Multi-Particle Collision Dynamics Algorithm for Nematic Fluids

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Research on transport, self-assembly and defect dynamics within confined, flowing liquid crystals requires versatile and computationally efficient mesoscopic algorithms to account for fluctuating nematohydrodynamic interactions. We present a multi-particle collision dynamics (MPCD) based algorithm to simulate liquid-crystal hydrodynamic and director fields in two and three dimensions. The nematic-MPCD method is shown to successfully reproduce the features of a nematic liquid crystal, including a nematic-isotropic phase transition with hysteresis in 3D, defect dynamics, isotropic Frank elastic coefficients, tumbling and shear alignment regimes and boundary condition dependent order parameter fields.

1 Introduction

As a state of soft condensed matter with intermediate symmetries between highly ordered crystals and disordered fluids, nematic liquid crystals are both phenomenologically fascinating and commercially valuable. No longer are liquid crystals of interest only to those producing liquid crystal display technology; now scientists interested in microfabricated systems, microelectromechanical devices, composite materials, biosciences and active gels are exploiting the unique properties of liquid crystals in novel applications. Interest in complex geometries (such as confining geometries, composite materials, biosciences and active gels) require versatile mesoscopic algorithms that can account for non-trivial boundary conditions. Likewise research into “hypercomplex liquid crystals” and self-assembly would benefit from efficient methods to simulate nematohydrodynamic baths for macromolecular and colloidal solutes.

Such elaborate systems present a considerable challenge for traditional particle-based numerical methods. Lattice Monte Carlo simulations have been very successful in simulating nematic liquid crystals and continue to be widely employed due to their computational frugality. However, out-of-equilibrium dynamics and relaxation mechanisms require more computationally costly methods. Off-lattice simulations of hard anisotropic particles and soft pair-potentials have played an important role in understanding generic liquid crystalline phases, but are limited to simple systems. Molecular dynamics simulations can account for molecular detail with a range of coarse-graining, including fully atomistic, generic molecules and the mesoscopic approach of dissipative particle dynamics. Even mesoscopic simulations can become computationally expensive when large numbers of constituent particles are required and so are generally limited to simplified systems.

Investigating hypercomplex fluids or dynamics within demanding geometries calls for the continued development of versatile and computationally efficient coarse-grained algorithms. One mesoscopic simulation technique that has shown promising capabilities in simulating fluctuating hydrodynamics of isotropic solvents is the multi-particle collision dynamics (MPCD) algorithm. MPCD has been used to simulate hydrodynamic interactions between macromolecules, colloids, vesicles and swimmers. It has even been extended to simulate viscoelastic fluids and electrohydrodynamics. In this work, we propose an extension to the MPCD method to efficiently simulate fluctuating nematohydrodynamics (nematic-MPCD).

2 Method

Multi-particle collision dynamics algorithms forgo simulating molecular-scale interactions between constituent molecules. Instead, the continuum description is discretised into many artificial, point-like MPCD particles that stochastically exchange momentum while respecting conservation laws for mass, momentum and energy. This is sufficient to reproduce the hydrodynamic equations of motion on sufficiently long length and time scales. Mesoscopic MPCD algorithms can dramatically reduce computational costs compared to simulations that explicitly calculate molecular pair-potentials and are well suited to simulating flowing systems involving non-trivial boundary conditions, finite Reynolds numbers, and fluctuating hydrodynamics, which are...
ideal for moderate Péclet number systems \(^{41,42}\).

Here, we develop a nematic-MPCD method to efficiently simulate fluctuating nematohydrodynamics, by assigning an orientation pseudo-vector to each MPCD point-particle and updating orientations through a local and stochastic nematic multi-particle orientation dynamics (MPOD) operator. Backflow and shear-alignement dynamics are ensured by coupling the MPCD and MPOD operators. In § 3, we demonstrate that nematic-MPCD reproduces the necessary physical properties to simulate a nematic liquid crystal when the velocity and director fields are coupled. In a very recent article, Lee and Mazza introduced an interesting hybrid, non-local MPCD method for liquid crystals\(^{43}\). The main difference to our approach is that their particles carry a director field that is coupled to the fluid through a discretisation of the stress terms in a simplified Ericksen-Leslie formalism of nematohydrodynamics.

In this section, we begin by reviewing a traditional Andersen-thermostatted MPCD algorithm that conserves angular momentum. We go on to describe the implementation of the MPOD operator for nematic fluids and the two-way coupling between the director and velocity fields. Finally, we describe how potentially complex boundary conditions can be implemented.

2.1 Traditional MPCD for Isotropic Fluids

The fundamental insight of MPCD algorithms is that continuous mass and momentum fields can be discretised into MPCD point-particles (labelled i). Each MPCD particle possesses a position \(\mathbf{r}_i\), mass \(m_i\) and velocity \(\mathbf{v}_i\), and which interact through multi-particle, near-equilibrium stochastic collision events within lattice-based cells (labelled c) defined by a size \(a\), population \(N_c\), centre of mass velocity centre \(\mathbf{v}_{cm.c}\) and moment of inertia \(I_c = \Sigma_k m_k \left( r_k^2 - \mathbf{r}_{cm,c} \cdot \mathbf{r}_k \right)\) of the point-particles in cell c relative to their centre of mass \(\mathbf{r}_{cm,c}\) where \(\mathbf{r}_c = \mathbf{r}_i - \mathbf{r}_{cm,c}\).

The MPCD algorithms consist of two steps. Each MPCD particle streams ballistically for a time \(\delta t\) such that its position at time \(t + \delta t\) becomes

\[
\mathbf{r}_i(t + \delta t) = \mathbf{r}_i(t) + \mathbf{v}_i(t) \delta t.
\]

Multiple particles then undergo collision events, in which momentum is transferred between MPCD particles. To exchange momentum, the simulation domain is partitioned into cubic cells of thermally varying number density \(\rho_c = N_c/a^d\) in d-dimensions. Discretising space into MPCD cells breaks Galilean invariance, though this can be remedied by randomly shifting the cell grid at each time step\(^{44}\). The collision operator \(\Xi_{i,c}\) is a non-physical exchange designed to be stochastic and also to conserve the net momentum within each cell c,

\[
\mathbf{v}_i(t + \delta t) = \mathbf{v}_{cm,c}(t) + \Xi_{i,c}.
\]

Many choices for the collision operator exist, which result in different versions of MPCD, including the original Stochastic Rotation Dynamics\(^{26,45}\) and a Langevin version of the algorithm\(^ {46}\). In this work, we utilise the Andersen-thermostatted collision operator\(^ {46,47}\)

\[
\Xi_{i,c} = \xi_j - \langle \xi_j \rangle_{N_c} + \left( I_c^{-1} \cdot \delta L_c \right) \times \xi_j. \tag{3}
\]

where \(\xi_j\) is a random velocity drawn from the Maxwell-Boltzmann distribution \(f_{vel}(\xi, k_BT)\) for thermal energy \(k_BT\) and \(\langle \xi_j \rangle_{N_c}\) is the average of the \(N_c\) random velocity vectors in the \(c^{th}\) cell during the instant of the collision event. Randomly generating the \(\xi_j\) from the equilibrium distribution \(f_{vel}\) in the moving reference frame ensures that the algorithm is locally thermostatted\(^ {46}\). The third term in the collision operator is a correction included to remove the angular momentum introduced by the collision operator

\[
\delta L_c = \sum_{j} m_j \{ \xi_j \times (\xi_j - \mathbf{v}_j) \}. \tag{4}
\]

Though the nematic-MPCD method does not strictly depend on this choice for \(\Xi_{i,c}\), coupling the velocity field to the director field is accomplished by respecting this conservation law (see § 2.3.2).

2.2 Multi-Particle Orientation Dynamics for Nematic Fluids

We now that propose nematic liquid crystals can be simulated via a nematic-MPCD algorithm by including an orientation field.

Each MPCD particle is assigned an orientation \(\Psi\), while each cell acquires a tensor order parameter

\[
\Psi_c = \frac{1}{d-1} \langle d\Psi - \frac{1}{d} \mathbf{1} \rangle_{N_c}. \tag{5}
\]

For a nematic fluid, the largest eigenvalue is the local scalar order parameter \(S_c\) of the cell and the local Frank director \(\mathbf{n}_c\) is parallel to the corresponding eigenvector.

Orientations interact through a positive, globally specified interaction constant \(U\). In physical liquid crystals, the energy \(U\) represents inter-molecular interactions and will be a non-constant function of temperature or molecular details such as nematogen dimensions and density. In this nematic-MPCD algorithm, the interaction constant \(U\) is the simulation specified energy that governs the local evolution of orientations. Taking inspiration from the Andersen-thermostatted MPCD collision operator, we implement a stochastic multi-particle orientation dynamics operator for orientation. The essential requirements are that the MPOD operator must be local and near equilibrium, with no gradient terms in the collision operator. Therefore, we propose the orientation collision event

\[
\Psi (t + \delta t) = \Psi_c \left( U, \Psi_c(t) \right), \tag{6}
\]

where the multi-particle orientation operator \(\Psi_c\) generates a random orientation \(\Psi (t + \delta t)\) drawn from the equilibrium probability distribution \(f_{on}(U, \Psi_c(t))\) about the local director \(\mathbf{n}_c(t)\) calculated from the tensor order parameter. The nematic collision operation causes the MPCD point-particles to change their orientation by \(\Psi_{col,i} = \delta \Psi_{col,i}/\delta t = [\Psi(t + \delta t) - \Psi(t)]/\delta t\) without altering the local director \(\mathbf{n}_c(t)\).
2.2.1 Maier-Saupe Distribution:

As in the traditional Andersen-thermostatted MPCD algorithm, the multi-particle orientation operator depends on the condition of local, near-equilibrium statistics. In this work, we assume that the local equilibrium distribution for the orientation field obeys the Maier-Saupe self-consistent mean-field theory and so is an exponential function of $u_n \equiv u \cdot \mathbf{n}_c$:

$$f_{\text{ori}}(U, S_c, \mathbf{n}_c) = e^{-\beta w_{\text{MF},c}(U, S_c, u_n)},$$

where $g$ is a normalisation constant, $\beta \equiv 1/k_B T$ and each cell’s mean-field interaction potential is

$$w_{\text{MF},c} = -US_c u_n^2 + \frac{U}{d} (S_c - 1).$$

The second term does not depend on $u_n$ and so the distribution of $u_n$ is determined by $e^{\beta BU_S c}$. When the scaled energy $BU_S c$ is small, all orientations are equally likely but when $BU_S c$ is large the distribution becomes sharply oriented about $\mathbf{n}_c$.

2.2.2 Generating the Maier-Saupe Distribution:

When $BU_S c \approx 1$, a Metropolis algorithm for $w_{\text{MF},c}$ generates the random orientations. However, the distribution can be more efficiently approximated in the limits of $BU_S c \gg 1$ and $BU_S c \ll 1$. In the strong mean field limit $BU_S c \gg 1$, $f_{\text{ori}}$ is sharply centred about $\mathbf{n}_c$ such that $u_n^2 = \cos^2 \theta_n \approx 1 - \theta_n^2$, which means that the distribution for $\theta_n$ can be approximated as Gaussian $f_{\text{ori}} \sim e^{-\beta BU_S c \theta_n^2}$. The Gaussian approximation is used when $BU_S c > 5$. On the other hand, when $BU_S c \ll 1$ the exponent can be expanded and the cumulative distribution function of $f_{\text{ori}}$ can be approximated as $W = \int_{-\delta_n}^{\delta_n} d\theta_n e^{\beta BU_S c \theta_n^2} \approx u_n + BU_S c u_n^3/3$. Random values of $u_n$ can be generated through the transformation $u_n = 2^{-1/3} \kappa/BU_S c - 2^{1/3}/\kappa$, where $\kappa(r) = (3r(BU_S c)^2 + 9(BU_S c)^4 r^2 + 4(BU_S c)^3 r^{3/2})^{1/3}$ and $r \in [0, 1]$. This expansion is used when $BU_S c < 0.5$.

2.3 Two-way Coupling

Coupling between the director and fluid flow is crucial for reproducing nematohydrodynamics since flows can rotate the nematogens (§2.3.1) and the rotation of nematogens in turn produces hydrodynamic motion, referred to as backflow (§2.3.2). We model the coupling by treating the nematic-MPCD particles as a dilute suspension of asymmetric particles rotating through an non-inertial fluid. The nematogens are implicitly envisioned as rotating through the viscous fluid that they themselves represent.

2.3.1 Shear Alignment: Velocity→Orientation Coupling:

We treat the nematic-MPCD particles as a dilute suspension of rods subject to a net torque

$$\Gamma_{\text{net},i} = \Gamma_{\text{H},i} + \Gamma_{\text{col},i} + \Gamma_{\text{ext},i} = 0,$$

where $\Gamma_{\text{H},i}$ is due to the flow field’s vorticity $\omega = [\nabla \times (\nabla \times \mathbf{V})]/2$ and shear rate $D = [\nabla \mathbf{V} + (\nabla \mathbf{V})^T]/2$ (Fig. 8; insets) $48, \Gamma_{\text{col},i} = \gamma_k \mathbf{u}_i \times \mathbf{u}_{col,i}$ is the fluctuating thermal torque due to the MPD collision and $\Gamma_{\text{ext},i}$ represents any external torques such as those due to magnetic or electric fields. In this algorithm, we assume viscously over damping such that the net torque is zero.

Solving Eq. 9 for the total rate of rotation of particle $i$ shows that $\dot{\omega}_i = \dot{\omega}_{\text{H},i} + \dot{\omega}_{\text{col},i}$, where $\dot{\omega}_{\text{H},i}$ obeys the discretised Jeffery’s equation for a slender rod

$$\frac{\delta \omega_{\text{H},i}}{\delta t} = \Gamma_{\text{H}} \left[ (\mathbf{u}_i \cdot \mathbf{D}) - \alpha (\mathbf{u}_i \cdot \mathbf{D} - \mathbf{u}_i \cdot \mathbf{D}) \right],$$

where $\lambda$ is the bare tumbling parameter and $\Gamma_{\text{H}}$ is a heuristic shear coupling coefficient, a simulation parameter that tunes the alignment relaxation time relative to $\delta t$. For the rotation of an individual prolate particle subject to shear flow, $\chi_{\text{H}} = 1$. When the shear coupling coefficient is set to zero ($\chi_{\text{H}} = 0$) there is no coupling of the director to the velocity field. Equation 10 is compatible with the torque used in Leslie-Ericksen continuum nematicohydrodynamics and has been utilized in molecular Doi theory to predict the Leslie viscosities of nematic fluids $49,50$.

2.3.2 Backflow: Orientation→Velocity Coupling:

In the nematic-MPCD algorithm, backflow coupling is accounted for by balancing the hydrodynamic torque $\Gamma_{\text{H},i}$ on each nematogen with an equal and opposite change in angular momentum to the velocity collision operator $\delta \mathbf{Z}/\delta t$. It is simplest to merely use Eq. 9 to identify $\Gamma_{\text{H},i} = -\Gamma_{\text{col},i} - \Gamma_{\text{ext},i}$ in the non-inertial limit.

To balance this torque with the hydrodynamic drag on the fluid, the opposite of the net change in the angular momentum $\delta \mathbf{Z}/\delta t = \sum_i \delta \mathbf{Z}_i = \sum_i \Gamma_{\text{col},i} - \Gamma_{\text{H},i} = \Gamma_{\text{col},i} - \Gamma_{\text{H},i}$ is transferred to the linear momentum portion of the algorithm. The MPCD collision operator $\Xi_{i,c}$ (Eq. 3) is thus modified to account for liquid crystal backflow becoming

$$\Xi_{i,c} = \xi - \sum_{N_i} \mathbf{E} \left[ \frac{1}{N_i} \right] + \left[ \mathbf{E} - \mathbf{E}_i \right] \times \mathbf{L}_c.$$
The boundary conditions also set the easy direction describing the preferred orientation of the liquid crystal director at a surface. During a bounce-back collision event with the surface, the orientation $\mathbf{n}_i$ of the impinging nematic-MPCD particle is set parallel to the surface's easy direction. This anchoring is not strong, as will be seen in § 3.5. For homeotropic boundary conditions, the easy direction is normal to the surface. For planar boundary conditions, the easy axis is parallel to the surface. In this case, all in-plane directions can be equivalent or a single preferred direction can be specified. If no preferred direction is specified then the boundary is said to be non-anchoring.

2.5 Units and Chosen Simulation Parameters

Values are expressed in MPCD simulation units — time, mass, energy and length are given respectively by time step $\delta t$, particle mass $m$, thermal energy $k_B T$ and cell size $a = \delta t \sqrt{k_B T / m}$. The new MPOD parameters are also stated these units. The interaction constant $U$ has units $k_B T$, while the rotational friction coefficient $\gamma_R$ has units $k_B T \delta t$. Both the bare tumbling parameter $\lambda$ and the shear coupling coefficient $\chi_{HH}$ are dimensionless.

Except when otherwise stated, the simulations presented in this manuscript vary input parameters about the following set of values: In this manuscript, simulations are carried out in 2D ($d = 2$) for a system of size $V = 50^2$ with periodic boundary conditions and a mean number density of $\rho = \langle N_e \rangle / 20$. The MPCD particles are randomly initiated with positions from a uniform distribution, velocities from the Maxwell-Boltzmann distribution and aligned nematic orientations. Parameter values are chosen to be $m = 1, k_B T = 1, \delta t = 1, a = 1, U = 15, \gamma_R = 0.01, \lambda = 2$ and $\chi_{HH} = 1$.

3 Results

Having described the implementation of the nematic-MPCD algorithm, we now characterise the resulting properties of the liquid crystal. We first consider how the isotropic-to-nematic phase transition depends on the simulation parameters, particularly the heuristic shear coupling coefficient and number density of MPCD particles. We measure the nematic-isotropic hysteresis and explore the dynamics of the defect annihilation rate as the system orders. Elastic free energy drives defect annihilation and we measure the isotropic Frank elastic coefficients to be a linear function of the interaction constant. The response of the isotropic phase to an ordering wall is characterised.

3.1 Nematic-isotropic transition

When $\beta U$ is small, the nematic-MPCD algorithm exists as an isotropic fluid state with a small global order parameter $S$ (Fig. 1; inset-left). When $\beta U$ is large a nematic state is formed (Fig. 1; inset-right). Maier-Saupe self-consistent theory predicts that the nematic-isotropic transition is first order (Fig. 1). Although the nematic-MPCD algorithm assumes near-equilibrium and so uses the Maier-Saupe distribution on the local cell level, the scalar order parameter and directors are spatially varying fields rather than mean-field values.

In 3D systems of large enough size, periodic boundary conditions and no shear coupling, the nematic-MPCD algorithm does exhibit a strongly first order nematic-isotropic phase transition (Fig. 1). In these simulations, the nematic fluid is initialised in the nematic state and resides in a periodic cube of size $50^3$. The system discontinuously jumps from zero to a global scalar order parameter $S^* = 0.860 \pm 0.003$ at $[\beta U]^* = 4.20 \pm 0.05$.

In 2D the nematic-isotropic transition is expected to become a Kosterlitz-Thouless-type transition. The present simulations demonstrate that the transition is no longer first order, increasing from zero at $[\beta U]^* = 4.1 \pm 0.1$ in a $50^2$ system (Fig. 1). The second order nature of the nematic-isotropic transition is a direct result of the nematic-MPCD’s ability to accommodate spatially varying fields. Future studies should more fully characterise the nature of the nematic-isotropic transition in 2D.

3.1.1 Global vs. Local Scalar Order Parameter:

By replacing the local scalar order parameter $S_e$ of each cell $e$ with the system’s globally determined order parameter $S$ in each cell’s local mean-field interaction potential $w_{MF,e}$ (Eq. 8), the 2D transition becomes first order (Fig. 1). The order parameter curve remains relatively unchanged except near the phase transition. The transition from the isotropically disordered state is retarded compared to the spatially varying case that uses the local order parameters but suddenly jumps to $S^* = 0.51 \pm 0.01$ at $[\beta U]^* = 5.00 \pm 0.05$.

3.1.2 Variation of Simulation Parameters:

In order to assess the impact of varying simulation parameters on the nematic-isotropic transition, we initially omit the velocity–orientation coupling by setting $\chi_{HH} = 0$ in Eq. 10 (Fig. 2; top row). We consider varying time step $\delta t$, mass $m$, temperature $k_B T$, rotational friction coefficient $\gamma_R$, bare tumbling parameter $\lambda$ and mean number density $\rho$. In the zero-coupling limit, Fig. 2
(top row) shows that none of the MPCD simulation parameters have a significant effect on the nematic ordering. Only the mean number density has an observable affect on the curve. At extremely low mean number densities, the transition occurs at a slightly larger interaction constant. It should be noted that when an individual nematic-MPCD particle is alone in an MPCD cell neither its velocity nor its orientation are altered.

3.1.3 Impact of Coupling Fluctuating Hydrodynamics:

When the shear coupling coefficient $\chi_{HI}$ is zero the global scalar order parameter $S$ rises from zero in the isotropic phase to $S = 1$ in the $\beta U \to \infty$ ordered limit. This is no longer true when $\chi_{HI} \neq 0$ (Fig. 2; bottom row). As the hydrodynamic coupling is restored by increasing $\chi_{HI}$, the value of the scalar order parameter decreases for a given interaction constant. This occurs because fluctuations in the velocity field introduce an additional source of noise through Eq. 10 when $\chi_{HI} \neq 0$. These fluctuations reduce the order in the director field and move the system away from the fully ordered state of $S = 1$. With full coupling, only the rotational friction coefficient $\gamma_R$ is seen to have no impact on the $S$ curve (Fig. 2; bottom row). This is because $\gamma_R$ controls the rotational relaxation dynamics and does not influence the equilibrium state.

When $\chi_{HI} = 0$, the mean number density is the only simulation parameter seen to have any observable effect on the isotropic to nematic transition and then only at extremely low values (Fig. 2; top row). At the lowest number density the transition is less sharp and occurs at a slightly higher interaction constant $\beta U$. While $S$ depends weakly on $\rho$ when $\chi_{HI} = 0$ (Fig. 3), it is a strong function of number density when $\chi_{HI} = 1$. Fig. 3 shows the global scalar order parameter as a function of density for $U = 15$. When the shear coupling parameter $\chi_{HI}$ is set to zero, the system remains in the nematic state even at quite low densities. On the other hand, when coupling is included, $S$ increases from zero with mean number density. In fact, the order parameter in Fig. 3 exhibits a continuous transition and the nematic-MPCD algorithm possesses a nematic-isotropic transition as a function of density when $\chi_{HI} = 1$.

Since there is a nematic-isotropic transition as a function of density (Fig. 3), it is clear that the shear coupling coefficient has a larger effect at lower number densities than it does at larger densities. Fig. 4 shows the strong interaction limit of $S$ (measured at $\beta U = 100$ and 500) for various densities as a function of coupling. For a low mean number density of $\rho = 5$, Fig. 4 shows that the strong limit drops from $\lim_{\beta U \to \infty} S \approx 1$ when $\chi_{HI} = 0$ to only $0.038 \pm 0.003$ when the algorithm is fully coupled. Fluctuations are pronounced because of the small number fluctuations of particles in each MPCD cell. By increasing the mean number density $\rho$, the continuum limit is approached and fluctuations become less severe. When $\rho = 20$ and the algorithm is fully coupled ($\chi_{HI} = 1$), the strong interaction limit is $S = 0.80 \pm 0.01$ (Fig. 4).

Throughout this work, we set the mean number density $\rho = 20$, though a lower density may suffice in many situations.
When the algorithm is fully coupled with $\chi_{HI} = 1$, the tumbling parameter can also increase the susceptibility of the order field to velocity fluctuations through Eq. 10. This can be seen in Fig. 2 (bottom row). When $\lambda = 5$, fluctuations in the shear rate $\nu$ fully disorder the system.

### 3.1.4 Hysteresis:

Hysteresis is expected in 3D due to the first order nature of the nematic-isotropic transition. By comparing 3D nematic-MPCD simulations initialised with the director field in the isotropic state (as in in § 3.1) to those initialised in the nematic state, a striking hysteresis loop is observed in Fig. 5. The interaction constant, $\beta U$, is fixed throughout the duration of individual simulations. At these system sizes, the width of the hysteresis is measured from Fig. 5 to be $\beta \Delta U^\ast = 0.70 \pm 0.03$ and the difference in order parameters at the transition points is $\Delta S^\ast = 0.20 \pm 0.04$.

### 3.2 Defect Annihilation Dynamics

The process of transitioning from the isotropic to nematic phase discussed in § 3.1 is controlled by the dynamics of topological defects. Though the increasing interaction constant $U$ generates local order along a spontaneous direction $\mathbf{n}_s$, neighbouring regions may break symmetry along any other direction. Therefore, many $\pm 1/2$ topological defects rapidly emerge from the disordered director field. Pairs of oppositely charged defects must approach each other and annihilate for global ordering.

Since the 2D number density $\rho_D = 0.0080 \pm 0.0005$ of defects is initially quite high, the annihilation rate $R_D = -\rho_D$ is large but falls rapidly (Fig. 6; inset a). As the density decreases, the average separation between topological defects increases and annihilation events become less frequent (compare Fig. 6a showing an example system at $t = 40$ to Fig. 6b showing the same system at $t = 400$). A variety of scaling relations for the annihilation rate $R_D(t) \sim t^{-(\nu + 1)}$ have been put forward. Mean-field arguments predict $\nu = 1$, purely diffusive kinetics suggest $\nu = 0.5$ and scaling arguments give $\nu = 6/7$\textsuperscript{57}. Furthermore, the scaling law possesses short-time logarithmic corrections\textsuperscript{58,59}. When measured on short times between $t \in [10^{-3}]$, the nematic-MPCD annihilation rate appears to decay as $\nu = 0.74 \pm 0.02$ (Fig. 6) but the exponent increases to $\nu = 0.83 \pm 0.04$ when evaluated over $t \in [10^{-1}, 10^0]$ (Fig. 6), which is in agreement with the $\nu = 6/7$ scaling prediction.

### 3.3 Frank Elastic Coefficients

We have considered how the nematic state arises from the isotropic state. Let us now consider the nematic response to distortions in the director field. Gradients in the director field $\mathbf{n}$ lead to the free energy density per unit volume $f = K_{\text{splay}} (\nabla \cdot \mathbf{n})^2/2 + K_{\text{twist}} (\mathbf{n} \cdot \nabla \times \mathbf{n})^2/2 + K_{\text{bend}} (\mathbf{n} \cdot (\nabla \times \mathbf{n}))^2/2$. Splay, bend and twist deformations are illustrate in Fig. 7; insets. Since distortion are typically large compared to molecular length scales, the Frank elastic coefficients $K_{\text{splay}}$, $K_{\text{twist}}$ and $K_{\text{bend}}$ are macroscopic material properties.

One technique for obtaining the Frank coefficients from particle-based simulations is to measure the equilibrium, orientational fluctuation spectrum\textsuperscript{60–63}. In reciprocal space, the tensor order parameter for each wave vector is $\hat{Q}(k) = \rho^{-1} \sum_{i=1}^{N} \frac{1}{2\pi^2} (d_{\mathbf{n}i} \cdot \mathbf{k}) \exp(i \mathbf{k} \cdot \mathbf{r}_i)$. We work in a varying director-based coordinate system, in which $\mathbf{n}_s = [0, 0, 1]$ and the wave vector is in the 13-plane, i.e. $\mathbf{k} = [k_1, 0, k_3]$. In this coordinate system, the equipartition theorem\textsuperscript{60} relates the the low $|k|$ limit of the orientational fluctuations to the Frank coefficients

$$\langle \hat{Q}_{\alpha} (k) \hat{Q}_{\alpha} (-k) \rangle = \frac{9}{4} \frac{SVk_B T}{K_\alpha k_1^2 + K_{\text{bend}} k_3^2}$$

for $\alpha = 1, 2$ (splay, twist). Through Eq. 12, the Frank coefficients may be determined as fitting parameters of the fluctuation spectrum in reciprocal space. A large system size of $V = 30 \times 30 \times 30$
and $N = 5.4 \times 10^5$ MPCD particles are used in the following simulations to ensure sufficient statistics for many near-zero $|k|$ values and accurate fits.

The resulting Frank coefficients of the nematic-MPCD fluid are shown in Fig. 7. Although splay, twist and bend deformations may possess differing coefficients in some physical systems, simple scaling suggests that all three elastic constants are of order $\sim U/\lambda$ and theoretical considerations of the Maier-Saupe self-consistent model$^{64}$ predict $K_i = \ell^2 \rho U S^2 [1 + C_i]/6$, where $i = \{splay, twist, bend\}$ and $\ell$ is a characteristic interaction distance. The different constants $C_i$ depend on the molecular details and higher moments of the orientation distribution$^{64}$. The nematic-MPCD simulations ostensibly exhibit isotropic elasticity. This is expected because, in the limit that the rod length is small compared to the interaction length, the constants $C_i$ are safely neglected and the Frank coefficients are predicted to converge$^{64}$. Since the nematic-MPCD algorithm simulates point-like nematogens with a characteristic interaction length equal to the finite cell size, the one-constant approximation applies.

In agreement with simple scaling and the Maier-Saupe self-consistent predictions, the measured elastic coefficients for the nematic-MPCD algorithm are linear with respect to the interaction constant $U$ and number density $\rho$ (Fig. 7). Together, they are fit to $K_i = \{84 \pm 4.113 \pm 6.142 \pm 7\} U$ for $\rho = \{15, 20, 25\}$, respectively, in Fig. 7.

### 3.4 Tumbling and Shear Alignment

Thus far, we have considered quiescent nematic fluids. We now turn our attention to flowing systems. Microscopically, the director field is influenced by shearing flows through Eq. 10 and as described schematically in Fig. 8; inset.

In the infinitely dilute limit of a suspension of spheroidal particles, the bare tumbling parameter is a geometrical entity that can be cleanly related to the particle aspect ratio $p$ by $\lambda = (p^2 - 1)/(p^2 + 1)$, which goes to unity as $p \to \infty$ and is zero for spheres ($p = 1$). However, interactions between nematogens in a nematic fluid allow the actual tumbling parameter to deviate from the isolated-slender-rod value and distributions of molecules can exhibit effective tumbling parameters that are larger than unity. Such fluids are referred to as aligning-nematics because there is a stable alignment angle, the Leslie angle $\theta_L$, between the director and shear field.

By considering a Fokker-Planck equation for the probability distribution of orientations, Archer and Larson$^{65}$ found that the flow tumbling behaviour of ellipsoidal particles with $\lambda = (p^2 - 1)/(p^2 + 1)$ is determined by the tumbling parameter

$$\lambda' = \lambda \frac{155 + 48S_4 + 42}{105S_4}. \tag{13}$$

In the nematic-MPCD algorithm, $\lambda$ is the specified simulation parameter for the bare tumbling parameter, the magnitude of which can be set larger than unity. We shall see that $\lambda'$ as given by Eq. 13 is the resulting tumbling parameter of the nematic-MPCD algorithm. In Eq. 13, $S_4$ is the fourth moment of the Maier-Saupe probability distribution. The distribution can be written as an expansion of orthogonal Gegenbauer polynomials $C_n^{(p)}(x)$ in $d$-

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Fig. 6 Number density of topological defects as a function of time with and without shear coupling. Inset (a): Annihilation rate of defects $\rho_D$. Simulation parameters from § 2.5 are used with a 2D system size of $500 \times 500$. In the defect maps, colour denotes local scalar order parameter and defects are mapped with red circles marking $-1/2$ defects and blue pentagons marking $+1/2$ defects.

Fig. 7 Frank elastic coefficients for splay, twist and bend as a function of interaction constant $U$. Simulation parameters from § 2.5 are used for a system of size $50 \times 50 \times 50$ and number densities $\rho = \{15, 20, 25\}$ ($N = \{1.875, 2.5, 3.125\} \times 10^6$ nematogens) with $\chi_H = 0$. Insets a,b,c respectively depict splay, twist and bend.
When the magnitude of the bare tumbling parameter $\lambda$ is larger than unity, the nematogens do not tumble but rather align with the shear. For these tumbling parameters, Eq. 10 has the solution
\[
\tan \theta_L = \pm \sqrt{\frac{|\lambda'| - 1}{|\lambda'| + 1}}.
\]

Good agreement is found between Eq. 16 and the simulations using Lees-Edwards boundary conditions when the tumbling parameter (Eq. 13) is greater than unity. As the tumbling parameter tends to 1+, the Leslie angle approaches zero. In this limit, the nematogens orient along the flow direction. When $\lambda' \gg 1$, the Leslie angle of the nematic-MPCD fluid approaches $\pi/4$ as predicted by Eq. 16.

### 3.5 Wall-Induced Ordering

Confining walls affect the nematic ordering. In the isotropic state, anchoring can cause ordering in the vicinity of the walls. We consider a 2D nematic-MPCD fluid confined between two no-slip plates separated by $L = 100$. The plate at $y = 0$ enacts homoeotropic boundary conditions, which order the nematic fluid. The plate at $y = L$ is a non-anchoring boundary, which does not set a condition for $\dot{y}$.

When the interaction constant is much less than the nematic-isotropic transition value $[\dot{\beta}U]$ (§ 3.1), the order decreases to the isotropic state far from the wall. As the interaction constant is increased, the value of the scalar order parameter $S_0(U) \equiv S(U, y = 0)$ at the wall increases (Fig. 10; inset). This signifies that the anchoring is not infinitely strong and is strongly effected by the value of $U$.

Additionally, the order extends further into the bulk fluid as $U$ increases. The characteristic distance the order extends from the wall is a coherence length $\xi$ (Fig. 10). One can predict that the order decays as $S(U, y) = S_0(U) e^{-y/\xi}$ by considering the total free energy functional to be the highest order term in the Landua-De Gennes free energy and the deformation free energy. The co-
Coherence length is a function of the elastic constant and the distance from the transition, \( \xi \propto \left( \frac{3K_{\text{elas}} + 2K_{\text{ani}}}{T - T_{\text{c}}} \right)^{1/2} \). As the nematic-isotropic transition is approached, the coherence length diverges. The nematic-MPCD simulations accurately reproduce the exponential decay far below the transition point (Fig. 10; inset).

It was seen in § 3.3 that \( K_i \sim U \) so the coherence length takes the form

\[
\xi = \left( \frac{c_1 \beta U}{1 - U/U_{\text{c}}} \right)^{1/2}. \tag{17}
\]

The theory captures the rapid growth of the coherence length near the nematic-isotropic transition but goes to zero as \( U \rightarrow 0 \), while the simulations do not (Fig. 10). The coherence length of the nematic-MPCD does not go to zero because MPCD algorithms are not able to resolve material properties on length scales comparable to the cell size \( a \). If a second fitting parameter \( c_2 = (1.505 \pm 0.005) a \) is included as in Fig. 10 to account for this discretisation effect, then Eq. 17 well-represents the divergence of the coherence length in the isotropic phase.

In the nematic phase, the order still decreases exponentially from \( S_0 \) but decays to a non-zero bulk value (Fig. 10; inset). Except near the nematic-isotropic transition, the order parameter falls steeply to its bulk value over a length scale comparable to a single MPCD cell.

4 Conclusions

We have proposed a nematic-MPCD algorithm for simulating fluctuating nematohydrodynamics. Nematic-MPCD uses traditional Andersen-thermostatted MPCD with conservation of angular momentum to integrate the velocity field and a novel multi-particle orientation dynamics (MPOD) collision operator to progress the director field. By stochastically drawing orientations from the local Maier-Saupe equilibrium distribution, the MPOD operator updates the orientations without numerically evaluating gradients. In addition, the two-way coupling between the MPCD and MPOD operators represents backflow and shear-alignment. We have shown that this nematic-MPCD algorithm reproduces the essential physical properties of a simple nematic fluid, such as the nematic-isotropic phase transition, topological defects, Frank elasticity and shear alignment.

The nematic-MPCD algorithm holds much promise as a tool for simulating nematohydrodynamics, but future studies should carefully investigate the anchoring strength (since modifications to the no-slip conditions were required in traditional MPCD\(^{52-54}\)) and work towards kinetic theories to quantitatively predict the nematic material properties, such as the Frank elastic and Leslie viscosity coefficients, as functions of simulation parameters. Though simple, the algorithm holds exciting potential for simulating a wide variety of soft matter systems. For example defect dynamics within topological microfluidic devices\(^9\) or porous media\(^66\) could be modelled, exploiting the ease with which the algorithm can handle complicated confining geometries. It would also be of interest to consider dispersed nanoparticles, carbon fibres\(^67\) or swimmers\(^68\) within a liquid crystal host, and it is relatively easy to imagine that generalized Maier-Saupe theories\(^69\) could be implemented in the MPOD collision operator to simulate cholesteric or biaxial liquid crystals.

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