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Compositional transferability of deep potential in molten LiF–BeF₂ and LaF₃ mixtures: prediction of density, viscosity, and local structure†

Xuejiao Li, *^{ab} Tingrui Xu^{ab} and Yu Gong *^{ab}

The accumulation of lanthanide fission products carries the risk of altering the structure and properties of the nuclear fuel carrier salt LiF–BeF₂ (Flibe), thereby downgrading the operating efficiency and safety of the molten salt reactor. However, the condition-limited experimental measurements, spatiotemporal-limited first-principles calculations, and accuracy-limited classical dynamic simulations are unable to capture the precise local structure and reliable thermophysical properties of heterogeneous molten salts. Therefore, the deep potential (DP) of LaF₃ and Flibe molten mixtures is developed here, and DP molecular dynamics simulations are performed to systemically study the densities, diffusion coefficients, viscosities, radial distribution functions and coordination numbers of multiple molten Flibe + xLaF₃, the quantitative relationships between these properties and LaF₃ concentration are investigated, and the potential structure–property relationships are analyzed. Eventually, the transferability of DP on molten Flibe + LaF₃ with different formulations as well as the predictability of structures and properties are achieved at the nanometer spatial scale and the nanosecond timescale.

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1. Introduction

The molten LiF–BeF₂ (Flibe) eutectic salt has been considered as one of the most promising fuel carriers and coolants for molten salt reactors (MSRs), due to its lower neutron absorption, high thermal conductivity, large heat capacity, appropriate viscosity, and good tolerance for actinides or fission products.^{1,2} The thermodynamic properties and structural characteristics of molten Flibe have been extensively investigated by experimental and theoretical methods.^{3–11} It is suggested that the thermophysical properties of molten fluorides are significantly affected by impurities,¹² so the impurity content should be strictly controlled in the application of MSRs. Lanthanide fluorides (LnF₃), as main nuclear fission products,¹³ dissolved in molten Flibe, may affect its thermophysical properties, thereby affecting the thermal energy storage and transfer performance of carrier salts. Therefore, studying the thermodynamic properties of molten Flibe + LnF₃ mixtures is critical to the design and development of MSRs, and the transport behavior of lanthanide products in molten Flibe is also essential to the daily operation of MSRs and the pyroprocessing of

spent fuels.¹⁴ Experimental measurements of the structure and properties of Flibe + LnF₃ molten salt are challenged by the high temperatures, toxicity, radioactivity, reactivity, corrosiveness, and data uncertainty. Particularly, the structural characteristics of multivalent cations are hardly accessed due to highly dynamic disorder and heterogeneity in a coordination environment.¹⁵

The molecular dynamics simulation technique has been developed rapidly as an alternative or auxiliary technology for experiments, making it possible to predict the macroscopic properties while revealing the microscopic structural information of molten salts.^{16,17} Classical molecular dynamics simulations based on empirical force field models have been applied to predict the thermodynamic properties of molten fluorides,^{18–23} whereas the accuracy depends on the expressive power of interionic potentials. In contrast, first-principles molecular dynamics (FPMD) simulations calculate the interionic forces on-the-fly and can achieve responses on the predictions of local structures and partial properties of molten salts with higher precision.^{24–29} However, the computationally intensive performance makes a typical FPMD simulation prohibitively expensive to run with larger, more complex, and longer times. Besides, the property calculations of viscosity, electrical and thermal conductivity are time-consuming, and the simulation box should be enlarged if the concentration range lanthanide fluoride is widened.^{30,31} In the age of artificial intelligence, several MD methods based on the machine-learning (ML)

^a Shanghai Institute of Applied Physics, Chinese Academy of Sciences, Shanghai 201800, China. E-mail: lixuejiao@sinap.ac.cn, gongyu@sinap.ac.cn

^b University of Chinese Academy of Sciences, Beijing 100049, China

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interatomic potentials have emerged and have been applied in molten salt systems.^{32–35} Among them, deep potential molecular dynamics (DPMD) eliminate the dilemma of accuracy *versus* spatiotemporal scales by using a sufficient function for describing the interionic interactions in a system.³⁶ At present, the DPMD simulations have been successfully performed on molten chlorides^{37–39} and fluorides.^{40,41}

In this work, LaF₃ as a representative lanthanide fluoride is added into molten Flibe eutectic salt to investigate the macroscopic properties and microscopic structure of molten mixtures. Firstly, diverse FPMD simulations of molten Flibe + LaF₃ are conducted at 823 K with small simulation cells under the complete consideration of the position of La. Secondly, the deep potential of molten Flibe + LaF₃ is trained, verified, and tested at the same temperature by evaluating the root-mean-square errors (RMSEs) of energy and force. Based on the optimized deep potential, the DPMD simulations of multiple molten Flibe + xLaF₃ are performed with an enlarged simulation cell and an extended simulation time to verify the transferability of the deep potential over wider concentration ranges. Thirdly, the densities, diffusion coefficients, and viscosities of molten Flibe + LaF₃ as well as the radial distribution functions and coordination numbers are predicted. Furthermore, the concentration dependencies of structures and properties for molten Flibe + LaF₃ are obtained. In summary, the development of a transferable deep potential model for Flibe + LaF₃ fulfills the prediction of structural and thermophysical properties, which also provides insights into the understanding of the physiochemical behavior of lanthanide fission products in molten Flibe.

2. Computational methods

2.1. FPMD simulations

The pure rhombohedral Li₂BeF₄ crystal ($a = b = c = 8.375 \text{ \AA}$, $\alpha = \beta = \gamma = 107.8^\circ$) composed of 12 LiF and 6 BeF₂ (total 42 atoms) is modelled and heated to 823 K.⁴² Based on the stable configuration, the initial configuration of molten Flibe + LaF₃ is built by adding one LaF₃. The FPMD simulation is performed based on the density functional theory and Born–Oppenheimer approximation as implemented in the Vienna *ab initio* simulation package (VASP).⁴³ The generalized gradient approximation (GGA) with the Perdew–Burke–Ernzerhof (PBE) exchange–correlation functional is used for simulations, and core electrons are approximated by pseudopotentials developed using the projector augmented wave method.⁴⁴ The wave functions for valence electrons, F(2s²2p⁵), Li(1s²2s¹), Be(1s²2s²), and La (5s²5p⁶5d¹6s²) are expanded using the plane wave basis set with a cutoff energy of 650 eV. A $1 \times 1 \times 1$ *k*-point mesh is chosen in FPMD simulations, which is a suitable compromise between computational accuracy and cost.⁴⁵ At first, a short-time FPMD simulation within the NVT ensemble *via* a Nosé thermostat is conducted aiming to bring the Flibe + LaF₃ system to the specified temperature. Then, 30 000 steps of simulations are carried out in the NPT ensemble with a time

step of 1 fs using the Langevin thermostat to achieve preliminary equilibrium, and the cell volume and lattice parameters of the molten mixture are optimized by using the method of Parrinello and Rahman. Finally, the equilibrium configuration is used as an input of the following simulations, and a 30 ps FPMD simulation of molten Flibe + LaF₃ (abbreviated as La1) are implemented within the NVT ensemble with the same time step. According to the last NVT configuration, the positions of Li or Be are manually exchanged with La to overcome the possible dynamic energy barriers while enriching sampling configurations. These new configurations are named by the number of Li (1–12) and Be (1–6), which are simulated by repeating the above NPT–NVT process, respectively.

2.2. Development of the deep potential

For establishing the deep potential (DP) of molten Flibe + LaF₃ for DPMD simulations, the datasets are adopted from the above FPMD results. All the FPMD phase–space trajectories simulated under both NVT and NPT ensembles are collected in the datasets (total 62 700), which are shuffled and split by about 2 : 1 : 1 as training, validation, and testing datasets for machine learning as shown in Fig. 1. The DeepMD-kit (version 2.0.1) package, a scheme for neural network training and evaluation of atomic energies and forces, is used to train the DP for molten Flibe + LaF₃.⁴⁶ The deep potential smooth edition (DeepPot-SE) is selected, which incorporates comprehensive atomic configuration information, and the hard cutoff radius (r_{cut}) is set to be 7.5 Å. The selected number of atoms in r_{cut} is {46, 46, 46, 46}, and four hidden layers {25, 50, 100, 200} are included to construct the residual networks architecture between them. The fitting net of {240, 240, 240, 240} is selected, which takes descriptors as inputs and predicts the atomic contribution to target properties.⁴⁷ And, the decay rate and step are set as 0.95

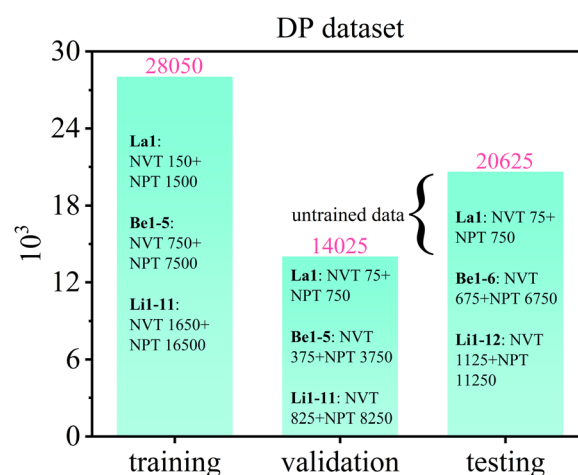


Fig. 1 Training (28 050), validation (14 025), and testing (20 625) datasets of molten Flibe + LaF₃ (La1, Be1–6, and Li1–12 represent 19 different initial configurations, for example, 'La1: NVT 150 + NPT 1500' represent 150 and 1500 samples from NVT and NPT simulations based on the initial La1 configuration, respectively, and the untrained data includes 6600 NVT and NPT data for Be6 and Li12).



and 5000, respectively. Meanwhile, the prefactors in the loss function are chosen as $\text{pref}_e^{\text{start/limit}} = 0.02/1$ and $\text{pref}_f^{\text{start/limit}} = 1000/1$, and no virial is incorporated in both training and testing stages based on previous study.³⁹

2.3. DPMD simulations

The trained DP can be directly used in the large scale atomic/molecular massively parallel simulator (LAMMPS) *via* the interface of the DeePMD-kit.⁴⁸ A larger 8050-atom box of Flibe + LaF₃ (La175:175 La, 1050 Be, 2100 Li, and 4725 F) is simulated, corresponding to the composition of the above FPMD simulations, and the simulation temperature is also set to be 823 K, higher than the inlet temperature of MSR. In DPMD simulations, the system is first relaxed for 1 ns under NVT ensemble using the Nose–Hoover thermostat. Then, another 2 ns simulation under the NPT ensemble is carried out, from which the average volume is obtained to calculate the equilibrium density. As for obtaining the long-term related viscosity properties, the system is simulated within the NVT ensemble at the target temperature by a production run of 5 ns to achieve convergence of the stress tensor, and the ionic self-diffusion coefficients and radial distribution functions of key ionic pairs are also analyzed according to the NVT simulations. Furthermore, the concentration dependences of these properties and structures are accessed by executing repetitive simulations of Flibe + LaF₃ systems with different LaF₃ concentrations (see Table 1), referring experimental dosages in ref. 49. For all simulations, the periodic boundary conditions are employed, and Newton's equation of motion is solved with a time step of 1 fs using the Verlet algorithm.

3. Results and discussion

3.1. Performance of DP

3.1.1. RMSEs of energy and force. The RMSEs of energy and force evaluated from validation datasets of molten Flibe + 5.26 mol% LaF₃ (La175) after the training stage are depicted in Fig. S1 of the ESI,[†] where the standard deviations of RMSEs of energy and force in the last 5000 training steps are less than 1%. It is suggested the continuous stabilization and equilibrium of the DP training has been achieved, and 1000 000 training steps are enough to reach convergence criteria.³⁷ At the testing stage, the trained DP is further examined by FPMD results as shown in Fig. 2, where all the ratios of energy and

force between FPMD and DPMD are close to unity, confirming the excellent accuracy of this DP for reproducing energy and force of molten Flibe + LaF₃. A new dataset with the consideration of trained and untrained trajectories is separately used as testing dataset and yields weighted RMSEs of energy (2.0 meV per atoms) and force (3.6×10^{-2} eV Å⁻¹) for molten Flibe + LaF₃. The deviations of RMSEs between validation and testing datasets are within the acceptable range, suggesting no occurrence of overfitting in the DP training.⁵⁰

3.1.2. RDFs and MSDs. The radial distribution functions (RDFs) of molten Flibe + LaF₃ (5.26 mol%) from DPMD simulations are compared with those of FPMD results, where the RDF curves of FPMD are evaluated from the average result of 19 different initial configurations as discussed in Section 2.1. In Fig. 3, the RDF curves of DPMD are obviously smoother due to the larger simulation cell, making it easy to identify peak positions and coordination numbers. Besides, all the DPMD RDFs are higher than those of FPMD due to the more particle numbers. The first peak positions (R_1) and corresponding coordination numbers (CN_1) of Be–F, Li–F and F–F from DPMD agree well with those from FPMD simulations, indicating that the robust DP is able to accurately predict the local structures of molten Flibe + LaF₃. From Fig. 4, all the mean square displacements (MSDs) at 823 K follow upward trends as time increases; therefore, the diffusion coefficient (D) from both FPMD (30 ps) and DPMD (5 ns) can be derived from the slope of MSDs with respect to time based on the Einstein equation. Both the orders of D from FPMD and DPMD simulations are $\text{Li}^+ > \text{F}^- > \text{Be}^{2+} > \text{La}^{3+}$, and the D values for the same ion are in the same order of magnitudes, consistent with the corresponding experimental and simulated results of pure molten Flibe.^{11,51} However, the D_{Be} and D_{F} are closer in large-scaled DPMD simulations, agreeing with the Maxwell–Stefan diffusivities,⁹ which is evident from the strong Be–F interaction. Overall, the reasonable results demonstrate the DP is well-represented and with expected accuracy comparable to those of FPMD results.

3.1.3. Densities and potential energies. The density and potential energy per atom of molten Flibe + LaF₃ at 823 K evaluated from DPMD simulations are displayed in Fig. 5. It is clear that the density linearly increases from 2.070 to 2.518 g cm⁻³ as the LaF₃ concentration increases from 0.19 to 5.26 mol%, and all densities of molten Flibe + LaF₃ are larger than that of pure molten Flibe (2.05 g cm⁻³).⁵² The similar phenomena also occur in the molten LiF–CaF₂–LaF₃ system, where the density increases from 2.071 to 2.423 g cm⁻³ at 1123 K as the LaF₃ concentration increases from 0 to 5 mol%;⁵³ but there is currently no comparable experimental density for molten Flibe + LaF₃. It is indicated that the mass increment of molten Flibe caused by lanthanide fission products is larger than volume increment. Besides, the linear relationship between density and LaF₃ concentration (c_{La}) can be expressed as shown in eqn (1), which is in excellent agreement with above experimental value of pure Flibe.

$$\rho_{\text{DP}} = 2.05 + 0.089c_{\text{La}} \text{ (mol\%)} \quad (1)$$

Table 1 Composition profile of multiple Flibe + LaF₃ systems (the system is abbreviated as Lax, x represents the number of La ions, mol% and wt% represents the mole and mass fraction of LaF₃, respectively, BeF₂, LiF, and Total represent the corresponding particle numbers)

System	mol%	wt%	LaF ₃	BeF ₂	LiF	Total
La6	0.19	1.12	6	1050	2100	7374
La18	0.57	3.28	18	1050	2100	7422
La36	1.13	6.36	36	1050	2100	7494
La76	2.36	12.54	76	1050	2100	7654
La175	5.26	24.84	175	1050	2100	8050



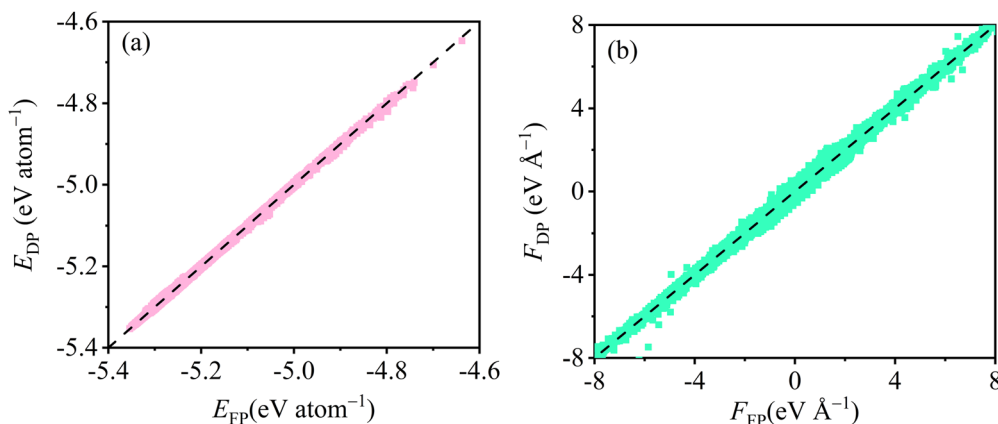


Fig. 2 DPMD (simplified to subscript DP) and FPMD (simplified to subscript FP) predicted (a) energy E and (b) force F for molten Flibe + LaF₃ (5.26 mol%).

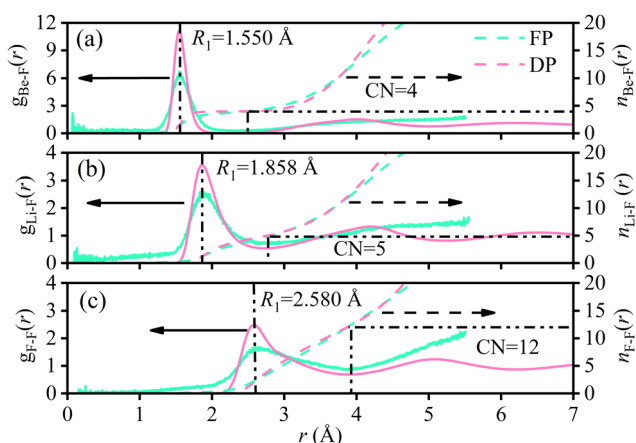


Fig. 3 FPMD (color code: light green) and DPMD (color code: light pink) simulated RDFs for (a) Be-F, (b) Li-F, and (c) F-F of molten Flibe + LaF₃ (5.26 mol%).

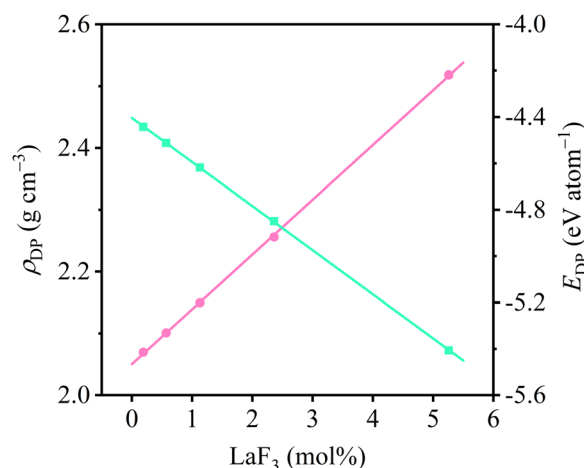


Fig. 5 DPMD predicted densities (color code: light pink) and potential energies per atom (color code: light green) of molten Flibe + LaF₃.

The potential energy per atom (E_{DP}) of molten Flibe + LaF₃ also linearly decreases from -4.4 to -5.4 eV per atom with the rising LaF₃ concentration, attributed to the decreased total

potential energy and increased atom numbers. The E_{DP-cLa} relationship can be obtained as follows,

$$E_{DP} = -4.40 - 0.19c_{La} \text{ (mol\%)} \quad (2)$$

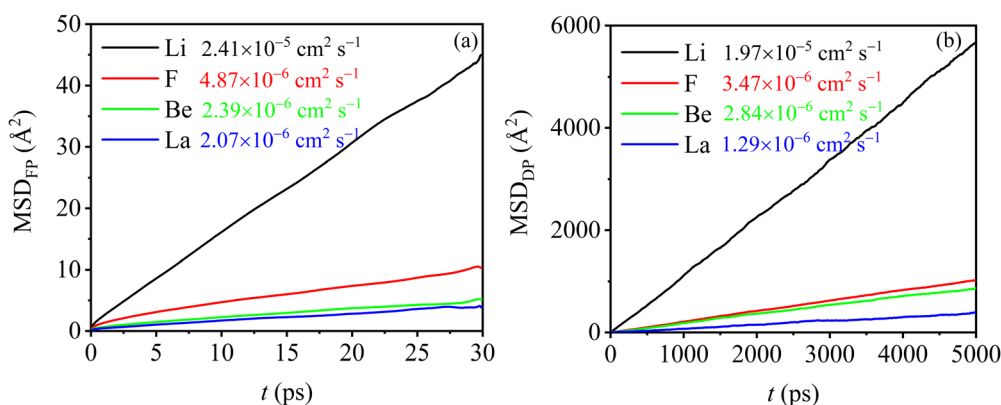


Fig. 4 (a) FPMD and (b) DPMD simulated MSDs of molten Flibe + LaF₃ (5.26 mol%) where the data after each ion are the corresponding ionic self-diffusion coefficient.



It is thus concluded that the DP transferability of molten Flibe + LaF₃ across different salt compositions satisfies the requirement for the accurate prediction of density and interaction energy. A molecular-level interpretation of the changes in properties with LaF₃ concentration will be discussed in Section 3.3.1.

3.2. Transport properties

3.2.1. Self-diffusion coefficients. As shown in Fig. 6, the ionic self-diffusion coefficients of multiple molten Flibe + LaF₃ systems are estimated from the MSD results (see Fig. S2, ESI[†]). Obviously, the D_{Li} is the largest due to its smallest ionic radius and molecular mass, and D_{Li} is more than 5 times that of D_{F} , hinting that the diffusivity of molten Flibe + LaF₃ mainly depends on Li ions. The D_{F} is similar to D_{Be} but slightly larger than D_{Be} , and the strong Be-F interaction leads to their synergistic movement, which will be further discussed in Section 3.3.2. Besides, the D_{Li} , D_{F} , and D_{Be} are basically decreased with the increasing LaF₃ concentration. The D_{La} is the lowest due to its highest mass among four ions, and significantly decreases from 4.0 to $1.3 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$, agree with the change in the trend of D_{La} in molten FLiNaK, where D_{La} is decreased with increasing concentration when the temperature is below 973 K.⁵⁴ Regrettably, no experimental D_{La} in molten FLiBe is available for direct comparison, and the diffusion behavior of lanthanide ions in similar molten salt systems can be used as references. The simulated diffusion coefficient of La³⁺ in molten Flibe at 823 K is consistent with that of Sm³⁺ ($3.0 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$) in molten Flibe,⁵⁵ lower than that of Eu³⁺ ($1.2 \times 10^{-5} \text{ cm}^2 \text{ s}^{-1}$) in molten Flibe,⁵⁶ but higher than La³⁺ in the FLiNaK molten salt ($1.8 \times 10^{-7} \text{ cm}^2 \text{ s}^{-1}$).⁵⁷ Notably, the repulsion force between La ions is not included in the trained DP because only one La in each FPMD model (see Section 2.1) and the La-La interaction is not involved, which may lead to underestimation of D_{La} and this phenomenon is more prominent in higher concentration LaF₃ systems.

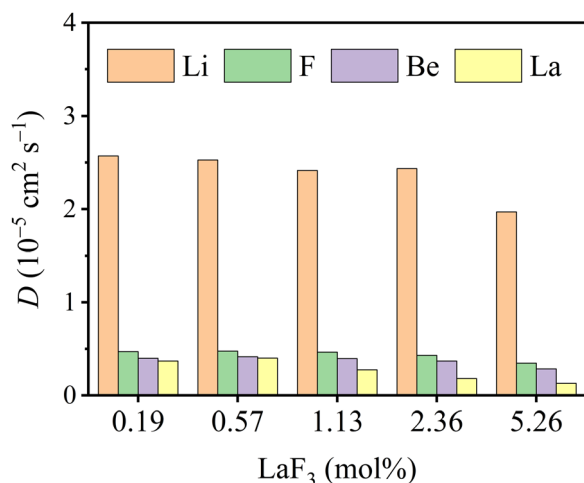


Fig. 6 DPMD predicted ionic self-diffusion coefficients (D) vs. LaF₃ concentrations of molten Flibe + LaF₃.

Table 2 Weighted self-diffusion coefficients (\bar{D} , $10^{-6} \text{ cm}^2 \text{ s}^{-1}$), equilibrium volumes (V , Å³), viscosities (η , cP), nearest neighbor distances (Å) of Be–Be ($R_1^{\text{Be-Be}}$) and La–F ($R_1^{\text{La-F}}$), and coordination numbers of La–F ($\text{CN}_1^{\text{La-F}}$) for molten Flibe + LaF₃ systems

System	mol%	\bar{D}	V	η	$R_1^{\text{Be-Be}}$	$R_1^{\text{La-F}}$	$\text{CN}_1^{\text{La-F}}$
La6	0.19	10.58	84 250	9.56	2.930	2.362	8.41
La18	0.57	10.48	84 854	9.58	2.930	2.328	8.51
La36	1.13	10.01	85 657	10.80	2.930	2.327	8.63
La76	2.36	9.69	87 383	11.66	2.950	2.320	8.82
La175	5.26	7.57	91 073	15.94	2.950	2.292	8.99

In addition, the weighted diffusion coefficient (\bar{D}) of molten Flibe + LaF₃ is calculated as,

$$\bar{D} = \frac{N_{\text{Li}}D_{\text{Li}} + N_{\text{F}}D_{\text{F}} + N_{\text{Be}}D_{\text{Be}} + N_{\text{La}}D_{\text{La}}}{N_{\text{tot}}} \quad (3)$$

and the values are listed in Table 2. The \bar{D} declines as the LaF₃ concentration increases, and the linear relationship is as follows,

$$\bar{D}(10^{-6} \text{ cm}^2 \text{ s}^{-1}) = 10.79 - 0.59c_{\text{La}} \quad (4)$$

The fitted index R^2 of \bar{D} and c_{La} is good, about 0.98, meaning that the eqn (4) can be used to predict the weighted diffusion coefficient of molten Flibe + LaF₃ with any LaF₃ composition.

3.2.2. Shear viscosities. The shear viscosity (η) of molten Flibe + LaF₃ is calculated *via* the Green–Kubo (GK) method related to the auto-correlation function of off-diagonal components of the stress tensor (SACF),³¹

$$\lambda = \frac{V}{k_{\text{B}}T} \int_0^\infty \langle S_{\alpha\beta}(t)S_{\alpha\beta}(t_0) \rangle dt \quad (5)$$

where V is the equilibrium volume of simulation model, k_{B} the Boltzmann constant, $S_{\alpha\beta}$ the stress tensor, the angle bracket denotes the average over all time origins t_0 , and t is the delay time of the SACF. Fig. 7 shows shear viscosities of multiple molten Flibe + LaF₃ over time at 823 K, which increase rapidly

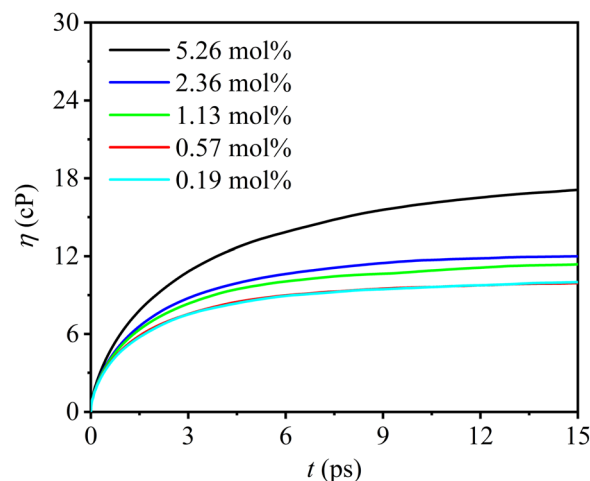


Fig. 7 DPMD predicted shear viscosities (η) of molten Flibe + LaF₃ where t is the correlation time.



before 5 ps, gradually level off after 10 ps, and thus the viscosities are assessed from the mean values between 10 and 15 ps. As summarized in Table 2, the viscosity almost linearly increases from 9.56 to 15.94 cP as rising LaF₃ concentration, and therefore, the concentration dependence of shear viscosity can be presented as,

$$\eta(\text{cP}) = 9.08 + 1.27c_{\text{La}} \quad (6)$$

The fitted index R^2 is more than 0.98, meaning that the eqn (6) is practical to estimate the shear viscosities for different compositions of Flibe + $x\text{LaF}_3$. According to eqn (6), the viscosity of pure Flibe at 823 K is calculated as 9.08 cP, lower than that of pure Flibe (11.5 cP for 66 LiF: 34 BeF₂), but consistent with that of 73 : 27 Flibe (9.10 cP).³ The experimental viscosity of molten Flibe with the addition of lanthanide compounds is missing, but it can be confirmed that the introduction of high valence ions will increase the viscosity of the blank salt.⁵⁸ In addition, structural factors or specific ion interactions that influence the viscosity trend will be further elaborated in Section 3.3.

3.3. Local structural descriptors

3.3.1. Nearest neighbor distances. Fig. S3 (ESI[†]) plots the comparisons of RDFs among different LaF₃ concentrations at the same temperature, where all the $g(r)$ curves quickly converge to unity with a cutoff distance of 7 Å, indicating that these simulation cells are large enough to accommodate a system that shows intermediate range Coulomb interactions.⁵⁹ The RDFs and integral curves of Be–F, Li–F, F–F, and Li–Li show minor changes with LaF₃ concentration, while the short-range interaction of La–F and Be–Be are evidently different, from which it can be inferred that the density and potential energy (in Fig. 5) are mainly influenced by La–F and Be–Be interactions. Therefore, the RDFs and integral curves of Be–Be and La–F are further magnified in Fig. 8 in the range of 1–5 Å. According to Fig. S3 (ESI[†]), the R_1 of Be–F, Li–F, F–F, and Li–Li are extracted as 1.550, 1.858, 2.580, and 3.118 Å. The average ionic radius of F[–] is evaluated as 1.290 Å, larger than its crystallographic radius of 1.246 Å in lanthanide fluorides,⁶⁰

indicating that the repulsive force between F ions is enhanced after salt melting. From Fig. 8a, the peak height of Be–Be greatly increases with the concentration increments, while the peak position slightly right-shifts by about 0.02 Å as shown in Table 2, suggesting that the repulsive force between Be ions is slightly reduced. In Fig. 8b, the peak height prominently increases and peak position left-shifts from 2.362 to 2.292 Å along with the rising LaF₃ concentration subjected to eqn (7),

$$R_1^{\text{La-F}}(\text{Å}) = 2.335 - 0.008c_{\text{La}} \quad (7)$$

The fitted R^2 index is relatively low (0.96), and the c_{La} cannot be zero. The shortening of La–F distance means the attraction force between two ions is enhanced; in other words, the dynamics of La ions to drag the surrounding F ions are weakened, which is more remarkable in high LaF₃ concentration. Based on the La–F distances, the average ionic radius of La³⁺ is evaluated at 1.036 Å, similar to the crystallographic radius (1.050 Å) of La³⁺ when CN = 6, but significantly lower than the radius of 1.186 Å when CN = 9,⁶⁰ from which it can be deduced that the electron clouds of F and La partially overlap accompanied by the salt melts. Overall, the reduced repulsive force between Be ions and enhanced attraction force between La and F ions are profitable for the structural stability of the molten system.

3.3.2. Coordination numbers. Similarly, the CN₁ of Be–F, Li–F, F–F, and Li–Li at the first shell layers remain almost unchanged with LaF₃ concentration, and the average CNs are 4, 5, 12, and 8, respectively. It is demonstrated that the LiF₅^{4–} clusters are network-type connected referring to the CN₁ of Li–F and Li–Li, which is beneficial for the transport of Li ions. However, the CN₁ of Be–Be is only 1 as shown in Fig. 8 and the Be–Be distance (2.94 Å) is less than twice the Be–F distance (3.10 Å), indicating that the BeF₄^{2–} clusters in Flibe + LaF₃ are chain-type and not network-type connected. The tight interaction between Be and F, stable tetrahedral configuration, and restricted transport path result in a much lower diffusion coefficient of Be compared to the Li cation. The CN₁ of La–F (0.19 mol%) is 8.41 with a cutoff radius of 3.370 Å, larger than that of 6 in KF-rich molten fluorides.⁵⁴ The CN₁ of La–F is

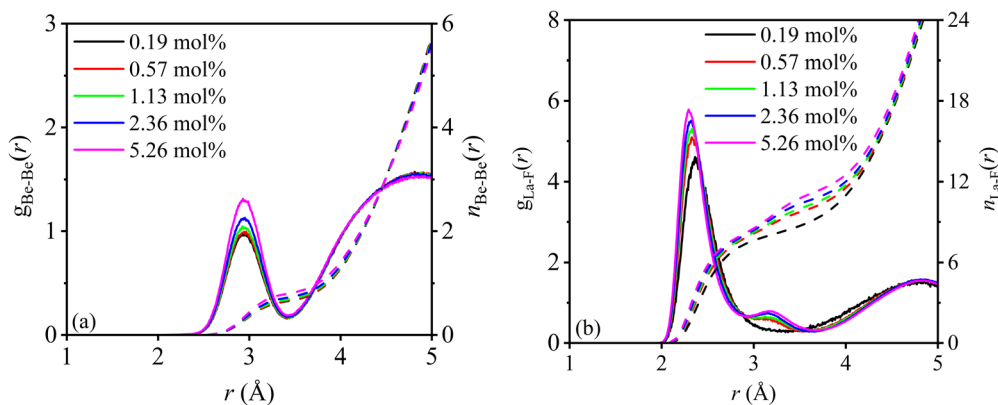


Fig. 8 RDFs and their integral curves of (a) Be–Be and (b) La–F for molten Flibe + LaF₃.



increased as the LaF₃ concentration increases and the relationship is subjected to eqn (8).

$$CN_1^{\text{La-F}}(\text{\AA}) = 8.367 + 0.255c_{\text{La}} - 0.026c_{\text{La}}^2 \quad (8)$$

The polynomial fitting index is up to 0.999, hinting that the CN₁ of La–F basically reached the maximum at 5 mol% LaF₃, and CN is maintained at 9 for more LaF₃ addition. Of interest is that the RDF curves of 0.57–5.26 mol% Flibe + LaF₃ are significantly different from that of low LaF₃ concentration system, where another shoulder peak appears at 3.15 Å and gradually enhances as the concentration increases. Therefore, the average CN of La–F will be increased if the cutoff radius covers the second small peak, meaning that 1–2 F ions may be stably distributed beyond the first coordination shell of LaF₈^{5–}, which is detrimental to the stability of the short-range ordered structure of molten Flibe + LaF₃. It should be pointed that the DP is trained without considering the repulsion between La–La, which will underestimate the La–La interaction and even shorten the La–La distance, thereby changing the RDF and CN of La–F. Furthermore, the increased coordination number and complex coordination surrounding for La ions lead to a decrease in diffusivity (eqn (4)) and an increase in the viscosity (eqn (6)) of the molten Flibe + LaF₃ system with the rising LaF₃ concentration.

4. Conclusions

The DP of molten Flibe + LaF₃ is developed for the first time based on machine learning methods and applied to investigate the LaF₃ concentration dependences of density, diffusion coefficient, viscosity, and structure. The database with 62 700 frames (ergodic 19 initial configurations) is collected from FPMD simulations, and excellent agreements of energies and forces as well as RDFs and MSDs demonstrate the accuracy and predictability of trained DP. The LaF₃ concentration dependences of density and potential energy are extremely linear, demonstrating the powerful transferability and flexibility of DP across different salt compositions. Furthermore, the ionic self-diffusion coefficients and shear viscosities of molten Flibe + LaF₃ are predicted by DPMD simulations, the relationships of weighted diffusion coefficient and shear viscosity on LaF₃ concentration are obtained, and the qualitative structure–property relationships are mined out. Finally, the DP provides more precise structural predictions over a wider concentration range, and the quantitative relationships between structural descriptors and concentration are accessed. Overall, these excellent results indicate the robustness of DPMD simulations, provide transferable DP for further studying other properties of Flibe with any LaF₃ concentration, and give insights for property predictions of other molten salts.

Author contributions

X. Li: conceptualization, investigation, methodology, writing-original draft, review & editing, and visualization funding

acquisition; T. Xu: methodology, investigation, and visualization; Y. Gong: supervision, project administration, funding acquisition.

Conflicts of interest

There are no conflicts to declare.

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