities while conserving linear momentum. (Collision rules can be extended to also enforce angular-momentum conservation [24, 55].) These stochastic collisions lead to a build up of hydrodynamic interactions, and the choice of collision rule and solvent properties determine the transport coefficients of the fluid.
In the stochastic rotation dynamics (SRD) collision rule [23], the center-of-mass velocity of each cell, $\mathbf{u}$, is first computed and the relative particle velocities are rotated by a fixed angle $\alpha$ about a randomly chosen axis for each cell,

$$
\mathbf{v}_i(t + \Delta t) = \mathbf{u}(t) + R(\alpha) \cdot (\mathbf{v}_i(t) - \mathbf{u}(t)),
$$

(2)

where $R(\alpha)$ is the rotation matrix for the cell. Malevanets and Kapral showed that the SRD rule yields the hydrodynamic equations for compressible flow, has an H-theorem, and gives the correct velocity distribution for the particles [23]. Although SRD in this form conserves linear momentum, eq. 2 requires modification in order to also enforce angular-momentum conservation [56, 57].

Because SRD is an energy-conserving collision rule, a cell-level thermostat is required to enforce isothermal conditions in nonequilibrium simulations. Huang et al. [58] recently explored several thermostat schemes and found the Maxwell-Boltzmann thermostat [59] to be most effective. Here, the current kinetic energy of each cell, $E_k$, in the reference frame of the cell center-of-mass velocity $\mathbf{u}$ is first determined,

$$
E_k = \frac{1}{2} \sum_{i=1}^{N_c} m (\mathbf{v}_i - \mathbf{u})^2,
$$

(3)

where the sum is taken over the $N_c$ particles in the cell. Then, a random energy $\hat{E}_k$ is drawn from the Maxwell-Boltzmann distribution consistent with $N_c$ at the desired temperature, and the relative particle velocities are scaled by $\xi = \sqrt{\hat{E}_k/E_k}$. This method generates average kinetic energies and fluctuations consistent with the canonical ensemble [58, 59].

Alternatively, the Andersen thermostat (AT) collision rule [27] implicitly generates isothermal conditions. In an AT collision, a random velocity $\delta \mathbf{v}_i$ consistent with the Gaussian distribution of velocities at the desired temperature is chosen for each particle in a cell, and the velocities are updated by

$$
\mathbf{v}_i(t + \Delta t) = \mathbf{u}(t) + \delta \mathbf{v}_i - \frac{1}{N_c} \sum_{j=1}^{N_c} \delta \mathbf{v}_j,
$$

(4)

where the sum is again taken over the particles in the cell. The last term enforces linear-momentum conservation in the cell. Angular-momentum conservation can also be enforced for the AT collision rule by applying an additional constraint [56].

Solute particles, such as the monomers of a polymer chain, can be coupled to the solvent through the collision step [30]. Embedded particles propagate using standard molecular dynamics methods [9, 10] between MPCD collisions. Typically the MD timestep is much shorter than the MPCD collision time in order to faithfully integrate the MD equations of motion. The embedded particles are then binned into cells with the solvent particles during the collisions. Care must be taken when computing the center-of-mass velocity or kinetic energy of a cell to appropriately weight the masses of the embedded particles, which are typically different from the solvent.
There are many additional variations and features of the MPCD algorithm \[24\], including the modeling of fluid-solid boundaries \[25, 33–37\], nonideal \[60\] and multiphase \[61\] fluids, nonequilibrium flows \[27, 35\], viscoelasticity \[62\], and deformable objects \[63\]. A thorough treatment of these aspects of the algorithm is beyond the scope of the current work. However, the open-source software presented here has been implemented in a way that can be readily extended to support these features.

3. Implementation

In this section, we present our implementation of the previously described MPCD algorithms for GPUs. Throughout, we will focus on optimizations for NVIDIA GPUs (Section 3.1) and use terminology from the CUDA programming model; however, many of the strategies described here can be generalized to other multithreaded programming frameworks. Figure 1 shows a flow diagram summarizing the steps of the MPCD algorithm required to complete one time step of streaming and collision. Each block represents an independent step in the algorithm, and blocks with dashed outlines require communication with nearest-neighbor ranks in MPI. By separating the steps of the algorithm in this way, we ensure a modular design that can be readily extended.

![Figure 1: Flow diagram of a single step of the MPCD algorithm. Each block represents an independent step of the algorithm. Blocks with dashed outlines require nearest-neighbor MPI communication for multi-GPU simulations. Particle sorting is an optional step that is performed periodically.](image-url)

An MPCD time step proceeds as follows. First, particles are streamed to their new positions according to eq. 1. Particles are then migrated onto the appropriate rank (Section 3.3) and binned into cells (Section 3.4). Cell properties such as the center-of-mass velocity are then computed (Section 3.5) before finally the particles undergo a multiparticle collision according to, e.g., eq. 2 or 4 (Sections 3.6 and 3.7). For performance reasons, the MPCD particle data (Section 3.2) is optionally resorted periodically after the cell list is constructed (Section 3.8). Each of these points will be discussed in further detail below.
3.1. Framework

We chose to implement the MPCD algorithm in the HOOMD-blue simulation package [50, 51]. HOOMD-blue is an open-source, general-purpose molecular dynamics code that was designed for NVIDIA GPUs. It is primarily used to study soft matter and perform coarse-grained simulations, and has a rich set of features for these purposes. Unlike other MD software that uses the GPU as an accelerator to the CPU at performance-critical steps, HOOMD-blue performs essentially all computation directly on the GPU [50]. This approach has the advantage that data structures can be optimized for the GPU, and only limited amounts of data must be migrated between the CPU and GPU. The efficient MD engine can be used to simulate particles embedded within the MPCD solvent completely on the GPU. (There is also a CPU-only version if a GPU is unavailable.) HOOMD-blue additionally supports simulations on multiple CPUs and GPUs with MPI domain decomposition and has been shown to scale to thousands of NVIDIA GPUs [51]. Moreover, it has a modular, object-oriented design and flexible Python user interface, which make it straightforward to implement MPCD, as outlined in Figure 1 alongside the existing MD engine and integrate it in users’ simulation scripts.

3.2. Particle data

We considered two designs for the MPCD particle data structures: using HOOMD-blue’s existing MD particle data or creating a standalone container. The first approach is more convenient for the programmer and is the design adopted in the LAMMPS SRD package. However, considerably more information is usually tracked for MD particles than is required for MPCD particles. This can lead to massive memory waste that restricts the problem sizes that can be studied. MPCD particles are also fundamentally different from MD particles, and so may have different optimal data structures. For example, MPCD particles nearly always have the same mass, and so it is more efficient to save this value once for the fluid rather than per particle. Accordingly, we implemented a standalone container for the MPCD particle data in HOOMD-blue.

![Particle data](image)

Figure 2: Particle data and memory requirements (in bytes) per MPCD particle. Each particle property is stored in a separate array. Position and velocity arrays are four-element vectors of 32-bit (or 64-bit) floating-point values, and tag and flag arrays are 32-bit integers. A duplicate is allocated for each array to facilitate sorting and particle removal, indicated by ×2.
The particle data is stored as a structure of arrays. The entries in these arrays for a single MPCD particle are schematically illustrated in Figure 2. The position and velocity vectors of each MPCD particle are stored as floating-point values. In order to promote coalesced memory accesses, the particle positions and velocities are saved as arrays of four-element floating-point vectors. The precision of these vectors can be selected during compilation. The fourth elements of the vectors are used to store additional data or as caches for data that may be commonly requested concurrently with the position or velocity. Currently, an integer identifying the type of the particle is stored in the position, and the current cell index of the particle is cached in the velocity. For purposes of saving and manipulating configurations, each particle is assigned a unique identifying integer tag. The particle tag is expected to be rarely accessed during the simulation, and is used primarily for manipulating particle data through the Python user interface. In MPI simulations, an additional integer is allocated per particle to be used as a flag for removing particles. A duplicate array is allocated for each property in order to facilitate fast swapping of data as particles are sorted or removed. Hence, the particle data requires a maximum of 80 bytes per particle for single-precision vectors and 144 bytes per particle for double-precision vectors.

3.3. Domain decomposition

HOOMD-blue employs a standard spatial domain decomposition using MPI for simulations on multiple GPUs [48, 51]. The simulation box is subdivided by planes along each of the box vectors, and one MPI rank is assigned to each local simulation box. When multiple GPUs are available in a node, a two-tier decomposition attempts to place spatially close ranks within the same node. The MPCD particles are initially decomposed onto each rank during initialization. They are subsequently migrated between ranks when communication is requested by part of the algorithm. The communication algorithm for the MPCD particles, which is similar to that employed for the MD particles [51], is described below. All steps are performed on the GPU except where explicitly noted otherwise.

Particles which have left the local simulation box are marked with an integer consisting of bitwise flags denoting the directions each particle should be sent. All marked particles are subsequently removed from the particle data and staged into a send buffer, while the unmarked particles are compacted into the local data arrays. After packing, the flags of particles in the send buffer are transformed into destination MPI ranks, and the buffer is sorted by destination. Nonblocking MPI calls then exchange particles with a maximum of 26 unique nearest neighbor ranks. If available, a CUDA-aware MPI implementation [64] could operate on buffers in device memory directly, which would permit the MPI library to optimize these transfers asynchronously and take advantage of GPUDirect technologies [65, 66]. However, in preliminary benchmarks, we found little performance gains from using CUDA-aware MPI for this communication, and so instead the entire send buffer is copied from the device to the host before the MPI calls are made. After the communication is completed, received particles are wrapped back into their destination simulation boxes.
and appended to the particle data arrays.

The CPU-only code employs a simpler communication pattern, as used in LAMMPS [48], where particles are exchanged with a maximum of 6 nearest neighbors along the cardinal directions in three rounds of MPI calls. Particles received in each round are checked to see if they require additional communication and, if so, are pushed into the send buffer for the next round of communication. This approach increases the message sizes in order to take advantage of the full network bandwidth and decrease communication latency. However, on the GPU, additional kernel calls and host-device data copies would be required in order to repack the send buffers during each round of communication, leading to a serial bottleneck which makes one round of communication with smaller messages more efficient.

### 3.4. Cell list

Binning particles into local collision cells in order to compute average cell properties like the center-of-mass velocity is central to the MPCD algorithm. In principle, such properties can be computed by direct summation of particle data. However, on the GPU, multiple atomic operations would be required to perform the summation without a read-modify-write race condition, and such operations are typically slow. Instead, it can be more efficient to first construct a list of particles belonging to each cell (Figure 3a) [49], and then perform the summation on particles within the cells. Moreover, a cell list is more flexible for applying different collision rules (see Section 3.7) and can be reused for efficient particle sorting (see Section 3.8).

Westphal et al. proposed a sophisticated method for constructing a cell list using hash tables and partial lists with shared memory staging [49]. In their benchmarks, this method performed well on NVIDIA GPUs with the Fermi architecture, but performance deteriorated for GPUs with the Kepler architecture that succeeded Fermi. Since subsequent GPUs more closely resemble the Kepler architecture, we require a different optimized solution. For example, the cell list could be constructed by first assigning a cell index to each particle, and then sorting the particle indexes using the cell index as a key. This approach has been applied successfully in the HALMD package [67], and it has the benefit that the generated cell list can be stored using an array whose length is exactly equal to the number of particles. However, sorting can be slow when the number of key-value pairs is large, as is the case for MPCD. Additionally, an auxiliary array is required to find the location of each cell’s particles in compact cell list storage, adding to the overhead of reading from the cell list.

In this work, we employ a method for generating the cell list based on limited use of atomic operations that is also currently used in HOOMD-blue for the MD particles. The cell list is stored as a two-dimensional matrix in row-major order [68]. Each row corresponds to a cell, and each column gives the index of a particle in that cell (Figure 3b). The row-major ordering is beneficial for computing cell properties (see Section 3.5). Because of the regular data layout, entries in the cell list can be straightforwardly accessed without an auxiliary array. However, enough memory must be allocated so that each cell can hold the
maximum number of particles in any cell. This layout leads to some memory waste, particularly if one cell has many more particles than another (Figure 3c). The number of MPCD particles in a cell should follow a Poisson distribution [24, 59], and we found that the typical cell list memory was only roughly 4 times larger than a perfectly compact array in our benchmarks (see Section 4). This overhead is comparable to the total memory requirements to build a cell list using an efficient parallel radix sort algorithm [69, 70].

Figure 3: (a) Schematic illustration of particles binned into MPCD cell list. (b) Two-dimensional row-major layout of cell list with one row per cell and columns containing particles. (c) One-dimensional cell list memory with numbers indicating region for each cell and white space showing unused storage per cell.

The cell list is built as follows. Each particle is first binned into a cell (Figure 3a). In order to support random grid shifting, particle positions are optionally shifted by a vector, subject to periodic boundary conditions, during binning. A particle is then inserted into the row of its cell using one atomic operation to determine the column. The cell index is also cached into the fourth element of the particle velocity vector for use in subsequent steps of the MPCD algorithm. The particle is not inserted if the determined column index would exceed the allocated size of the cell list. However, in this case, the maximum number of particles that should be inserted into any cell is still implicitly tracked, and the cell list is subsequently reallocated and recomputed. Reallocation occurs infrequently and primarily during the first few times the cell list is built.

In MPI simulations, cells may overlap the boundaries of domains. This is further complicated by grid shifting, since particles may be shifted outside the local simulation box. To accommodate these boundaries, we introduce a layer of ghost cells around the domain (Figure 4). We compute the minimum number of cells required along each dimension so that a particle shifted from the local simulation box by ±a/2 will still be binned into a cell. Because of this assumption, particles must be migrated to lie within their local simulation boxes before the cell list is built, effectively requiring that particle migration occurs before every MPCD collision. In order to reduce the frequency of particle migration, it has been suggested to use additional ghost cells as a buffer so that particles can diffuse farther off rank [71]. We found that using additional ghost cells was not effective for the GPU because increased cell communication posed a bigger bottleneck.
3.5. Cell properties

After the cell list has been built, the cell properties can be computed by iterating over particles in each cell. The main quantities of interest are the center-of-mass velocity and the kinetic energy. The temperature in the cell, which is defined by the kinetic energy relative to the center-of-mass velocity (see eq. 3), can be determined from these properties. The simplest GPU implementation of this calculation would use one thread per cell to loop over the member particles and accumulate the momentum and kinetic energy [49]. However, this approach has limited parallelism, especially when scaling to multiple GPUs, because the number of cells may be too small to fully occupy a GPU.

Instead, we compute the cell properties using an array of \( w \) threads to process the particles in a cell. Using multiple threads increases parallelism and also promotes coalesced reads from the row-major ordered cell list. Each thread reads from the cell list in a strided fashion to compute the total momentum, mass, and kinetic energy of the particles it has accessed. The number of threads \( w \) is restricted to be a power of two less than the size of a CUDA warp so that these quantities can then be reduced using shuffle instructions available on Kepler and newer NVIDIA GPUs that allow threads within a warp to efficiently read from each other’s registers. The first thread within each array then computes the center-of-mass velocity from the momentum, determines the temperature, and writes the final result. The optimum value of \( w \) depends on the MPCD particle density, the volume of the simulation box, and the specific GPU hardware and is determined using runtime autotuning [51].

It is essential to accurately compute the cell center-of-mass velocity because the MPCD collisions are performed relative to this quantity, and inaccuracies can lead to momentum drift and poor energy conservation. Accordingly, all reductions are performed with double-precision floating-point values even if the particle velocities are stored in single precision. Alternatively, we could perform the operations using double-single
floating-point arithmetic or fixed-point arithmetic. These strategies replace 64-bit floating-point arithmetic with multiple equivalent 32-bit floating-point operations or 64-bit integer fixed-point arithmetic, respectively, which can be faster for certain GPU architectures due to differences in instruction throughput. However, we found that the primary bottleneck in the cell property calculation was actually the memory bandwidth rather than arithmetic operations, and so chose to simply use double-precision floating-point operations for reducing the cell properties (see Section 4.1).

In MPI simulations, communication between ranks is required in order to compute properties in cells overlapping the boundaries. This communication pattern is regular provided that the domain decomposition does not change during the simulation. We adopt a similar communication pattern as for the particle migration on the GPU, and employ nonblocking MPI calls to a maximum of 26 nearest neighbors for both CPU and GPU code pathways. All buffers are packed and unpacked on the GPU if available, and only the minimum required amount of data is communicated. Still, the amount of data that must be transferred from the device to host is large, leading to a significant latency. To mask this latency, we divide the computation of cell quantities into two stages. First, we compute the properties for “outer” cells requiring communication (shaded cells in Figure 4). We then begin the MPI communication, and calculate properties of cells lying fully within the local simulation box (white cells in Figure 4). Last, we finalize communication from the outer cells and finish reducing their properties. We found this strategy to be much more efficient than performing three rounds of communication with fewer, larger messages because it (1) does not lead to serial bottlenecks of host-device data migration and (2) allows computation to overlap with network communication.

The total momentum, energy, and temperature of the system may be required for measurement purposes. These quantities can be obtained by summation of the cell properties, which are reduced using a parallel device-wide reduction scheme. In MPI simulations, we ensure that cells overlapping the boundary are only included in this sum by one of the ranks. The net properties are then reduced across all MPI ranks with a collective communication call.

3.6. Stochastic rotation dynamics

Once the MPCD cell list and cell properties have been computed, it is straightforward to apply the SRD collision rule to the particle velocities. The most important technical challenge to address is how to randomly draw and store the rotation matrix for each cell. The rotation vector must be picked randomly and uniformly from the surface of the unit sphere. We employ the cylindrical projection method for picking points, which requires drawing only two uniform random numbers. This method is particularly efficient on the GPU compared to rejection sampling methods because there is no branch divergence. The rotation axis for each cell is stored in double precision to ensure numerical accuracy of the rotation.

We generate the uniform random numbers using a cryptographic-hash approach to create unique independent microstreams of random numbers. This approach has proven to be highly useful for massively
parallel computing because it allows independent threads to generate random numbers without needing to access or advance a shared state. In this work, we employ the Saru random number generator used throughout HOOMD-blue, although other generators could be straightforwardly substituted. Saru takes three seeds to its hash to initialize a compact state. To generate random vectors for the cell, we feed three seeds: (1) the unique global index of the cell (within the entire simulation box), (2) the current simulation timestep, and (3) a user-supplied seed. This choice of seeding ensures a unique stream (1) for each cell, (2) at each time step, and (3) between simulations. Moreover, because the global cell index is used, the chosen rotation vector for any given cell can be reproduced at any given time step on any rank, regardless of the previous history of drawing numbers. This is particularly useful for MPI simulations because it eliminates the need to communicate random rotation vectors between ranks, significantly decreasing latency.

Particle velocity rotation simply proceeds as outlined in Algorithm 1. Each particle velocity is loaded into memory, including its cell index, which was stored as the fourth element of its velocity (line 1). The average velocity (line 2) and rotation axis (line 3) for the cell are read from memory, and the rotation is applied to the relative particle velocity (lines 6–7). The center-of-mass velocity is added back to the rotated relative velocity (line 11), and the updated velocity is stored (line 12). We found that, regardless of the precision of the particle velocities, it was necessary to perform the rotations in double precision in order to obtain good momentum conservation. We accordingly upcast the particle velocities to double precision before performing any steps in the rotation (line 4). The precision of the rotated velocity vector is then downcast for storage (line 12). We will discuss the ramifications of this type conversion for accuracy of the simulations in Section 4.1.

**Algorithm 1 SRD collision rule.**

1: for each particle $i$ in parallel
2:    \{${v_i, c}$\} $\leftarrow$ particle velocity and cell
3:    $u$ $\leftarrow$ center-of-mass velocity for $c$
4:    $R$ $\leftarrow$ rotation vector for $c$
5:     Cast $v_i$ to double precision.
6:    $\Delta v_i$ $\leftarrow$ $v_i - u$
7:    Rotate $\Delta v_i$ by angle $\alpha$ around $R$.
8:    if thermostat enabled then
9:        $\xi$ $\leftarrow$ scale factor for $c$
10:       $\Delta v_i$ $\leftarrow$ $\xi \Delta v_i$
11:       $v_i$ $\leftarrow$ $u + \Delta v_i$
12:     Store $v_i$ in native precision.
As discussed in Section 2, the SRD collision rule optionally applies a cell-level thermostat. This step requires first computing the rescaling factor $\xi$ per cell according to eq. 3 and then applying this factor to the relative particle velocities. If thermostatting is enabled, we compute $\xi$ for each cell at the same time that we draw the rotation vectors. A random kinetic energy is drawn for each cell from a $\Gamma$ distribution using Marsaglia’s efficient rejection sampling method [75]. This method requires drawing a Gaussian random variable, which we generate by the Box-Muller transformation [76] of two uniform variables to avoid additional branch divergence [77]. Cells having fewer than two particles are assigned a scale factor of $\xi = 1$ because the temperature is not defined in these cases. Velocity rescaling is then applied to the relative velocities after rotation but before they are shifted by the center-of-mass velocity and stored (lines 8–10 in Algorithm 1).

Solute particles are coupled to the MPCD solvent during the collision step. This necessitates including the solute particles in all stages of the algorithm, including cell list construction, cell property calculation, and then the velocity rotation. In HOOMD-blue, the coupled particles are treated as a subset (group) of MD particles. The solute particles evolve according to the standard MD equations of motion between collisions using the velocity Verlet algorithm [9, 10]. Since the solute is coupled only during the collision step, the MPCD particles are only required to be streamed every MPCD collision time $\Delta t$ and not every MD time step.

3.7. Andersen thermostat

The AT collision rule requires the generation of a set of random velocities per MPCD particle and the subsequent reduction of these velocities within each cell in order to apply eq. 4. We first draw random velocities for each particle from a Gaussian distribution using the Box-Muller transformation. The uniform random values are generated using Saru with the timestep, particle tag, and a user-supplied seed. These velocities are then summed using the methods described in Section 3.5. The particle velocities are then updated by adding the randomly drawn velocity for each particle to the cell center-of-mass velocity and subtracting the summed random contribution. The velocity update step is simpler than the SRD algorithm (Algorithm 1) because only simple summation is required and the AT collision rule implicitly thermostats the solvent. However, we found that the AT collision rule performed slower than SRD overall for two reasons: (1) more random numbers must be drawn (per-particle rather than per-cell), and (2) an additional reduction is performed for each cell. This second step especially incurs a performance penalty in MPI simulations because additional communication is required compared to SRD. To help mask some of this latency, we overlap the process of drawing the random velocities with communication during the initial calculation of the center-of-mass velocities for each cell.

3.8. Particle sorting

Many steps of the MPCD algorithm involve processing particles within one cell. Performance of these steps can be improved by first reordering the particle data into cell order [49], which improves data locality.
HOOMD-blue uses a three-dimensional Hilbert curve to sort the MD particles, which is beneficial for evaluating pair forces between particles. The MPCD particles can be sorted more simply into the order they reside in the cell list since this strategy gives optimal ordering for cell property calculation during the first MPCD collision after sorting. Random grid shifting, particle diffusion, and insertion order of particles into the cell list will decrease this ordering on subsequent collisions, but particles still retain some locality. Using the cell list to sort the particles is particularly convenient because it does not require significant additional calculations, and sorting can be injected into the usual MPCD algorithm (see Figure 1). The ability to sort the MPCD and MD particles independently is another advantage of the self-contained MPCD particle data structure described in Section 3.2.

The particles are sorted as follows. First, the cell list is constructed as described in Section 3.4. If sorting should occur at the current MPCD step, the two-dimensional cell list array (Figure 3b) is compacted into a one-dimensional list of MPCD solvent particle indexes. The compaction step is necessary in order to remove any empty entries from cells with fewer particles than the maximum allocation per cell (Figure 3c). Embedded solute particle indexes are also removed to preserve the Hilbert-curve ordering already applied independently to the MD particles. The embedded particles usually comprise only a small fraction of particles in the cells, and so their ordering should have negligible impact on performance. The MPCD particle data is then sorted according to the compacted list, and the indexes in the cell list are updated to the new ordering.

Although particle sorting improves performance of other steps of the MPCD algorithm, there is a significant cost associated with data movement during the sort, especially on the GPU. There is accordingly an optimum frequency of sorting that balances the improved performance from data locality with this added cost. The optimum will depend on the specific properties of the MPCD fluid, the number of particles per GPU, and also the GPU architecture. We therefore advocate tuning the sorting frequency with a series of short simulations to achieve maximum performance.

4. Performance

Performance was benchmarked using the SRD collision rule with typical simulation parameters. MPCD particles having unit mass $m$ were randomly placed into a cubic simulation box of edge length $L$ at number density $\rho = 10/a^3$. Here, the MPCD cell size $a$ defines the unit of length. The particle velocities were drawn randomly from the Gaussian distribution consistent with temperature $T = \varepsilon/k_B$, where $\varepsilon$ is the unit of energy. The SRD rotation angle was $\alpha = 130^\circ$, and the time between collisions was $\Delta t = 0.1 \tau$, where $\tau = \sqrt{ma^2/\varepsilon}$ is the unit of time. Random grid shifting was applied to ensure Galilean invariance. With this choice of parameters, the solvent viscosity was $8.7 \varepsilon \tau / a^3$ and the estimated Schmidt number was 14 [78]. These values are consistent with a liquid-like solvent [29, 79]. The MPCD particle data was sorted every 25
We performed four benchmarks of our MPCD implementation. We first tested the accuracy and performance of the mixed-precision model (Section 4.1) for this SRD fluid. We then assessed the weak and strong scaling efficiency of our implementation (Section 4.2). Finally, we performed a research-relevant benchmark of polymer chains in solution (Section 4.3). These benchmarks are described in detail next and summarized in Table 1.

Table 1: Summary of benchmarks. $L$ is the edge length of the simulation box, and $N$ is the number of MPCD particles. For the weak scaling benchmark, $L$ refers to the size of a cubic box per node. The MPCD parameters for all benchmarks are density $\rho = 10/a^3$, temperature $T = \varepsilon/k_B$, SRD rotation angle $\alpha = 130^\circ$, and collision time $\Delta t = 0.1 \tau$.

| benchmark                  | $L$  | $N$                | simulated time | hardware        |
|----------------------------|------|--------------------|----------------|-----------------|
| precision (Section 4.1)    | 50 $a$ | 1.25 million      | $10^5 \tau$    | Tesla P100 /   |
|                            | 100 $a$ | 10 million        |                | GeForce GTX 1080|
| weak scaling (Section 4.2) | 50 $a$/node | 1.25 million/node | $500 \tau$     | Blue Waters (Tesla K20x) |
| strong scaling (Section 4.2)| 400 $a$ | 640 million       | $500 \tau$     | Blue Waters (Tesla K20x) |
| polymer (Section 4.3)      | 50 $a$ | 1.25 million      | $2000 \tau$    | Tesla P100     |

4.1. Precision

HOOMD-blue’s MD codes can be compiled to use single-precision or double-precision floating-point values for its data structures and calculations. Single precision can give a sizable performance increase compared to double precision due to reduced data size and higher 32-bit floating-point arithmetic throughput. Our MPCD implementation is often bottlenecked by accessing particle data in global memory, and so MPCD should similarly benefit from using single-precision floating-point values when possible. In what we will refer to as the mixed-precision model for our implementation, particle positions and velocities are stored in single precision instead of double precision (Section 3.2), and single-precision floating-point operations are used when possible in the algorithm. However, some key steps always require double-precision values and arithmetic for numerical accuracy (Sections 3.5 and 3.6), prohibiting the exclusive use of single-precision floating-point operations throughout. In these steps, the single-precision particle data is upcast to double precision, and intermediate calculations are performed using double-precision values before downcasting the result to single precision. Here, we study the numerical accuracy and performance of the SRD algorithm for this mixed-precision model compared to a fully double-precision model.

We measured the $x$-component of the center-of-mass velocity $v_{c,x}$ of the benchmark SRD fluid for box sizes $L = 50 \, a$ and $L = 100 \, a$ over $10^6$ MPCD time steps. (Comparable results were obtained for the $y$ and $z$ components.) Random initialization of the particle velocities imparted a nonzero initial value, which we aimed to remove by distributing a constant correction to each particle, as is the typical strategy [9, 10].
For the double-precision model, \( v_{c,x} \) was essentially zero for all times considered after the initial correction was applied (\( |v_{c,x}| \lesssim 10^{-16} \text{ ma}/\tau \)). For the mixed-precision model, however, this process still left a small nonzero value due to loss of precision when the velocities were stored in single precision after the correction. The center-of-mass velocity then fluctuated around zero during the simulation (Figure 5). Importantly, we did not observe any significant systematic drift over the simulation times considered, corresponding to good global momentum conservation. The absolute value of \( v_{c,x} \) remained small and bounded, and decreased for larger numbers of particles (larger \( L \)). It appeared that \( v_{c,x} \) for the mixed-precision model was essentially a random variable due to the digits that were lost when casting the precision of the particle velocities for rotation since a single-precision floating-point value typically contains only 7 or 8 significant digits [80]. Comparable values of \( v_{c,x} \) are obtained in MD simulations with good momentum conservation [67]. We accordingly judged that the accuracy of the mixed-precision model was acceptable and that mixed precision could be used provided that it improved performance.

![Figure 5: Center-of-mass velocity of benchmark SRD fluid along x direction, \( v_{c,x} \), in mixed-precision model over 10^6 MPCD time steps for box sizes \( L = 50 \text{ a} \) and 100 \( \text{a} \).](image)

We tested the performance of the double- and mixed-precision models on two recent NVIDIA GPUs, Tesla P100 and GeForce GTX 1080, using CUDA 8.0. The average performance in time steps per second (larger is faster) is shown in Figure 6 for the \( L = 100 \text{ a} \) box. For the Tesla P100, the mixed-precision model was 1.7x faster than the double-precision model. This difference is close to the maximum 2x speedup that could be obtained from the difference in peak theoretical floating-point performance in single and double precision. For the GeForce GTX 1080, the speedup from the mixed-precision model was slightly smaller at 1.5x. Our benchmarks indicate performance is mostly limited by data accesses, which is perhaps unsurprising for the MPCD algorithm where many key steps involve only little computation. The MPCD algorithm benefits accordingly from the mixed-precision model, which reduces the size of the particle data while delivering reasonable accuracy and performance.
4.2. Scaling

Efficiently utilizing multiple GPUs is a requirement for studying problems too large to reside in the memory of a single GPU and also decreases the time required to complete a simulation. We performed scaling benchmarks of our MPCD implementation on Blue Waters, hosted by the National Center for Supercomputing Applications at the University of Illinois at Urbana-Champaign. HOOMD-blue was compiled using the GNU Compiler Collection (version 4.9.3) and CUDA 7.5 with the double-precision model to provide a performance baseline. The GPU code was tested using Blue Waters’s Cray XK7 nodes, each of which contains one NVIDIA Tesla K20x GPU and one AMD Interlagos 6276 CPU with 8 floating-point cores. Previous benchmarks of HOOMD-blue’s MD code found no benefit of multiplexing multiple MPI processes onto the same GPU, and we expect the same to hold true for MPCD. Accordingly, one MPI rank was assigned per XK7 node. The CPU code was tested using Blue Waters’s Cray XE6 nodes with 16 MPI ranks per node. Each XE6 node has two AMD Interlagos 6276 CPUs (16 total floating-point cores). We chose to compare the CPU and GPU code using these configurations to obtain a practical test of the maximum performance that could be achieved per node hour.

In order to test scaling of the MPCD code to problems with large length scales, we performed a weak scaling test with a constant number of particles per node. A cubic box with edge length $50a$ was assigned per node so that there were 1,250,000 particles per node on average. We first performed 2,000 SRD time steps in order to allow all runtime autotuners to determine their optimal parameters. These parameters were then fixed, and the performance was measured over 5,000 SRD time steps with HOOMD-blue’s internal profiler enabled. We found that overhead from profiling incurred a small performance penalty of roughly 5% for the GPU in this test. Each benchmark was repeated three times to determine the average performance, and the 95% confidence interval was estimated from these measurements.

Both the CPU and GPU codes gave excellent weak scaling performance up to 1,024 nodes, as shown in Figure 7. At this largest node count, there were over one billion MPCD solvent particles in the simulation.

Figure 6: Time steps per second for SRD collision rule in double precision and mixed precision on NVIDIA Tesla P100 and GeForce GTX 1080 GPUs for the $L = 100a$ simulation box.
box. The CPU weak scaling efficiency relative to two nodes was greater than 90% for all node counts tested, while the GPU scaling efficiency was greater than 65%. Most of the loss of efficiency in the GPU code came from an increase in the time required to compute cell properties, which requires the most communication in our implementation. There was also a large initial drop in efficiency for the GPU code as the domain decomposition was increased from one dimension (at 2 nodes) to three dimensions (at 8 nodes). This drop was expected because the amount of communication per node increased. The CPU code did not show this initial drop because there were 16 MPI ranks per node, and so the simulation box was three-dimensionally decomposed for all node counts. The inset to Figure 7 shows the absolute performance of the two codes in time steps per second. The GPU code was roughly 3 times faster than the CPU code on 2 nodes and 2 times faster on 1,024 nodes. For reference, the theoretical peak double-precision performance of the K20x is roughly 4 times the XE6 node.

We then tested the strong scaling performance of the MPCD code for a large simulation box with \( L = 400 \alpha \), which corresponded to 640 million solvent particles. Efficient strong scaling is required to fully utilize computational resources and decrease the time to solution. As for the weak scaling tests, we measured the time required to complete 5,000 SRD time steps with internal profiling enabled after runtime autotuning was completed. The performance overhead of this profiling was again roughly 5% on the GPU for this test, and was most significant at the largest node counts. We measured the scaling from 32 nodes up to 1,024 nodes. Although the CPU code could be run on smaller node counts, the GPU code required a minimum of 32 nodes due to the limited capacity of the GPU memory (6 GB). Each benchmark was repeated three times.

Figure 8 shows the strong scaling performance and efficiency (inset) of both the CPU and GPU codes.
The CPU code exhibits excellent strong scaling, with over 80% efficiency on 1,024 nodes (16384 MPI ranks). The GPU code exhibits similarly good scaling, with some efficiency lost at high node counts. In order to investigate this loss of efficiency further, we separated the time required per step into three main components: particle migration, cell property calculation, and all other steps (including streaming and collision) that require no communication. Figure 9 shows these components for various node counts. It is clear that the communication steps, especially cell property calculation, limit the scaling efficiency at high node counts.

Figure 8: Strong scaling performance for benchmark SRD fluid in a cubic simulation box with $L = 400\,a$ on NCSA Blue Waters. The total number of particles was $N = 640,000,000$. GPU benchmarks (circles) were performed using 1 MPI rank (1 NVIDIA Tesla K20x GPU) per XK7 node, while CPU benchmarks (squares) used 16 MPI ranks (16 floating-point cores) per XE6 node. Solid lines indicate ideal scaling, and the inset shows the strong scaling efficiency.

Figure 9: Profile of GPU strong scaling for SRD fluid on NCSA Blue Waters (see Figure 8). Required time per step is shown for the total simulation (circles), core SRD methods (squares), particle migration (triangles), and computing cell properties (diamonds). Solid black line indicates ideal scaling of the cell property communication.

At small node counts, the cell property calculation is in principle dominated by the inner cells, which can effectively mask most of the latency from communication. At larger node counts, however, the latency associated with both host-device data migration as well as the MPI communication itself become significant.
Theoretically, it is expected that the time for communication of cell properties should scale with $1/P^{2/3}$ for $P$ MPI ranks due to the surface-to-volume ratio of the domain decomposition \[71\]. Initially, we did indeed observe this scaling as indicated in Figure \[9\] but it is clear that for $P > 256$ much weaker scaling was obtained. More detailed profiling revealed that this bottleneck was due to host-device data migration. This latency might be reduced using an MPI library that operates on device buffers directly since the library can optimize copies asynchronously to pipeline messages or take advantage of GPUDirect RDMA technologies \[64\]-\[66\]. Unfortunately, we found essentially no performance benefits from these optimizations on Blue Waters due to limited support for GPUDirect RDMA.

Overall, the GPU code is faster than the CPU code by a factor between 2 and 3. As a point of reference, we note that our CPU code also outperforms a comparable benchmark with LAMMPS SRD (31 March 2017 release) \[47\]-\[48\] by nearly a factor of 2 when running on 128 XE6 nodes on Blue Waters. (There are some minor SRD parameter differences in the LAMMPS benchmark due to limitations of that implementation.)

Our weak and strong scaling benchmarks clearly demonstrate the feasibility of using GPUs to efficiently perform and accelerate MPCD simulations at massive scale.

### 4.3. Polymer solution benchmark

In addition to the pure SRD fluid benchmarks, we also performed a more complex simulation of a solution of polymers embedded in the benchmark SRD solvent on an NVIDIA Tesla P100 GPU using the double-precision model and CUDA 8.0. The polymers were represented by a bead-spring model as described in \[78\]. Each polymer chain consisted of 50 beads, and the number of chains was varied from 16 to 1,024, giving between 800 and 51,200 total monomers in the simulation box with $L = 50a$. The equations of motion for the polymers were integrated between collisions with the solvent using the velocity Verlet algorithm with a time step of 0.002$\tau$. (An MPCD collision occurred every 50 MD timesteps.) The MD neighbor list used to compute nonbonded pair forces had a buffer radius of 0.4$a$. The MPCD solvent particles were streamed every 50 MD timesteps because their updated positions were only required for collisions. We performed a short run of 200$\tau$ to determine the optimal runtime kernel launch parameters and then profiled performance for 2,000$\tau$.

In order to measure the contribution of the MPCD solvent to the simulation time, we performed simple Langevin dynamics simulations \[73\] of the polymers as a baseline. To obtain comparable long-time polymer dynamics, we adjusted the monomer friction coefficient \[81\] to give a long-time polymer diffusion coefficient consistent with the value predicted by the Zimm model for dilute polymer solutions \[82\]-\[84\]. We emphasize that this baseline does not include any hydrodynamic interactions, which is a clear advantage of the MPCD simulations, and simply serves as a point of comparison of the contributions of MD and MPCD to the overall simulation time.
The total required simulation time increased nearly linearly with the monomer density, shown in Figure 10. The density spanned from the dilute to semidilute polymer concentration regimes. Initially, the Langevin dynamics simulation time did not increase much with density due to limited occupancy of the GPU for small numbers of MD particles. The required time for the MPCD simulations closely tracked the Langevin dynamics simulations, indicating that the performance of the MPCD algorithm itself did not depend significantly on the polymer concentration. This trend is reasonable because the embedded particles comprise only a small fraction of the particles in an MPCD cell. The MPCD simulations were at most 1.4x slower than the Langevin dynamics simulations, which is only a modest cost to incorporate hydrodynamic interactions. Even though there were many more solvent particles than monomers, the MPCD streaming and collision steps happened infrequently compared to the MD step.

![Figure 10: Required time to simulate polymer solutions at various concentrations for 2,000 \( \tau \) using MPCD (circles) and Langevin dynamics (squares) on an NVIDIA Tesla P100 GPU.](image)

This benchmark demonstrates the strengths of implementing MPCD within an existing MD package designed to run exclusively on the GPU. We were able to easily perform simulations with up to 51,200 monomers embedded in 1,250,000 solvent particles on a single GPU and obtained excellent performance. Moreover, the MD algorithms could be used as a “black box” without special considerations for hiding latency of migrating MD particle data to the GPU.

5. Conclusions

We developed an implementation of the MPCD algorithm for simulating mesoscale hydrodynamics that is optimized to exploit the massively parallel computational capabilities of GPUs. Spatial domain decomposition onto multiple GPUs is supported with MPI, enabling simulations of mesoscale hydrodynamics at length and time scales that would be otherwise challenging or impossible to access. We showed that our MPCD implementation efficiently scales to hundreds of GPUs. Here, the performance was primarily bottlenecked by host-device data transfers, which we speculate may be mitigated in computing environments.
having multiple GPUs within a node and using a CUDA-aware MPI library in conjunction with GPUDirect RDMA. We found that adopting a mixed-precision computing model for the MPCD particle data improved performance on a single GPU with acceptable numerical accuracy. We also showed for a benchmark polymer solution how MPCD can be used to incorporate hydrodynamics into research-relevant simulations with only modest additional computational cost. Our MPCD implementation is available open-source as part of the HOOMD-blue project (version 2.3.0) and should prove useful for conducting simulations of soft matter and complex fluids when hydrodynamic interactions are important.

In this work, we have focused on implementing the most fundamental components of the MPCD algorithm. However, there are many extensions of MPCD that are of interest for areas of active research, including coupling the solvent to solid boundaries such as walls and imposing external fields to drive solvent flow. We have designed our software to be modular in order to readily support these extensions. For example, fluid-solid coupling and flow could be added by modification of the streaming step. We hope to continue to expand our software with support from the community to incorporate these and other features in the future.

Note.—During the publication process, we determined an improved method for packing particle data for migration compared to the version that we benchmarked. In the improved version, we partition the indexes of the particles to be migrated from the particles that are to be retained. Holes in the particle data arrays left by migrated particles are backfilled from the list of retained particles to maintain compact arrays, reducing the number of read and write operations compared to our previous implementation based on stream compaction. For the SRD benchmark fluid with $L = 200a$ on 8 NVIDIA Tesla P100 GPUs, this optimization reduced the time to pack the MPI buffers by a factor of 12.5x, the overall particle migration time by a factor of 2.8x, and the total simulation time by a factor of 1.3x. The performance timings reported in Figures 7-9 should also be similarly improved by this optimization.

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