The ENUF method—Ewald summation based on nonuniform fast Fourier transform: Implementation, parallelization, and application

Sheng-Chun Yang1 | Bin Li 2 | You-Liang Zhu3 | Aatto Laaksonen4,5,6,7 | Yong-Lei Wang4

1School of Computer Science, Northeast Electric Power University, Jilin, China
2School of Chemical Engineering and Technology, Sun Yat-sen University, Zhuhai, China
3State Key Laboratory of Polymer Physics and Chemistry, Changchun Institute of Applied Chemistry, Chinese Academy of Sciences, Changchun, China
4Department of Materials and Environmental Chemistry, Arrhenius Laboratory, Stockholm University, Stockholm, Sweden
5State Key Laboratory of Materials-Oriented and Chemical Engineering, Nanjing Tech University, Nanjing, China
6Centre of Advanced Research in Bionanoconjugates and Biopolymers, Petru Poni Institute of Macromolecular Chemistry Aleea Grigore Ghica-Voda, Iasi, Romania
7Department of Engineering Sciences and Mathematics, Division of Energy Science, Luleå University of Technology, Luleå, Sweden

Correspondence
Yong-Lei Wang, Department of Materials and Environmental Chemistry, Arrhenius Laboratory, Stockholm University, SE-10691 Stockholm, Sweden.
Email: wanyong@auestu.se

Funding information
Education Department of Jilin Province, China, Grant/Award Number: JJKH20190696KJ; Fundamental Research Funds for the Central Universities, Grant/Award Number: 2019G7076; Knut and Alice Wallenberg Foundation, Grant/ Award Number: KAW 2018-0380; Ministry of Research and Innovation of Romania, Grant/ Award Number: PN-III-P4-ID-PCCF-2016-0050; National Natural Science Foundation of China, Grant/Award Number: 21774129; State Key Laboratory of Supramolecular Structure and Materials of Jilin Universtit, Grant/Award Number: SKLSSM202031; Vetenskapsrådet, Grant/ Award Number: 2019-03865

Abstract
Computer simulations of model systems are widely used to explore striking phenomena in promising applications spanning from physics, chemistry, biology, to materials science and engineering. The long range electrostatic interactions between charged particles constitute a prominent factor in determining structures and states of model systems. How to efficiently calculate electrostatic interactions in simulation systems subjected to partial or full periodic boundary conditions has been a grand challenging task. In the past decades, a large variety of computational schemes has been proposed, among which the Ewald summation method is the most reliable route to accurately deal with electrostatic interactions between charged particles in simulation systems. In addition, extensive efforts have been done to improve computational efficiencies of the Ewald summation based methods. Representative examples are approaches based on cutoffs, reaction fields, multi-poles, multi-grids, and particle-mesh schemes. We sketched an ENUF method, an abbreviation for the Ewald summation method based on the non-uniform fast Fourier transform technique, and have implemented this method in particle-based simulation packages to calculate electrostatic energies and forces at micro- and mesoscopic levels. Extensive computational studies of conformational properties of polyelectrolytes, dendrimer-membrane complexes, and ionic fluids demonstrated that the ENUF method and its derivatives conserve both energy and momentum to floating point accuracy, and exhibit a computational complexity of $O(N \log N)$ with optimal physical parameters. These ENUF based methods are attractive alternatives in molecular simulations where high accuracy and efficiency of...
simulation methods are needed to accelerate calculations of electrostatic interactions at extended spatiotemporal scales.

1 | INTRODUCTION

Computer simulations of molecular systems have rapidly expanded over the past decades, and have been widely used to study phase behaviors of materials and biological systems.\textsuperscript{[1,2]} Electrostatic interactions between charged particles play a prominent role in determining structures, dynamics, and states of these physical systems, leading to many important applications in academia and industrial communities.\textsuperscript{[3–8]} An accurate description of electrostatic interactions in model systems is a nontrivial task in computer simulations. The slow decay feature of electrostatic interactions with respect to particle distance poses a significant challenge to model charged simulation systems as the computation of electrostatic interactions between charged particles is extremely time consuming.\textsuperscript{[1,2,4,5]} The spherical cutoff treatment used for short-ranged interactions (such as Lennard–Jones potentials) ignoring particle interactions beyond a certain range is inadequate for electrostatic interactions because any arbitrary truncation leads to nonphysical artifacts. Therefore, one has to take electrostatic interactions between all pairs of ion species into consideration, which is termed as direct summation method, leading to an unfavorable computational complexity of $O(N^2)$ (where $N$ is the number of charged particles in simulation systems). As simulation system size expands, the calculation of electrostatic interactions becomes the major computational bottleneck for a thorough understanding of phase behaviors of charged physical systems at extended spatiotemporal scales.\textsuperscript{[1,2,4,8]}

A traditional way to sum electrostatic interactions between charged particles and all their infinite periodic images is the Ewald summation method.\textsuperscript{[9]} By introducing a differentiable and localized function, the Ewald summation method recast the total electrostatic interaction, a single slowly and conditionally convergent series, into a short range particle–particle interaction part that can be calculated using spherical cutoff treatment in real space and a long range interaction part for smeared charges that can be computed by solving the Poisson’s equation in reciprocal space.\textsuperscript{[1,2]} Although this representation is exact, it contains infinite summation terms and therefore, calls for error-controlled approximations for their applicable utilization in molecular simulations of charged model systems. The parameters entering the Ewald summation method, that is, the interaction range of short range part, the number of Fourier modes in long range part, and the splitting parameter controlling the relative weight of short range and long range terms, can be optimized in such a way that the overall performance of the Ewald summation method is reduced to $O(N^{3/2})$.\textsuperscript{[1,2,4]}

Although the Ewald summation method represents a substantial improvement respect to the direct summation method and removes the quadratic complexity, the numerical effort is still too large for simulation systems extending to millions of charged particles and long-time simulations. Alternative splitting methods having the same underlying idea as the Ewald summation method were developed to accelerate the solvation of the Poisson’s equation in reciprocal space by taking advantages of the fast Fourier transform (FFT) technique, leading to an $O(N\log N)$ scaling of computation time. Examples include the particle–particle particle-mesh (PPPM) Ewald summation method,\textsuperscript{[10–12]} the particle-mesh Ewald summation method (PME),\textsuperscript{[13,14]} and a variety of derivatives from these splitting methods.\textsuperscript{[15–19]} These methods have been successfully employed in the past decades at varied levels of molecular simulations, but the related physical parameters should be carefully optimized for speedy and accuracy. Although these splitting methods are substantially faster than the standard Ewald summation method, their accuracy is inferior. Errors are inevitably introduced in the particle-mesh scheme, which first interpolates particle charges onto a uniform mesh, and thereafter extrapolates the solution of the Poisson’s equation represented on mesh back to charged particles.\textsuperscript{[13,14,17–19]} Therefore, the quality of interpolation and extrapolation plays a critical role in determining computational accuracy of these splitting methods.\textsuperscript{[11,12,17–19]}

In previous works, we sketched an ENUF method\textsuperscript{[20]} an abbreviation for the Ewald summation method based on the nonuniform fast Fourier transform (NFFT) technique,\textsuperscript{[21,22]} to calculate electrostatic energies and forces between charged particles in molecular simulations. The ENUF method is easy-to-implement and efficient for calculating long range electrostatic interactions, and additionally, both electrostatic energy and momentum are conserved to floating point accuracy. Indeed, the ENUF method is the starting point of the subsequent development of particle–particle NFFT with periodic boundary condition \textsuperscript{[23]} By choosing a set of optimal physical parameters, the ENUF method gives a good precision as desired and bears a computational complexity of $O(N\log N)$.\textsuperscript{[8,20,24]} Later, the ENUF method was implemented in the dissipative particle dynamics (DPD) framework to calculate electrostatic interactions between charge density distributions at mesoscopic level.\textsuperscript{[24–26]} The ENUF and ENUF-DPD methods were adopted to explore the dependence of conformational properties of polyelectrolytes on charge fraction, ion concentration and counterion valency of added salts,\textsuperscript{[26]} to investigate specific binding structures of dendrimers on bilayer membranes and the corresponding permeation mechanisms,\textsuperscript{[27]} and to study heterogeneous structures and dynamics in ionic liquids (ILs) and how electrostatic interactions between charged particles affect these properties at extended spatiotemporal scales.\textsuperscript{[8]} In addition, great endeavors have been made in recent years to accelerate scientific computation using, for example, dedicated and
specialized hardware and high-performance accelerator processors.\cite{28} The commodity graphics processing unit (GPU) and the compute unified device architecture (CUDA) represent a disruptive technology advance in simulation hardware, providing a mature programming environment, and have been the majority of investigations in computational material science.\cite{29} At present, most molecular simulation packages support GPU acceleration.\cite{35-39} We have implemented the ENUF and ENUF-DPD methods in an open source GALAMOST (GPU-accelerated large-scale molecular simulation toolkit) package.\cite{8,40,41} In addition, several (hybrid) parallelization strategies based on gridding\cite{42} and Near-Distance algorithms\cite{43} have been developed to accelerate the evaluation of electrostatic energies and forces using GPU and CUDA technology.\cite{44,45} These derivatives of the ENUF and ENUF-DPD methods exhibit distinct computational efficiencies in handling long range electrostatic interactions between charged particles and charge density distributions at multiple spatiotemporal scales.

In the current contribution, we present a comprehensive review on the detailed implementation of the ENUF method in molecular simulations and DPD frameworks based on CPU nodes, the hybrid parallelization of the ENUF and ENUF-DPD methods using GPU and CUDA toolkit, the determination of effective interaction parameters to achieve an optimal computational complexity, and representation applications of the ENUF and ENUF-DPD methods in treating long range electrostatic interactions between point charges and charge density distributions in inorganic crystals, polyelectrolytes, dendrimer-membrane complexes, and IL systems.

2 | IMPLEMENTATION AND PARALLELIZATION OF THE ENUF AND ENUF-DPD METHODS IN CPU FRAMEWORK

2.1 | The Ewald summation method

We consider a simple cubic simulation system with box length $L$ consisting of $N$ charged particles, each one carrying partial charge $q_a$ at position $r_a$ and interacting with each other according to the Coulomb's law. An overall charge neutrality is assumed in such a simulation system and the boundary condition without cutoff is represented by replicating the simulation box in three dimensional (3D) space. The total charge-charge electrostatic interaction energy is given as

$$U^E(r^N) = \frac{1}{4\pi\varepsilon_0} \sum_{n=0}^\infty \sum_{a}^{N} \sum_{b \neq a}^{N} \frac{q_a q_b}{|r_{ab} + nL|},$$

where $n = (n_x, n_y, n_z)$, and $n_x$, $n_y$, and $n_z$ are arbitrary integers representing a replication of charged particles in the whole 3D space. The summation over $n$ takes into account all periodic images, and the $\dagger$ symbol indicates that the self-interaction terms for all charged particles are omitted when $n = 0$. The variable $\varepsilon_0$ is the permittivity (dielectric constant) of the vacuum space.

The direct summation of Equation (1), although simple to implement, suffers a major drawback as a direct numerical evaluation of Equation (1) is excessively computational demanding.\cite{1,2,4} In the triply periodic case, the direct and infinite summation in Equation (1) is conditionally convergent for charge neutral systems, and the computational results (electrostatic energies and forces between charged particles) depend on the order of summation. In fact it was discovered that any conditionally convergent series can be rearranged to yield a series which converge to any prescribed summation.\cite{1,2} Such a situation is very similar to the case when a linear equation has an infinite number of solutions because it is under-determined; by adding a set of conditions a unique solution will be defined. In other words, the summation result is not well defined unless one specifies a detailed procedure to sum up these terms. For electrostatic energies and forces between charged particles, a physically relevant summation order has to be prescribed, and the boundary condition (spherical, cubic, cylindrical, etc.) of the surrounding medium has to be specified.\cite{1,2,4}

The Ewald summation formula for the triply periodic case was derived by Ewald in 1921.\cite{9} The resulting formula imposes two choices: a spherical summation order and an assumption that the dielectric constant of the surrounding medium is infinite, that is, it is a conductor, which is often referred to "tin foil" boundary condition. To gain more physical insights, we consider the electric field potential generated by a charged particle with partial charge $q_a$ located at $r_a$, and the electric field potential is

$$\phi_a(r) = \frac{1}{4\pi\varepsilon_0 |r-r_a|}.$$  

The electric field potential generated by $N$ charged particles and their periodic images at $r_a$ is

$$\phi(r_a) = \frac{1}{4\pi\varepsilon_0} \sum_{b=0}^N \sum_{b \neq a}^N \frac{q_b}{|r_{ab} + nL|}.$$  

Herein, we define $\phi_a(r_a)$ as the electric field potential generated by all other charged particles and their periodic images at $r_a$, excluding ion $a$ itself,

$$\phi_a(r_a) \equiv \phi(r_a) - \phi_a(r_a) = \frac{1}{4\pi\varepsilon_0} \sum_{b=0}^N \sum_{b \neq a}^{N} \frac{q_b}{|r_{ab} + nL|}.$$  

Therefore, the charge-charge electrostatic interaction energy in Equation (1) can be rewritten as

$$U^E(r^N) = \frac{1}{2} \sum_a q_a \phi_a(r_a).$$
The Ewald summation method splits the slowly convergent in Equation (1) into two terms that exhibit exponentially fast and absolute convergence at fixed level of accuracy. The partial charges, described by a collection of delta functions \( \rho_0(r) = q_a \delta(r - r_a) \), are decomposed into two parts by adding and subtracting a set of Gaussian-type charge density distributions. \[^{[46]}\]

\[
\rho_b(r) = \rho_b^R(r) + \rho_b^\sigma(r),
\]

\[
\rho_b^R(r) = q_a \delta(r-r_a) - q_a G_a(r-r_a),
\]

\[
\rho_b^\sigma(r) = q_a G_a(r-r_a).
\]

where

\[
G_a(r) = \frac{1}{(2\pi\sigma^2)^3/2} e^{-\frac{r^2}{2\sigma^2}}.
\]

\( \sigma \) is the standard deviation parameter of the Gaussian-type charge density distribution. In literature, \( \alpha \equiv 1/\sqrt{2\sigma} \) denoted as the Ewald convergence parameter is always used in molecular simulations. This splitting scheme leads to the electric field potential \( \phi_a(r) \) being described by two terms,

\[
\phi_a(r) = \phi_a^R(r) + \phi_a^\sigma(r),
\]

\[
\phi_a^R(r) = \frac{q_a}{4\pi\varepsilon_0} \frac{\delta(r-r_a) - G_a(r-r_a)}{|r-r_a|} d^3r,
\]

\[
\phi_a^\sigma(r) = \frac{q_a}{4\pi\varepsilon_0} G_a(r-r_a) d^3r.
\]

The electric field potential \( \phi_{\text{ion}}(r_a) \) generated by all other charged particles and their periodic images at \( r_a \) excluding ion \( a \) itself can be decomposed in a similar way,

\[
\phi_{\text{ion}}(r_a) = \phi_{\text{ion}}^R(r_a) + \phi_{\text{ion}}^\sigma(r_a).
\]

Correspondingly, the charge–charge electrostatic energy in Equation (5) is given as

\[
U^E(r_a) = \frac{1}{2} \sum_{a} q_a \phi_{\text{ion}}^R(r_a) + \frac{1}{2} \sum_{a} q_a \phi_{\text{ion}}^\sigma(r_a).
\]

The electric field potential \( \phi_G(r) \) generated by the Gaussian-type charge density distribution \( G_a(r) \) can be obtained by solving the Poisson’s equation,

\[
\nabla^2 \phi_G(r) = - \frac{q_a}{\varepsilon_0} G_a(r).
\]

In spherical coordinate system, \( \phi_G(r) \) is symmetric and only depends on the magnitude of \( r = r_a \), and therefore, it is described as

\[
\phi_G(r) = \frac{q_a}{4\pi r_a} \text{erf}(\alpha r_a).
\]

where \( \text{erf}(x) = \frac{2}{\sqrt{\pi}} \int_0^x e^{-t^2} dt \) is the well-known error function. Therefore, the electric field potential \( \phi_b^R(r) \) and \( \phi_b^\sigma(r) \) in Equation (8) can be expressed as

\[
\phi_b^R(r) = \frac{q_a}{4\pi\varepsilon_0} \text{erf}(\alpha |r-r_a|),
\]

\[
\phi_b^\sigma(r) = \frac{q_a}{4\pi\varepsilon_0} \text{erf}(\alpha |r-r_a|).
\]

in which \( \text{erf}(x) \equiv 1 - \text{erf}(x) \) is the complementary error function. As \( \lim_{\alpha \to \infty} \text{erf}(x) = 1 \), the \( \phi_b^R(r) \) is a long-ranged nonsingular potential and the \( \phi_b^\sigma(r) \) is a short-ranged singular potential. The short-ranged singular potential \( \phi_b^\sigma(r) \) generated by all other charged particles and their periodic images at \( r_a \) excluding ion \( a \) itself is given by

\[
\phi_b^\sigma(r_a) = -\frac{1}{4\pi\varepsilon_0} \sum_{a} q_a \text{erf}(\alpha |r_a|) = -\frac{1}{4\pi\varepsilon_0} \sum_{a} q_a \text{erf}(\alpha |r_a + nl|).
\]

The short range part of electrostatic energy \( U^E(r_a) \) in Equation (10) can be rewritten as

\[
U^E(r_a) = \frac{1}{2} \sum_{a} q_a \phi_{\text{ion}}^R(r_a) = \frac{1}{24\pi\varepsilon_0} \sum_{a} q_a \text{erf}(\alpha |r_a + nl|).
\]

Choosing a suitable value of the Ewald convergence parameter \( \alpha \), the short range part of electrostatic energy extends no longer than a cutoff distance and can be expressed as

\[
U^E(r_a) = \frac{1}{24\pi\varepsilon_0} \sum_{a} q_a \text{erf}(\alpha |r_a + nl|).
\]

and therefore, this part can be obtained from a direct summation in real space calculations.

As \( \phi_b^R(r) \) is a long-ranged nonsingular potential, the long range part of electrostatic energy \( U^E(r_a) \) in Equation (10) is not feasible via a direct computation in real space summations. It is noteworthy that all Gaussian-type charge density distributions constitute a periodic function and can be described as

\[
\phi^\sigma(r) = \sum_{n} \phi^\sigma(r + nl).
\]

Considering electrostatic contributions from all charged particles, \( \phi^\sigma(r) \) is the electric field potential generated by a periodic array of charged particles. This indicates that \( \phi^R(r) \) and \( U^E(r_a) \) are smooth periodic functions and hence their Fourier transformations exhibit fast decay in reciprocal space.

By solving the Poisson's equation in reciprocal space.
we can get the electric field potential generated by all Gaussian-type charge density distributions as

$$\phi^K(r) = \frac{1}{V_0 \sum_{a} N_a \sum_{b} \frac{d_{ab}}{k^2} e^{-k^2/4\alpha^2}}$$

where $V = L^3$ is the volume of the simulation system. The long range part of electrostatic energy can be expressed as

$$U^{\text{LR}}(r_0) = \frac{1}{2V_0 \sum_{a} N_a \sum_{b} \frac{d_{ab}}{k^2} e^{-k^2/4\alpha^2}} \xi_{ab}$$

Similarly, the long range part of electrostatic force acting on charged particle $a$ can be obtained as

$$F_a^{\text{LR}}(r_0) = -\nabla_a U^{\text{LR}}(r_0)$$

From electrostatic energy we can easily obtain the electrostatic force $F_a^\text{E}$ acting on charged particle $a$ by taking a partial derivative of the electrostatic energy $U_a^\text{E}$ with respect to its position $r_a$.

$$F_a^\text{E}(r_a) = -\nabla_a U_a^\text{E}(r_a)$$

Splitting electrostatic force using a similar procedure as recasting electrostatic energy, we can get the total electrostatic force acting on charged particle $a$ from real and reciprocal summations,

$$F_a^\text{E}(r_a) = -\nabla_a \left[ U_a^\text{ER}(r_a) + U_a^\text{EK}(r_a) + U_a^\text{ES}(r_a) \right]$$

The short range part of electrostatic force acting on charged particle $a$ can be obtained as

$$F_a^\text{SR}(r_a) = -\nabla_a \left[ \frac{1}{4\pi \epsilon_0} \sum_{b} \left( \frac{1}{r_{ab}} \right) \frac{q_a q_b e^{-r_{ab}/\alpha}}{r_{ab}} \right]$$

The implementation of the Ewald summation method requires the Ewald convergence parameter $\alpha$, a preset energy accuracy parameter $\delta \ll 1$, and two cutoffs ($r_c$ for real space and $r_e$ for reciprocal space). These parameters are inter-correlated with the following conditions:

$$e^{-r_c^2/(4\alpha^2)} \leq \delta \Rightarrow r_c = \frac{\alpha}{\pi} L \Rightarrow r_e \propto \xi N^{1/3},$$

with

$$\xi = \frac{\alpha}{\pi} \sqrt{-\log(\delta)} \Rightarrow r_e \propto \frac{\alpha}{\pi} L$$

In practical applications with presetted $\delta$, it is straightforward to choose a suitable value for $n_c$, and thereafter one can determine $\alpha$ and $r_c$ directly from Equation (32).
2.2 Discrete Fourier transforms for nonequispaced data

The FT for nonequispaced data points is a generalization of FFT.\cite{21,47}

The basic idea of NFFT is to combine standard FFT algorithm with various window functions, which are well localized both in spatial and in frequency domains. Representative window functions include Gaussian, B-spline, Sinc-power, and Kaiser-Bessel types. A controlled approximation using a cutoff scheme in frequency domain and a limited number of terms in spacial domain results in an aliasing error and a truncation error, respectively. The aliasing error is controlled by an oversampling factor \( \sigma_p \) and the truncation error is determined by the number of terms, \( m \), in spatial approximation.\cite{21}

For a finite number of Fourier coefficients \( f_k \in \mathbb{C} \) with \( k \in I_M \), we wish to evaluate the trigonometric polynomial \( f(x) = \sum_{k \in I_M} f_k e^{-2\pi i k x} \) at \( N \) nonequispaced points \( \{x_i \in \mathbb{R} \mid i = 0, 1, ..., N-1 \} \). The space of the \( d \)-variable function \( f \in \mathbb{D}^d \) is restricted to the space of \( d \)-variable trigonometric polynomial \( e^{-2\pi i k \cdot x} \) with a degree of \( M \) \((t = 0, 1, ..., d-1)\) in the \( t \)-th dimension. The possible frequencies \( k \) are well collected in the multi index set \( I_M \) with

\[
I_M = \left\{ k = (k_t)_{t=0}^{d-1} \in \mathbb{Z}^d : -\frac{M_t}{2} \leq k_t \leq \frac{M_t}{2} \right\}.
\]

The dimension of the \( d \)-variable function or the total number of data points in the index set is \( M = \prod_{t=0}^{d-1} M_t \). As such, the resulting FFT algorithm has a computational complexity of \( O(M \log M + \log(N/\delta)) \), where \( \delta \) is the desired computational accuracy.\cite{48}

With these preliminary definitions we can perform discrete FT for nonequispaced data. The trigonometric polynomial for \( N \) given data points can be described by

\[
f_j = f(x_j) = \sum_{k \in I_M} f_k e^{-2\pi i k x_j} \quad (j = 0, 1, ..., N-1).
\]

Using a matrix-vector notation, all trigonometric polynomials can be rewritten as \( f = A f \), where \( f = (f_j) = 0, 1, ..., N-1 \), \( A = [e^{-2\pi i k x_j}]_{j=0}^{N-1} \cdot [k \in I_M] \), and \( f = \hat{f} \). In the following implementations, the related matrix-vector products are the conjugated form

\[
\hat{f} = A^\dagger f, \quad \hat{f}_j = \sum_{k \in I_M} f_k e^{-2\pi i k x_j},
\]

and the transposed form

\[
f = A^T \hat{f}, \quad f_k = \sum_{j=0}^{N-1} f_j e^{-2\pi i k x_j},
\]

in which \( A \) and \( A^T \) are the conjugated and transposed complex of matrix \( A \), respectively. With given Fourier coefficients \( \hat{f} \), the Fourier samples \( f \) can be transformed with suitable FFT algorithms in both directions. More in-depth discussion and technical details can be found in publications\cite{20,23,24,26,48-50} and references therein.

2.3 The ENUF method

2.3.1 Implementation of the ENUF method

The ENUF method combines the standard Ewald summation method with the nonuniform FFT technique to handle electrostatic interactions between charged particles. It is noteworthy that in the ENUF method the NFFT only makes approximations for computing the long range part of electrostatic energies and forces between charged particles in reciprocal space summations,\cite{20,24,26} which is similar to the other particle-mesh based schemes in handling the long-ranged part of electrostatic interactions.\cite{10-13,17,19,51,52} Therefore, in the following subsections, we mainly present the detailed procedures to calculate reciprocal space summations of electrostatic energies and forces using NFFT.

For electrostatic energies obtained from reciprocal space summations (Equation(22)), replacing the parameter \( k \) with \( n \) in the structure factor \( S(k) \), and then normalizing particle positions \( x_a = r_a/L \), the lattice structure factor in Equation (26) is rewritten to another form \( S(n) \) as

\[
S(n) = \sum_a^n q_a e^{-i \pi n x_a} = \sum_a^n q_a e^{-i \pi n x_a} = \sum_a^n q_a e^{-2\pi m x_a} = S(n).
\]

For a fixed vector \( n \), the lattice structure factor \( S(n) \) is just a complex number.

It is clear that the structure factor \( S(n) \) in Equation (37) and the transposed FFT form in Equation (36) have similar structures. Substituting \( q_a \) with \( f_k \), the structure factor \( S(n) \) is then a 3D sample of the transformed FFT form. By viewing the structure factor \( S(n) \) as a trigonometric polynomial \( f_k \), the long range part of electrostatic energy determined from reciprocal space summations can be rewritten as

\[
U^{\text{E}}(r^2) = \frac{1}{4\pi\epsilon_0} \sum_{n=0}^{N-1} \sum_{m=0}^{M-1} \sum_{l=0}^{L-1} e^{-i (\pi n^2 r^2 / a^2)}\frac{1}{n^2} |f_n|^2.
\]

The reciprocal space summations are approximated by a linear combination of window functions sampled at nonequidistant \( M_t \) grids. These grids are used as input to transposed FFT, with which we can calculate each component of the structure factor \( S(n) \), and thereafter the reciprocal space summations of electrostatic energy. Such an approximation can effectively minimize errors initiated during interpolation and extrapolation processes, and therefore, the ENUF method is distinct to the other particle-mesh based approaches in handling the long-ranged part of electrostatic interactions.\cite{10-12,14,17,19,51}

The reciprocal space part of electrostatic force acting on charged particle \( a \) can be obtained via a similar procedure.
F_a^{E}_\text{K} = -\nabla_a U_a^{E}_\text{K} = \frac{-1}{4\pi \epsilon_0 r_c} \sum_{n \neq 0} \frac{e^{-(|m|/n)l^2}}{n^2} \frac{4\pi e \rho_n}{L} \left( -\sin \left( \frac{2\pi}{L} n r_c \right) \sum_b q_b \cos \left( \frac{2\pi}{L} n r_b \right) + \cos \left( \frac{2\pi}{L} n r_c \right) \sum_b q_b \sin \left( \frac{2\pi}{L} n r_b \right) \right)
+ \cos \left( \frac{2\pi}{L} n r_c \right) \sum_b q_b \sin \left( \frac{2\pi}{L} n r_b \right)
= \frac{1}{4\pi \epsilon_0} \frac{2q_b}{L^2} \sum_{n \neq 0} \frac{e^{-(|m|/n)l^2}}{n^2} \left( \sin \left( \frac{2\pi}{L} n r_c \right) \text{Re}(S(n)) + \cos \left( \frac{2\pi}{L} n r_c \right) \text{Im}(S(n)) \right).

(39)

Since the structure factor is a complex number, the expression in the bracket of Equation (39) can be written as the imaginary part of a product

\[\sin \left( \frac{2\pi}{L} n r_c \right) \text{Re}(S(n)) + \cos \left( \frac{2\pi}{L} n r_c \right) \text{Im}(S(n)) = \text{Im} \left\{ e^{2\pi i n r_c} S(n) \right\}.\]

(40)

Following this expression, Equation (39) can be expressed as

\[F_a^{E}_\text{K} = \frac{1}{4\pi \epsilon_0} \frac{2q_b}{L^2} \sum_{n \neq 0} \frac{e^{-(|m|/n)l^2}}{n^2} \text{Im} \left\{ e^{2\pi i n r_c} S(n) \right\}
= \frac{1}{4\pi \epsilon_0} \frac{2q_b}{L^2} \sum_{n \neq 0} \frac{e^{-(|m|/n)l^2}}{n^2} S(n) e^{2\pi i n r_c}
= \frac{1}{4\pi \epsilon_0} \frac{2q_b}{L^2} \left\{ \sum_{n \neq 0} \hat{g}_n e^{2\pi i n r_c} \right\},\]

where \(\hat{g}_n = n^{-|m|/|n|l^2} S(n)\) with \(n \neq 0\). Again, Equation (41) is a 3D sample of the conjugated FFT form (Equation (35)). Assuming \(n \in \mathbb{I}_H\) and \(\hat{g}_0 = 0\), we can reformulate Equation (41) into a series of Fourier terms

\[F_a^{E}_\text{K} = \frac{1}{4\pi \epsilon_0} \frac{2q_b}{L^2} \text{Im} \left\{ \sum_{n \in \mathbb{I}_H} \hat{g}_n e^{2\pi i n r_c} \right\} = \frac{1}{4\pi \epsilon_0} \frac{2q_b}{L^2} \text{Im} \left\{ \hat{g}_n \right\}.

(42)

Therefore, we can calculate the reciprocal space summations of electrostatic force acting on charged particle \(a\) using the conjugated FFT algorithm based on the structure factor \(S(n)\) obtained from the transposed FFT algorithm in the calculation of electrostatic energy.

### 2.3.2 Determination of optimal parameters for the ENUF method

Since the ENUF method makes an approximation on the calculation of reciprocal space summations of electrostatic energy and force, it is reasonable to expect that the ENUF method behaves in a consistent manner with the standard Ewald summation method. For the ENUF method, besides the parameters used in the standard Ewald summation method (Equation (31) and Equation (32)), there are additional two parameters from NFFT controlling the approximation errors: the oversampling factor \(\sigma_r\) and the number of terms \(m\) in spatial domain approximation.\(^{[20,21,23]}\) These two parameters are regarded as “knobs” controlling how accurately the structure factor is approximated in NFFT.

Through the calculation of electrostatic energies between charged particles using the standard Ewald summation method and the ENUF method, it is shown in Figure 1 that for a fixed oversampling factor \(\sigma_r > 1\), the relative error for electrostatic energy exhibits a significant decrease with \(m\).\(^{[20]}\) For a cutoff of \(m \geq 2\), the relative error of electrostatic energy determined using the ENUF method is consistent with that calculated using the standard Ewald summation method in comparison with the electrostatic reference energy. This indicates a strong similarity of the ENUF method with the standard Ewald summation method in handling electrostatic interactions between charged particles. Similar computational results were also observed in the relative error and the maximum relative error of electrostatic force acting on charged particle \(a\).\(^{[20]}\) It should be noted that the computational accuracy of the ENUF method can be further improved, either increasing \(m\) or \(\sigma_r\), but in the meantime, the total computational time used for handling electrostatic interactions will also be exponentially increased. Compromising the computational accuracy and the computational speed of the ENUF method, \(m = 2\) and \(\sigma_r = 2.0\) were adopted in ensuing simulations and applications to study phase behaviors of charged soft matter systems.\(^{[20,24,26]}\)

The scaling character of the ENUF method was estimated by varying the number of charged particles in simulation systems. For the physical parameters related to the standard Ewald summation method shown in Equations (31) and (32), we fix \(\delta\) and \(n_0\), and thereafter determine \(\sigma_r\) and \(r_c\) using constraints as listed in these two equations. The optimization procedure iterates from the smallest \(n_0\) permitted by the simulation system size, and stops when the resulting real space cutoff \(r_c\) is compatible with that for short-ranged interactions (such as Lennard–Jones potential). With optimal physical parameters, the ENUF method exhibits an \(O(N \log N)\) scaling behavior.

### 2.4 The ENUF-DPD method

#### 2.4.1 The DPD method

The DPD method is a particle-based approach, originally introduced by Hoogerbrugge and Koelman in 1992 as a novel scheme to simulate hydrodynamic phenomena of complex fluids at mesoscopic level.\(^{[53,54]}\) One important conceptual difference between DPD and atomistic molecular dynamics (MD) approach is the adoption of coarse-graining (CG) procedure allowing a mapping of several atoms or molecules in atomistic simulation systems onto large dissipative particles. The time evolution of dissipative particles is governed by the Newton’s equation of motion

\[\frac{\partial \mathbf{r}_a}{\partial t} = \mathbf{v}_a, \quad m_a \frac{\partial \mathbf{v}_a}{\partial t} = \mathbf{f}_a,\]

(43)

where \(\mathbf{r}_a\), \(\mathbf{v}_a\), and \(m_a\) denote the coordinate, velocity, and mass of dissipative particle \(a\), respectively. The total force \(\mathbf{f}_a\) acting on the
dissipative particle \(a\) is normally composed of three different pairwise additive forces: the conservative force \(F_{C}^{a}\), the dissipative force, and the random force \(F_{R}^{a}\).

\[
f_{a} = \sum_{b \neq a} \left( F_{C}^{ab} + F_{D}^{ab} + F_{R}^{ab} \right).
\]

with

\[
F_{C}^{ab} = \alpha_{ab} \omega_{C}(r_{ab}) \hat{r}_{ab},
\]

\[
F_{D}^{ab} = -\gamma \omega_{D}(r_{ab}) \hat{v}_{ab} \hat{r}_{ab},
\]

\[
F_{R}^{ab} = \sigma \omega_{R}(r_{ab}) \theta_{ab} \hat{r}_{ab},
\]

where \(r_{ab} = r_{a} - r_{b}\), \(r_{ab} = |r_{ab}|\), \(\hat{r}_{ab} = r_{ab}/r_{ab}\), and \(\hat{v}_{ab} = \hat{v}_{a} - \hat{v}_{b}\). The parameters \(\alpha_{ab}, \gamma,\) and \(\sigma\) determine the strength of conservative, dissipative, and random forces, respectively. \(\theta_{ab}\) is a random fluctuating variable with zero mean and unit variance.

The pairwise conservative force is usually written as a weight function \(\omega_{C}(r)\) with the form of

\[
\omega_{C}(r) = \begin{cases} 
1 & \left(1/r_{C} - |r - r_{C}| \right) \\
0 & \left(r > r_{C}\right) 
\end{cases}
\]

Compared with the Lennard–Jones 12-6 potential, the conservative force adopted in the DPD method is a soft repulsive force, and hence it allows a large time step in the integration of the equation of motion of all dissipative particles. The unit of length \(r_{C}\) is related to the size of dissipative particles and can be determined from specific CG schemes.

Two weight functions \(\omega_{D}(r)\) and \(\omega_{R}(r)\) for dissipative and random forces are coupled together via the fluctuation-dissipation theorem

\[
\omega_{D}(r) = \frac{\left(1 - |r|/r_{C}\right)}{r < r_{C}} \left(1 - |r|/r_{C}\right) \left(r > r_{C}\right)
\]

\[
\sigma_{2}^{2} = 2\gamma k_{B} T
\]

to form a thermostat and generate natural canonical distribution.\(^{[55]}\) In most applications, the weight function \(\omega_{D}(r)\) adopts a simple form as\(^{[56]}\)
density distributions meshed around dissipative particles to remove Equation (51) over the whole space gives the total charge \( \rho \) charge density distributions of Lennard-Jones interactions than soft repulsive conservative interactions. [70] Other, forming artificial ion clusters due to stronger electrostatic interactions in DPD model because dissipative particles carrying opposite point charges tend to collapse onto each other, a dimensionless parameter \( r' = r/R_c \) as the reduced center-to-center distance between two Slater-type charge density distributions and \( \beta = R_c/\lambda_e \) the reduced electrostatic energy and force are given by

\[
U_{ab}^{\text{DPD}}(r_{ab}) = \frac{1}{4\epsilon\varepsilon_0} \frac{q_a q_b}{r_{ab}^2} \left( 1 - \frac{R_{ab}}{\lambda_e} \right) e^{-2\lambda_e r_{ab}}. 
\]

(53)

The electrostatic force acting on the Slater-type charge density distribution \( \rho_e^s(r) \) is obtained by taking the negative of the derivative of electrostatic energy \( U_{ab}^{\text{DPD}} \) respect to its position \( r_a \)

\[
P_{ab}^{\text{DPD}}(r_{ab}) = -\nabla_r U_{ab}^{\text{DPD}}(r_{ab}) = \frac{1}{4\epsilon\varepsilon_0} \frac{q_a q_b}{r_{ab}^2} \left( 1 - \frac{2R_{ab}}{\lambda_e} \left( 1 + \frac{R_{ab}}{\lambda_e} \right) e^{-2\lambda_e r_{ab}} \right). 
\]

(54)

By defining a dimensionless parameter \( r' = r/R_c \) as the reduced center-to-center distance between two Slater-type charge density distributions and \( \beta = R_c/\lambda_e \) the reduced electrostatic energy and force are given by

\[
U_{ab}^{\text{DPD}}(r_{ab}) = \frac{1}{4\epsilon\varepsilon_0} \frac{q_a q_b}{r_{ab}^2} \left( 1 - (1 + \beta R_{ab}) e^{-2\lambda_e r_{ab}} \right), 
\]

(55)

\[
P_{ab}^{\text{DPD}}(r_{ab}) = \frac{1}{4\epsilon\varepsilon_0} \frac{q_a q_b}{r_{ab}^2} \left( 1 - (1 + 2\lambda_e R_{ab}) e^{-2\lambda_e r_{ab}} \right). 
\]

(56)

Now it is clear that the electrostatic energy and force between Slater-type charge density distributions in DPD simulations are those between point charges in MD simulations scaled with correction factors of

\[
B_U = 1 - (1 + \beta r') e^{-2\lambda_e r'}, 
\]

(57)

\[
B_F = 1 - (1 + 2\lambda_e r' (1 + \beta r')) e^{-2\lambda_e r'}. 
\]

(58)

Such similarities between electrostatic energies and forces in MD and DPD simulations imply that once we get electrostatic energies and forces between point charges in MD simulations, from which the electrostatic energies and forces between Slater-type charge density distributions in DPD simulations can be directly scaled with the corresponding correction factors. In the limit of \( r_{ab} \rightarrow 0 \), the reduced electrostatic energy and force between the Slater-type charge density distributions are described by

\[
\lim_{r_{ab} \rightarrow 0} U_{ab}^{\text{DPD}}(r_{ab}) = \frac{1}{4\epsilon\varepsilon_0} \frac{q_a q_b}{r_{ab}} \beta, 
\]

(59)

\[
\lim_{r_{ab} \rightarrow 0} P_{ab}^{\text{DPD}}(r_{ab}) = 0. 
\]

(60)

From Equations (59) and (60), we can specify that the adoption of Slater-type charge density distributions in DPD simulations removes the divergence of electrostatic interactions at \( r_{ab} = 0 \), indicating that both electrostatic energies and forces between Slater-type charge density distributions are finite quantities.

By matching electrostatic interactions between Slater-type charge density distributions at \( r_{ab} = 0 \) with previous work \([70]\) it gives \( \beta = 1.125 \). From the relation of \( \beta = R_c/\lambda_e \), we can get \( \lambda_e = 6.954\text{Å} \),
which is consistent with the electrostatic smearing radii used in González-Melchor’s computational model.\textsuperscript{[71]} It should be noted that as charge density distributions in simulation systems are affected by hydrodynamic flow,\textsuperscript{[77]} these proposed methods provide a natural coupling between electrostatics and fluid motion.

Figure 2 presents the reduced electrostatic energy and force between equal sign Slater-type charge density distributions and point charges. It is clearly demonstrated that at short distance \( r < 3.0R_c \) the electrostatic energy and force calculated using the ENUF-DPD method are comparable with those obtained from the Ewald summation method. In addition, both the ENUF-DPD and Ewald summation methods give indistinguishable electrostatic energy and force at \( r \geq 3.0R_c \). Therefore, the ENUF-DPD method can capture essential characteristics of electrostatic interactions as the Ewald summation method does in DPD simulations in describing phase behaviors of charged soft matter systems at extended spatiotemporal scales.\textsuperscript{[70–76,78–80]}

### 2.4.3 Determination of physical parameters for the ENUF-DPD method

The implementation of the ENUF-DPD method uses similar parameters as the ENUF method does, and the correlations between these physical parameters are described by Equations (31) and (32) with predetermined oversampling factor \( \sigma_s \) and parameter \( m \) controlling the number of terms in spatial domain approximation.\textsuperscript{[20,21,23]} However, due to the fact that \( n_c \) should be an integer and \( r_c \), which is the cutoff for the short-ranged part of electrostatic interactions between charged dissipative particles, should be a suitable value for the link-cell list update scheme in DPD simulations, we adopted another procedure to determine these parameters for the ENUF-DPD method.

First, due to the soft repulsive feature of conservative force \( F^c \) in the DPD method, we adopted \( \delta = 10^{-4} \) for the computational accuracy parameter, which is enough to keep acceptable accuracy in describing electrostatic interactions between charge density distributions in DPD simulations.\textsuperscript{[20–24,26]}

Second, we determined a suitable value for the cutoff \( r_c \) for the short-ranged part of electrostatic interactions between charged dissipative particles. In the ENUF-DPD method, as specified in Equations (55) and (56), the electrostatic energy \( U_{ab}^{\text{DPD}}(r_{ab}) \) and force \( F_{ab}^{\text{DPD}}(r_{ab}) \) are scaled with correction factors, \( B_U \) and \( B_F \), respectively, both of which are \( r \)-dependent. This indicates that the reciprocal space summations of electrostatic energy \( U_{ab}^{\text{Ewald}}(r_{ab}) \) and force \( F_{ab}^{\text{Ewald}}(r_{ab}) \) should also be scaled with the corresponding correction factors. It is noteworthy that the electrostatic energy and force we obtained from conjugated and transposed FFT algorithms are the total influence of the other charged dissipative particles on the central dissipative particle \( a \). It is difficult to differentiate their individual contributions since the corresponding correction factors for the other charged dissipative particles are related to their relative distance to the central dissipative particle \( a \). However, if we choose a suitable cutoff \( r_c \), beyond which two correction factors \( B_U \) and \( B_F \) approximate to 1.0, the total reciprocal space summations of electrostatic energy \( B_U U_{ab}^{\text{Ewald}}(r_{ab}) \) and force \( B_F F_{ab}^{\text{Ewald}}(r_{ab}) \) can be approximately expressed as \( U_{ab}^{\text{EFDP}}(r_{ab}) \) and \( F_{ab}^{\text{EFDP}}(r_{ab}) \), respectively. Such an approximation enables us to directly use conjugated and transposed FFT results as reciprocal space summations. It was found that both \( B_U \) and \( B_F \) converge to unity when \( r \geq 3.0R_c \), which is consistent with those for electrostatic energy and force between Slater-type charge density distributions shown in Figure 2. Therefore, in the ENUF-DPD scheme, \( r_c = 3.0R_c \) was taken as the cutoff for real space summations of electrostatic interactions. Such an adoption indicates that both \( B_U \) and \( B_F \) are only applied on real space summations of electrostatic interactions within a cutoff of \( r_c = 3.0R_c \).

Since the Fourier-based Ewald summation methods utilize FFT to evaluate reciprocal space summations, it is more appropriate to choose a suitable value for the Ewald convergence parameter \( \alpha \), with which we can minimize the total computational time in the calculation of electrostatic interactions from real and reciprocal space summations. The choice of \( \alpha \) is system-dependent and is related to trade-offs between accuracy and computational speed. Based on Equations (31) and (32), as well as the determined \( r_c = 3.0R_c \), one can deduce that \( \alpha = 0.12\,\text{Å}^{-1} \). In order to find a suitable value for \( \alpha \), we carried out a set of trial DPD simulations to evaluate the Madelung constant of a face-centered cubic (FCC) crystal lattice consisting of 4000 charged particles, half of which are positively charged with partial charge of +1.0 and the other half have negative charge of −1.0, respectively. It was revealed that for a wide range of \( \alpha \) values the calculated Madelung values coincide with the theoretical value.\textsuperscript{[81]} The lowest acceptable value, \( \alpha = 0.2\,\text{Å}^{-1} \), is then adopted in the subsequent DPD simulations to minimize the total computational effort.

![Figure 2](https://example.com/figure2.jpg)

**Figure 2** Electrostatic energy and force between two equal sign Slater-type charge density distributions calculated from the ENUF and Ewald summation methods. The Coulombic energy and force for two equal sign point charges, both of which diverge at \( r = 0 \), are included for a comparative purpose. Reproduced with permission from ref. [26]. Copyright 2013 Elsevier [Color figure can be viewed at wileyonlinelibrary.com]
In the last step, the parameter $n_c$ was determined together with additional two parameters ($\sigma_c$ and $m$) controlling approximation errors in NFFT. It has been shown in MD simulations that $\sigma_c = 2.0$ is adequate to provide a reliable computational accuracy in the calculation of electrostatic interactions between charged particles.[20] Therefore, this value is used in DPD simulations to keep a comparable accuracy in handling electrostatic interactions between charge density distributions at mesoscopic level.

Additional trial simulations were performed on bulk electrolyte systems to determine parameters $m$ and $n_c$. It is shown in Figure 3(a) that the relative errors of electrostatic energies and forces calculated from the ENUF-DPD and Ewald summation methods with $m = 2$ and $n_c \geq 7$ fluctuate within predetermined accuracy values, indicating that these two methods behave in a same manner in describing electrostatic interactions between Slater-type charge density distributions with adopted physical parameters. With larger $m$ and $n_c$ values, we can further increase the accuracy of electrostatic interactions in DPD simulations, which, however, leads to increased computational time in handling electrostatic interactions. By compromising the accuracy and computational efficiency of the ENUF-DPD method, $m = 2$ and $n_c = 7$ were used in following DPD simulations.

A third set of trial simulations were performed to study structural properties of dilute aqueous electrolyte solution with a salt concentration of 0.6 mol using the standard Ewald summation method with reference parameters, and the ENUF-DPD and Ewald summation methods with above determined parameters. It is shown in Figure 3(b) that pair correlation functions for the same pair particles calculated from three methods exhibit similar tendencies. A peculiar feature is that there is no ion cluster formation at $r = 0$. In addition, pair correlation functions between Slater-type charge density distributions satisfy $g_{-...} (r) g_{...+} (r) = g_{00} (r)$, where $+, -$ and $0$ correspond to positive, negative, and neutral dissipative particles in simulation systems, respectively. This indicates that microstructures of charged dissipative particles in aqueous solution are related to the effective electrostatic potentials between different particle pairs.[70]

By a systemic variation of the number of charged dissipative particles in simulation systems, we estimated the scaling behavior of the ENUF-DPD method with predetermined physical parameters. Figure 4 presents the averaged time per $10^3$ steps in DPD simulations as a function of the number of charged dissipative particles in simulation systems. It is shown that the standard Ewald summation method with reference parameters scales as $O\left(N^2\right)$, and its computational complexity is reduced to $O\left(N^{2.2}\right)$ with predetermined optimal parameters, which is consistent with that observed in previous studies.[71,72] The ENUF-DPD method with the above determined parameters exhibits an excellent computational efficiency in describing electrostatic interactions between charge density distributions at extended spatiotemporal levels, and scales as $O(N \log N)$, which is in line with the scaling behavior of FFT in treating electrostatic interactions. Theoretically, the computational complexity of NFFT is $O(M_1 \log M_1 + \log(N/\delta))$, where $M_1$ is the total number of data points in the index set, and $\delta$ is the desired computational accuracy and also a function of $m$ for fixed over-sampling factor $\sigma_c$.[21] Combining the definition of $M_1$ and the relationship in Equations (31) and (32), we can get $M_1 \propto n_c^2 \times N$. Therefore, the theoretical complexity of the ENUF-DPD method is $O(N \log N + \log(N/\delta))$, which is well reproduced from DPD simulations.

It is worth mentioning that the ENUF-DPD method has comparable computational efficiency as that of the PPPM-DPD method.

![FIGURE 3](image-url)  
(a) Relative errors of electrostatic energy ($\Delta U = \langle U_{\text{ref}} - U_{\text{Ewald}} / U_{\text{ref}} \rangle$) and force ($\Delta F = \langle F_{\text{ref}} - F_{\text{Ewald}} / F_{\text{ref}} \rangle$) calculated from the ENUF-DPD and Ewald summation methods with $m = 2$ and various $n_c$, compared with those calculated from the standard Ewald summation method with reference parameters. $U_{\text{ref}}$ and $F_{\text{ref}}$ is the total (averaged) electrostatic energy (force) on charged dissipative particles calculated from the Ewald summation method with reference parameters, and $U_{\text{Ewald}}$ is the total (averaged) electrostatic energy (force) on charged dissipative particles calculated via either the ENUF-DPD or Ewald summation methods with determined parameters. (b) Pair correlation functions between different types of dissipative particles calculated from the ENUF-DPD and Ewald summation methods with determined parameters, as well as the standard Ewald summation method with reference parameters. Reproduced with permission from ref. [26]. Copyright 2013 Elsevier [Color figure can be viewed at wileyonlinelibrary.com]
implemented by Groot\cite{70} and the EW3DC method implemented by González-Melchor et al.\cite{71} in their respective studies of charged soft matter systems. In the PPPM-DPD method,\cite{70} Groot used a linear function to spread out point charges over neighboring lattice nodes. The long-ranged part of electrostatic interactions between charge density distributions was calculated by solving the Poisson’s equation on regular lattice nodes in real space calculations using real space successive over-damped relaxations, akin to the multi-grid method.\cite{82} This method was referred to PPPM method, but is distinct to the original PPPM algorithm in which the far field part was solved using a fast Fourier transform.\cite{10} Additionally, in the EW3DC method,\cite{71} González-Melchor and coworkers considered a similar Slater-type charge density distribution on charged dissipative particles. The magnitude of electrostatic energy and force between two Slater-type charge density distributions is then given by the sum of a pairwise-additive contribution coming from real space calculations and a nonpairwise-additive contribution from the reciprocal space calculations. A z-component force was added for each charged dissipative particles to remove the net dipole moment in simulation systems. These methods have comparable computational complexities in treating long range electrostatic interactions between charged dissipative particles at mesoscopic level.\cite{26,70-72}

3 | PARALLELIZATION OF THE ENUF METHOD IN CPU AND GPU FRAMEWORKS

3.1 | Development of the CU-NFFT

It is shown in the previous section that the ENUF and ENUF-DPD methods are mainly responsible for the computation of reciprocal space summations of electrostatic energies and forces via conjugated and transposed NFFTs. More specifically, the estimation of electrostatic energies in the ENUF and ENUF-DPD methods needs a forward NFFT, and the evaluation of electrostatic forces on charged particles or charge density distributions needs three inverse NFFTs. These two processes overwhelmingly dominate the computational efficiency of NFFT. Therefore, the speedup of NFFT calculations is another procedure to improve the efficiency of the ENUF and ENUF-DPD methods in handling electrostatic interactions at extended spatiotemporal scales.

Following the work of Greengard and Lee,\cite{83} we proposed a gridding algorithm to accelerate NFFT calculations.\cite{42} The main idea of the gridding algorithm is to transform nonequispaced data points in 3D space into equivalent equispaced ones via an approximation scheme, and thereafter to accelerate NFFT calculations using standard FFT algorithms.\cite{42} The approximation scheme mapping data from nonequispaced to equispaced matrices is performed on the basis of a window function $\phi$ that is well localized both in spatial and in frequency domains, respectively. The forward NFFT for the estimation of electrostatic energies between charged particles or charge density distributions is decomposed into spreading, FFT, and scaling steps (upper panel in Figure 5), which first spreads the values of $f(x)$ from nonequispaced data points $x_i$ to equispaced and over-sampled cells using a periodic window function $\phi$, then performs forward NFFT on equispaced cells using standard FFT algorithms, and finally scales the computational FFT results to obtain $f(k)$ in a given frequency domain.

Correspondingly, the inverse NFFT for the estimation of electrostatic forces on charged particles or charge density distributions is divided into subdividing, inverse FFT, and interpolating steps, respectively (lower panel in Figure 5). In the inverse NFFT calculations, the values of $f(k)$ are first subdivided into $g(k)$ in an over-sampled cells, which are further converted to $g(x)$ in real space using the inverse FFT algorithm. Then an interpolation process is executed on the given data points $x_i$ to obtain the values of $g(x)$ using the periodic window function $\phi$. In the implementation of spreading and interpolation steps, a Gaussian window function is adopted due to the fact that its periodic version has a uniformly convergent Fourier series and this window function is also well localized both in spatial and frequency domains.\cite{42}

The NFFT and the proposed gridding algorithm have been implemented and paralleled using NVIDIA GPU via CUDA-C language and is named as CU-NFFT.\cite{42} The parallel codes of CUDA threads called kernels are designed for data-parallel processing to speed up computations using GPU. As shown in a representative diagram in Figure 6a, the kernels map data points to parallel processing threads, which are executed s times in parallel using s CUDA threads with high arithmetic intensity. In CU-NFFT, all CUDA threads are performed on physically separated devices (GPU) that operate as co-processors of the host (CPU) to run CUDA-C program. The CU-NFFT algorithm is composed of three main steps: allocating global memory in device (GPU) and loading data $f(x_i)$ from host (CPU) to device (GPU), executing kernel functions using CU-FFT, and retrieving computational results $f(k)$ from device (GPU) to host (CPU) before releasing global memory. More specifically, the spreading kernels divide the task into threads that are evenly partitioned into $s$ steaming multiprocessors (SMs) for concurrent computations (Figure 6a). Once all SMs finalize
their spreading processes, the forward CU-NFFT in CUDA library starts to conduct a standard Fourier transformation. Subsequently, all scaling kernels execute their tasks with $m$ threads in a similar manner. Correspondingly, the inverse CU-NFFT is performed in a similar manner for subdividing, inverse CU-FFT, and interpolating kernels. It is noteworthy that an advantage of the proposed CU-NFFT algorithm is that both the forward and inverse procedures are executed concurrently in GPU without interruption of transferring data between CPU and GPU so as to improve the computational efficiency of CU-NFFT.

3.2 Implementation of the CU-ENUF method

Based on the CU-NFFT algorithm, we have implemented the ENUF method in GPU, which is termed as CU-ENUF because in this method both real and reciprocal space summations are handled using GPU and CUDA technology, that is, the CU-ENUF method is an essential ENUF method paralleled with GPU and CUDA threads.[43] In real space summations, a Near-Distance algorithm was developed to effectively reduce neighbor list size so as to minimize computational time in searching interacting particle pairs within a desired cutoff, whereas
in reciprocal space calculations, the CU-NFFT replaces traditional NFFT for an efficient evaluation of electrostatic interactions among charged particles. Both real and reciprocal space summations are accelerated using GPU and CUDA technology.

In real space summations of electrostatic energies and forces among charged particles (Equations (17) and (29)), we need to sum up all contributions of interacting particles within a cutoff sphere with a radius of $r_c$. In the linked-cell list scheme, the cubic simulation box is decomposed into a regular lattice of small cells, and the side length of these cells is slightly larger than the cutoff $r_c$. Herein, we adopted a simple neighbor-list scheme to refine the size of these cells, with which one can further decrease the computational time in searching interacting particles nearby. A set of trial MD and DPD simulations indicated that in dense granular systems the adoption of an average distance between interacting particles as the side length of the decomposed cells would be an optimal balance of cell partitioning and particle counting. The proposed neighbor-list scheme embedded in linked-cell list scheme was implemented in real space summations of electrostatic interactions using the CU-ENUF method.

For the calculations of reciprocal space summations of electrostatic energies and forces between charged particles using the CU-ENUF method, the computations of structure factor $S(q)$ and $F(q)$, corresponding to a forward FFT and three inverse FFTs, are calculated using NFFT (for CPU) and accelerated using CU-NFFT (for GPU), respectively. In the CU-ENUF method, reciprocal space summations of electrostatic interactions are transformed into a parallel structure capable of running GPU acceleration based on CUDA technology. The parallel structure accepts particle charges and positions as input from CPU, and produces total electrostatic energies and forces on charged particles, the latter of which are thereafter combined with those determined from real space summations as output to CPU.

Additional art of programming were included in the CU-ENUF method to further optimize its efficiency in handling electrostatic interactions. For examples, some shared data, that is, mainly the values of window functions in CU-NFFT, are computed in local memory instead of being calculated and stored in global memory so as to avoid sacrificing efficiency when accessing to global memory of GPU. Many pre- and postprocessing computations in CPU are computed using GPU. In addition, the calculations of electrostatic forces in 3D space are packed into a single calculation entity in the inverse CU-NFFT procedure, which can reduce a large amount of execution time for initialization and redundant loops for three independent inverse NFFT calls. These improvements, together with predetermined optimal simulation parameters, render the CU-ENUF method a pure CUDA-based program and have a comparable or a better performance than the particle-mesh Ewald summation method and the ENUF method using CPU for computations.

The CU-ENUF method has been implemented as a computational module in the GALAMOST package for the computation of electrostatic interactions among charged particles in model simulation systems. Both real and reciprocal space summations are accelerated using one GPU card without any participation of CPUs. With the GALAMOST package, the CU-ENUF method can be adopted to perform CG MD and DPD simulations of charged soft matter systems at microcanonical, canonical, and isothermal-isobaric ensembles under periodic boundary conditions. In each step calculation, the GALAMOST package maximizes the amount of computations using GPU and minimizes communications between GPU and CPUs, except compulsory I/O performance. Benchmarks on representative IL systems demonstrated that the performance of the PPPM Ewald summation method appears better than the CU-ENUF method by roughly 50% for small and intermediate simulation systems with the number of ion pairs less than 0.2 million, whereas these two methods exhibit comparable computational efficiencies in handling electrostatic interactions in large simulation systems with the number of ion pairs exceeding 0.5 million. In addition, the performance of the ENUF method in the GALAMOST package is better than the PME method in the GROMACS package using one GPU and up to 28 CPU processors for the computation of electrostatic interactions in small simulation systems. However, at current stage, it is difficult to accurately quantify the computational performances of the GALAMOST and GROMACS packages for charged soft matter systems since these two packages support different features with different computational demands. Additional parallelization strategy will be explored to further improve the computational efficiency of the ENUF related methods, and the compatibility of the GALAMOST and GROMACS packages such that we can perform consistent multiscale modeling of charged soft matter systems at micro- and mesoscopic levels to explore their striking phase behaviors at extended spatiotemporal scales using GPU and CUDA technology.

### 3.3 Architecture of the HP-ENUF method

Although the CU-ENUF method achieves a qualitative leap compared with other particle-mesh based methods in handling electrostatic interactions, its computational efficiency is limited to the throughput capacity of GPU for simulation systems at extreme spatiotemporal scales. Therefore, we proposed a hybrid parallel scheme combining multiple CPU and GPU devices to upgrade the CU-ENUF method, which is described as HP-ENUF method. Similar to the CU-ENUF method, a GPU-optimized particle-data structure is employed in the HP-ENUF method so that all computations are performed using multiple GPU devices. The HP-ENUF method enables direct communications between GPU devices within different computer nodes via NVIDIA GPUDirect technology supported by CUDA-aware MPI (Message Passing Interface) library, which eliminates unnecessary data transfer between CPU and GPU devices.

The hybrid parallel scheme in the HP-ENUF method consists of multiple MPI ranks, each of which includes a CPU node and a GPU node responsible for the calculation of electrostatic energies and forces between charged particles in specific domains (subcells) in simulation systems. A schematic diagram of the hybrid parallel scheme for the HP-ENUF method is illustrated in Figure 6b. Both real and reciprocal space summations of electrostatic interactions are first parallelized via a domain decomposition scheme, which is implemented using MPI library on multiple CPU nodes, and thereafter paralleled via GPU threads in each CPU node. Using the domain decomposition scheme, each simulation task (large simulation system) is uniformly decomposed into several subtasks (small simulation subcells) that are
partitioned to different CPU nodes for CPU-parallel computation. Each CPU node is responsible for the calculation of electrostatic interactions between charged particles in its own subcell and its communication with other CPU nodes is relatively low. The MPI library is used to implement CPU parallel strategy (process level), and each MPI rank delivers subtask (the data of its own subcell) to the corresponding GPU card, and this subtask is executed in GPU-parallel computation (thread level) using CUDA technology.

For each subcell, the real space summations of electrostatic interactions in the HP-ENUF method is the same as that in the CU-ENUF method. However, unlike independent calculations of real space summations, the computations of reciprocal space summations of electrostatic interactions in each subcell depend on interactions of charged particles with those in the other subcells, indicating that CU-NFFT must be performed for charged particles in all subcells of the simulation systems. It should be mentioned that an additive property of NFFT for the calculation of structure factor $S(n)$ is that it can be decomposed into several sub-NFFTs with arbitrary particle distributions, which makes the parallel computation of CU-NFFT feasible in the HP-ENUF method. Assuming that $N$ charged particles are uniformly distributed in $n$ subcells, there are $C = N/n$ charged particles in each subcell. The total structural factor for the whole simulation system is mathematically decomposed into $n$ partial structural factors determined from $n$ subcells as

$$S(n) = \sum_{a=1}^{N} q_a e^{-2\pi n \alpha}$$

$$= \sum_{a=1}^{C} q_a e^{-2\pi n \alpha} + \sum_{a=C+1}^{2C} q_a e^{-2\pi n \alpha} + \cdots + \sum_{a=(n-1)C+1}^{NC} q_a e^{-2\pi n \alpha}$$

$$= S_1(n) + S_2(n) + \cdots + S_n(n).$$

Once all partial structural factors are calculated from different nodes, these partial structural factors will be collected and summarized in a specific computer node, and thereafter the total $S(n)$ will be broadcasted to all computer nodes to calculate electrostatic interactions between charged particles in each subcell.

All in all, as a significant extension of the CU-ENUF method, the HP-ENUF scheme successfully removes the throughput capacity of a single GPU, and is capable of conducting efficient simulations of charged soft matter systems at extended spatiotemporal scales. In addition, the HP-ENUF method is constructed with concise software architectures using C and CUDA C language, which makes it pretty transferable to other popular CUDA-accelerated packages, such as HOOMD and LAMMPS.

## 4 | APPLICATIONS OF THE ENUF RELATED METHODS IN MODELING CHARGED SOFT MATTER SYSTEMS

The ENUF method and its derivatives can capture essential characteristics of electrostatic interactions between charged particles and charge density distributions at extended spatiotemporal scales, and have been adopted to explore representative properties of charged soft matter systems, such as the effect of charge fractions of polyelectrolytes, ion concentration and counterion valency of added salts on conformational properties of polyelectrolytes, the binding structures of dendrimers on bilayer membranes and the corresponding permeation mechanisms, and the heterogeneous structures and dynamics in ILs matrices and how electrostatic interactions between charged particles affect these properties at extended spatiotemporal scales.

### 4.1 | Polyelectrolyte conformational properties

Electrostatic interactions between charged particles on polyelectrolytes lead to rich conformational properties of polyelectrolytes, which are qualitatively different from those of neutral polymers. CG MD simulations demonstrated that the size of polyelectrolytes increases with increasing the degree of ionization of polyelectrolytes, exhibiting a structural transition of polyelectrolytes from collapse (Figure 7a) to fully extended conformation (Figure 7c). These computational results are qualitatively consistent with experimental observations and theoretical predictions for weakly charged polyelectrolytes.

The intramolecular pair correlation functions between charged particles on polyelectrolytes reveal distinct tendencies with an increase of charge fraction of polyelectrolytes. The intramolecular correlations in initial zone ($r < 1$) are dominated by soft conservative repulsions. In the regime of $r > 1$, two striking tendencies are observed in simulations and shown in Figure 7g. For polyelectrolytes with a small charge fraction, a small scaling-like domain is observed and then followed by a terminal correlation range. In contrast, polyelectrolytes with a large charge fraction exhibit a scaling behavior over the entire range. These simulation results are consistent with the theoretical description of weakly charged polyelectrolytes deduced from scaling theory.

In the absence of inorganic salts, polyelectrolytes adopt extended conformations, owing to strong electrostatic repulsions between charged particles on polyelectrolytes. However, these electrostatic interactions are partially screened upon addition of salts into solution. Both ion concentration and valency of salt ions can significantly affect conformational properties of polyelectrolytes due to strong electrostatic correlations between multivalent ions and charged particles on polyelectrolytes. The condensation ability of trivalent ions on polyelectrolytes (Figure 7f) is much stronger than that of monovalent ions (Figure 7d), leading to a decrease of osmotic pressure and a conformational collapse of polyelectrolytes in solution. It is noted that a gradual increase in multivalent counterion concentration after a threshold leads to a structural transition of polyelectrolytes from fully collapsed to semiswelled conformations, which is akin to the redissolution behavior of multichain aggregates and is attributed to a competitive feature of counterions with varied charge valencies in condensing polyelectrolytes.
The effect of ion concentration and charge valency of counterions on polyelectrolyte conformations can be specified by the Debye screening length in polyelectrolyte solution. The addition of inorganic salts with multivalent counterions leads to a short Debye screening length, demonstrating that electrostatic interactions between charge particles beyond a certain distance are screened and hence are no longer long-ranged interactions. Therefore, it is very likely that a finite cutoff for electrostatic interactions, or a screened interaction potential (like the Yukawa potential) between charge particles, can be used in describing electrostatic interactions in ion-concentrated simulation systems.

4.2 | Dendrimer-lipid membrane complexes

The ENUF-DPD method were adopted to investigate specific binding structures of dendrimers on amphiphilic bilayer membranes. Polyamidoamine (PAMAM) dendrimers have hollow core and dense shell structures, and are promising nano-vehicles to protect small drug molecules during delivery process. Moreover, PAMAM dendrimers undergo conformational transitions from dense shell to dense core under external stimuli, facilitating the release of drug molecules to specific targets. When PAMAM dendrimers are used as nanodevices for nonviral gene delivery and antitumor therapeutics, the central issue is how they interact with cell membranes and how to control structures of dendrimer-membrane complexes during drug delivery.

Mutually consistent CG models of PAMAM dendrimers and dimyristoylphosphatidylcholine (DMPC) lipid molecules were constructed based on volume criteria and chemical identities. These CG models could qualitatively describe conformational properties of charged dendrimers and surface tension of atomistic DMPC lipid membranes, respectively. DPD simulation results revealed that the permeability of dendrimers across tensionless DMPC bilayer membranes is enhanced upon increasing dendrimer size. The 3rd generation (G3) PAMAM dendrimer rests on DMPC bilayer membranes without any impact on membranes, whereas G5 dendrimer spreads on bilayer membranes and attracts some lipid head groups into the vicinity of dendrimer cations. For large dendrimers, significant bending of bilayer membranes is observed due to a strong coordination of dendrimers with membranes (Figure 8b). The dendrimer-membrane contact region is characterized with high surface tension, facilitating the permeation of dendrimers across membranes.

In addition, the conformational properties and binding structures of dendrimers on membranes are influenced by the length of hydrophobic moieties of lipid amphiphiles. For membranes consisting of amphiphiles with shorter hydrophobic tails than DMPC molecule, G3 dendrimer can easily penetrate into membranes due to a small energy barrier formed by the corresponding amphiphiles. Significant binding structures of G5 dendrimer onto membranes are observed, which destroys the integrity of membranes and induce a big hole on membranes. By increasing dendrimer sizes, the dendrimer-filled vesicles (Figure 8c) are formed, which are suggested to be a possible mechanism for charged dendrimers removing amphiphiles from membranes in atomic force microscope experiments. While for dendrimers on membranes consisting of amphiphiles with longer hydrophobic moieties than DMPC molecule, small dendrimers exhibit planar conformations on membranes (Figure 8a), and large dendrimers induce small
cavities on membranes but can neither rupture membranes nor initiate isolated pores on membranes due to a large energy barrier formed by the hydrophobic moieties of corresponding amphiphiles. An illustrative phase diagram and representative binding structures of charged dendrimers on bilayer membranes are shown in Figure 8.

### 4.3 Heterogeneous structures and dynamics in ILs

Additional CG MD simulations were performed using the CU-ENUF method to address heterogeneous structures and dynamics in ILs, and how electrostatic interactions among charged particles affect these properties in IL matrices. Room temperature ILs are fascinating molten salts solely composed of ion species with distinct molecular symmetry and charge delocalization, having their melting points below 100°C. Recent years have seen a great enthusiasm for ILs regarding their utilities as facilitating functional materials in diverse applications including material synthesis and catalysis, micro-lubrication and nanotribology, gas adsorption and separation, and electrochemical devices for energy storage and harvesting. Compared with traditional molten salts, like sodium chloride, one fascinating feature of ILs is that they exhibit distinct heterogeneous microstructures and dynamics spanning multiple length and time scales in bulk region and in confined environments. Both experimental and computational characterizations revealed that mesoscopic liquid organization of ILs is characterized by either sponge-like interpenetrating polar and apolar networks or segregated polar (apolar) domains within apolar (polar) framework depending on the relative ratios of polar groups over apolar moieties in ion species. Recent atomistic simulations of ILs demonstrated that both simulation size and simulation time do matter to get reliable collective structural and dynamical quantities of ions in IL matrices.

As simulations of ILs should be performed over long time scales due to sluggish dynamics of ion species in heterogeneous IL matrices, it imposes severe fundamental challenges for atomistic simulations to accurately predict dynamics and transport properties of ILs. In this regard, we performed extensive CG MD simulations with a modest computational cost to explore the effects of neutral chain length in cations, molecular sizes of anions, and temperatures on microstructural and dynamical quantities of ion groups in 1-alkyl-3-methylimidazolium tetrafluoroborate ([CnMIM][BF4]) ILs at extended spatiotemporal scales. It was found that lengthening cation alkyl chains leads to an aggregation of neutral beads, promotes the formation of spatially heterogeneous apolar domains dispersed in ionic channels, and thereafter results in a remarkable transition of mesoscopic liquid morphologies in model IL systems from dispersed neutral (apolar) beads in a 3D framework of ion channels to that characterized by bi-continuous interpenetrating polar and apolar networks in liquid matrices. Such a microstructural evolution in model IL matrices can be rationalized by a competition of short range collective interactions between neutral beads and long range electrostatic interactions among charged particles in constituent ions.

For dynamical quantities, translational diffusion of ion groups presents a gradual decrease upon lengthening cation alkyl chains and enlarging molecular sizes of anions. The temperature dependence of diffusion coefficients of all representative groups in ion models is...
described by a classical Arrhenius feature. The rotational dynamics of cations in varied IL matrices are characterized by a bi-exponential structural relaxation behavior. The correlation times for these two rotational modes are significantly temperature dependent, and are strongly related to cation structures, indicating a rotational heterogeneity of charged and neutral moieties of cations in heterogeneous ionic environments.

Additional CG MD simulations were carried out on neutral analogues of model [C10MIM][BF4] IL (Figure 9a) in which all electrostatic interactions between charged particles were switched off. The removal of electrostatic interactions between ion models tends to loosen liquid structures leading to an almost homogeneous and interleaved distribution of neutral ion models resembling a distorted ion lattice in simulation systems.[8,111,113] This visual microstructural change is manifested in detailed characterizations, such as liquid densities, radial distribution functions, and translational dynamics of representative beads in neutral and ion models, as clearly shown in Figure 9. These observations highlight a critical role of electrostatic interactions in describing collective structural and dynamical quantities of model IL systems.

5 | CONCLUDING REMARKS AND OUTLOOK

Computer simulations provide a unique insight into molecular interactions and structural and dynamical quantities responsible for many peculiar properties of materials and biological systems at multiple spatiotemporal scales. The quality of computer simulations is essentially determined by an accurate description of intra- and intermolecular interactions in these molecular systems, among which the electrostatic interactions between charged particles deserve special attention because of these long-ranged interactions play a prominent role in determining structures and states of simulation systems. The efficient calculations of electrostatic interactions in model systems subjected to partial or full periodic boundary conditions have been a daunting task. A wide variety of theoretical approaches, ranging from quantum mechanical ab initio methods, classic Maxwell theory of electromagnetism, generalized Born algorithms, to phenomenological modifications of Coulomb's law, and computational methods, including particle-mesh based methods and multi-poles/grids approaches, have been developed for electrostatic analysis at different resolution levels.[1,2]

The standard Ewald summation method does a remarkable job in splitting the very slowly converged Coulomb potential into two parts converging fast exponentially, which makes the calculations of electrostatic interactions in computer simulations feasible, and therefore, this method has been widely used to handle long range electrostatic interactions in modeling charged simulation systems. However, the standard Ewald summation method suffers from its computational demanding feature due to the long range nature of electrostatic interactions. Although plenty of methods have been proposed with different computational schemes to tackle long range interactions in simulation systems with partial or full periodic boundary conditions, only a certain subset of them has entered into widely used MD codes for scientific computing with varied computational efficiencies. This
fact is related to the long standing and continuously improved implementations of selected methods and also the large effort needed to propose and implement new approaches.

As an attractive alternative approach to particle-mesh based schemes, we sketched an ENUF method, which has been implemented in particle-based simulation packages to speedup calculations of electrostatic energies and forces between charged particles at micro- and mesoscopic levels without resorting to a nonphysical truncation of the Coulomb potential. The ENUF method and its derivatives conserve both energy and momentum to floating point accuracy, capture essential characteristics of electrostatic interactions, and exhibit an outstanding computational complexity of $O(N \log N)$ with optimal physical parameters. The ENUF method are further adopted to investigate the dependence of conformational properties of polyelectrolytes on charge fraction of polyelectrolyte and counterion valency and concentration of added salts, and the effect of dendrimer sizes and variation of hydrophobic tails of lipid molecules on specific binding structures of dendrimers on amphiphilic membranes.

Taking advantages of GPU and CUDA technology, we have upgraded NFFT and the ENUF method to their GPU versions, which are termed as CU-NFFT and CU-ENUF, respectively, and are specialized to improve computational efficiencies in computations of electrostatic interactions using the GALAMOST package. In addition, several (hybrid) parallelization strategies based on gridding and Near-Distance algorithms were developed to effectively partition hardware memories and balance computational loads between CPU and (multi) GPU nodes. In addition, the CU-ENUF and HP-ENUF methods are developed using C and CUDA C language and are constructed with concise software architecture, which render them having significant transferabilities to other popular CUDA-enabled packages, such as GROMACS, LAMMPS, and HOOMD. It is expected that the ENUF related methods will be used in varied computational communities and updated by researchers to expand their visibilities and applications in handling electrostatic interactions between charged particles at extended spatiotemporal scales.

ACKNOWLEDGMENT

We thank Prof. Zhong-Yuan Lu (Jilin University, China) for insightful discussion. S.-C. Yang acknowledges financial support from Science and Technology Research Project Fund from Education Department of Jilin Province, China (JJKH20190696KJ), and from State Key Laboratory of Supramolecular Structure and Materials of Jilin University (SKLSSM202031). B. Li acknowledges financial support from the Fundamental Research Funds for the Central Universities (No. 20lgpy76). Y.-L. Zhu acknowledges financial support from the National Natural Science Foundation of China (21774129). A. Laaksonen acknowledges Swedish Science Council for financial support (2019-03865), and partial support from a grant from Ministry of Research and Innovation of Romania (CNCS - UEFISCDI, project number PN-III-P4-ID-PCCF-2016-0050, within PNCDI III). Y.-L. Wang gratefully acknowledges financial support from Knut och Alice Wallenbergs Stiftelse (KAW 2018.0380).

ORCID

Yong-Lei Wang https://orcid.org/0000-0003-3393-7257

REFERENCES

[1] M. P. Allen, D. J. Tildesley, *Computer Simulation of Liquids*, Oxford University Press, New York, NY 1989.
[2] D. Frenkel, B. Smit, *Understanding Molecular Simulation: From Algorithms to Applications*, Academic Press, New York 1996.
[3] K. Sharp, B. Honig, *Annu. Rev. Biophys. Chem.* 1990, 19, 301.
[4] C. Holm, P. Kézích, B. Podgornik, *Electrostatic Effects in Soft Matter and Biophysics*, Springer, Heidelberg 2001.
[5] A. Naji, S. Jungblut, A. G. Moreira, R. R. Netz, *Phys. A* 2005, 352, 131.
[6] S. K. Sainis, J. W. Merrill, E. R. Dufresne, *Langmuir* 2008, 24, 13334.
[7] M. N. Kobrak, H. Li, *Phys. Chem. Chem. Phys.* 2010, 12, 1922.
[8] Y.-L. Wang, Y.-L. Zhu, Z.-Y. Lu, A. Laaksonen, *Soft Matter* 2018, 14, 4252.
[9] P. Ewald, *Annales der Physik* 1921, 369, 253.
[10] J. W. Eastwood and R. W. Hockney, *New York: Mc GrawHill* (1981).
[11] M. Deserno, C. Holm, *J. Chem. Phys.* 1998, 109, 7694.
[12] W. M. Brown, A. Kohlmeyer, S. J. Plimpton, A. N. Tharrington, *Comput. Phys. Commun.* 2012, 183, 449.
[13] T. Darden, D. York, L. Pedersen, *J. Chem. Phys.* 1993, 98, 10089.
[14] C. F. Batcho, D. A. Case, T. Schlick, *J. Chem. Phys.* 2001, 115, 4003.
[15] U. Essmann, L. Perera, M. L. Berkowitz, T. Darden, H. Lee, L. G. Pedersen, *J. Chem. Phys.* 1995, 103, 8577.
[16] Z.-H. Duan, R. Krasny, *J. Chem. Phys.* 2000, 113, 3492.
[17] Y. Shan, J. L. Klepeis, M. P. Eastwood, R. O. Dror, D. E. Shaw, *J. Chem. Phys.* 2005, 122, 054101.
[18] M. Harvey, G. De Fabritiis, *J. Chem. Theory Comput.* 2009, 5, 2371.
[19] H. Wang, F. Dommett, C. Holm, *J. Chem. Phys.* 2010, 133, 034117.
[20] F. Hedman, A. Laaksonen, *Chem. Phys. Lett.* 2006, 425, 142.
[21] A. Dutt, V. Rokhlin, *SIAM J. Sci. Stat. Comput.* 1993, 14, 1368.
[22] A. Dutt, V. Rokhlin, *Appl. Comput. Harmon. Analysis* 1995, 2, 85.
[23] R. Weeber, F. Nester, F. Weik, M. Pippig, D. Potts, C. Holm, *J. Comput. Phys.* 2019, 391, 243.
[24] Y.-L. Wang, F. Hedman, M. Porcu, F. Mocci, A. Laaksonen, *Appl. Math.* 2014, 5, 520.
[25] Y.-L. Wang, Ph.D. thesis, Department of Materials and Environmental Chemistry, Stockholm University (2013).
[26] Y.-L. Wang, A. Laaksonen, Z.-Y. Lu, *J. Comput. Phys.* 2013, 235, 666.
[27] Y.-L. Wang, Z.-Y. Lu, A. Laaksonen, *Phys. Chem. Chem. Phys.* 2012, 14, 8348.
[28] D. E. Shaw, M. M. DePeters, R. O. Dror, J. S. Kuskin, R. H. Larson, J. K. Salmon, C. Young, B. Batson, K. J. Bowers, and J.C. Chao, ACM SIGARCH Comp Arch News 2007, 35, 1.
[29] D. Kirk, *ISMM 2007*, 7, 103.
[30] W. Liu, B. Schmidt, G. Voss, W. Müller-Wittig, *International Conference on High-Performance Computing*, Springer, Heidelberg 2007, p. 185.
[31] J. S. Vetter, R. Glassbrook, J. Dongarra, S. McNally, J. Meredith, J. Rogers, P. Roth, K. Spafford, S. Yamamoto, *Comp. Sci. Eng.* 2011, 13, 90.
[32] A. L. Blumers, Y.-H. Tang, Z.-Y. Lu, A. Laaksonen, *Soft Matter* 2010, 6, 1922.
[33] A. Deserno, C. Holm, *J. Mol. Graphics* 2005, 23, 520.
[34] W. M. Brown, S. Hampton, P. Agarwal, P. Wang, P. Crozier, and S. Plimpton, *Sandia National Laboratories, Tech. Rep* (2010).
[35] J. E. Stone, D. J. Hardy, I. S. Ufimtsev, K. Schulten, *J. Mol. Graphics Modell.* 2010, 29, 116.
