Temperature-dependent potential for the molecular dynamics of the superionic conductor $\beta$-PbF$_2$

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A B S T R A C T

Molecular dynamic (MD) calculations were performed to investigate the thermodynamic and structural properties of lead fluoride (PbF$_2$) by using a proposed inter-ionic temperature-dependent potential. This potential allows calculating with high precision the linear thermal expansivity and the lattice parameter as a temperature function. In addition, the potential can be represented as a sum of two contributions, a temperature-independent potential added to another temperature-dependent potential, considered last as a correction justified by the one-dimensional Newtonian quantum equation. Two fitting regions were considered, the first region from 300 to 700 K and the other one from 700 to 900 K. These regions arise naturally due to the smooth and continuous transition that PbF$_2$ undergoes until it reaches the superionic state and, allows us to model with high precision the anomaly in the dependence of the lattice parameter with the temperature of this material, a feature that until now under the molecular dynamic method has not been studied. These results are all in good agreement with the experimental measurements.

1. Introduction

Computational simulations applied to ionic systems are a powerful tool that allows us to study and predict their behavior when they are subjected to external stimuli, which in most cases cannot be achieved in the laboratory. Particularly, molecular dynamics (MD) is a classical method that bases its algorithm on the resolution of Newtonian motion equations [1], which allows the study of large ionic systems at a low computational cost. The quantum effects of each ion can be disregarded (i.e., electronic structure) because the choice of an effective interaction potential, acts manifesting the characteristics as a global average (see Eq. (1)). Therefore, these potentials are derived from quantum or empirical considerations. For this reason, a right choice in the mathematical form of the potentials is crucial to guarantee a behavior like the real one. In molecular dynamics (MD), these interaction potentials are adjusted to describe the actual behavior of physical systems employing constants, and it depends explicitly on the distance between the ions or atoms. However, if a temperature gradient is applied to the system, the effect of the temperature only manifests itself as a contribution to the kinetic energy of each atom [2]. Nevertheless, it is expected that the temperature slightly modifies the interactions between the ions. Due to this, it is necessary to include temperature-dependent corrections to the interaction potential. Currently, few studies employ a methodology that reveals a dependence on interaction potentials with temperature variation. Subramaniyan and Sun [3] developed an alternative to classical molecular dynamics called engineering molecular mechanics (EMM), which allows simulating physical properties in materials at high temperatures ($T > 0$ K). They employed the coefficient of thermal expansion (CTE), which is a well-studied property, converting the interaction potential into a temperature-dependent potential by allowing the adjustment constants to depend on the CTE. Although the authors manage to optimize the computation time, this method is based on the molecular statics (MS) and not on the MD. Instead, Nasehzadeh et al. [4] reevaluated the adjustment parameters of the Lennard-Jones potential and improve the calculation of the thermodynamic properties of liquids. They used the Chandler-Week-Andersen perturbation theory, which provides a precise analytical expression and allows determining that both $\sigma$ and $\epsilon$ can be expressed as a linear function of temperature. Other works that respond to more particular problems and that consider this type of dependency can be found in Refs. [5, 6, 7].
On the other hand, the study of ionic conductors through the use of MD is a current scientific topic, since there is still no precise understanding of the mechanisms involved in the ionic conductivity of the superionic state [8, 9, 10, 11]. One of the most relevant superionic conductors with fluoride structure is β-PbF2, which has a structure described as an FCC array of lead cations (Pb2+), where all the tetrahedrally coordinated interstitial sites are filled with fluoride anions (F−), being F− the mobile species. The lattice parameter, their density and specific heat capacity are 0.593 nm, 7.750 g/cm3 and 301 J Kg−1 K−1, respectively. The phase transition to the superionic state, undergone by this compound, is accompanied by a λ-type peak in the specific heat, no reorganization of the crystalline frame structure, and a smooth and continuous change in conductivity as the temperature increases [12]. Likewise, this material has an anomaly in its thermal expansion when it undergoes a reorganization of the crystalline frame structure, and a smooth and continuous change in conductivity as the temperature increases [12].

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In this work, the parameter $\rho_{F-Pb}$ was considered as a function of the temperature, with a linear relation:

$$\rho = \rho_0 (1 + \eta T) \quad (5)$$

In a similar way to Lennard-Jones coefficients [4] and the pre-exponential factor of the compact representation of Buckingham's potential [7]. By replacing the Eq. (5) in $U(r_{F-Pb})$ given by Eq. (4) obtains:

$$U(r_{F-Pb}) = A_0 \exp \left[ -\frac{r_{F-Pb}}{\rho_0 (1 + \eta T)} \right] \approx A_0 \exp \left[ -\frac{r_{F-Pb}}{\rho_0} \right] (1 - \eta T) \rho_0 \quad (6)$$

The result of the right side (Eq. (6)) is obtained by expanding the argument of the exponential in a series of power of temperature and using the fact that it is possible to rewrite this equation, therefore, the temperature-dependent part is:

$$A_0 \exp \left[ -\frac{r_{F-Pb}}{\rho_0} \right] \exp \left[ \frac{\alpha r_{F-Pb}}{\rho_0 T} \right] \approx A_0 \exp \left[ -\frac{r_{F-Pb}}{\rho_0} \right] \left( 1 + \frac{\alpha r_{F-Pb}}{\rho_0 T} \right) \quad (7)$$

However, Eq. (7) shows that $U(r_{F-Pb}) \approx U(r_F) \pm \delta U(r_{F-Pb}, T)$, which is the condition imposed in Eq. (3), that the potential must satisfy.

### 3.1. Determination of the temperature-dependent parameter

To determine the temperature dependence of the $\rho_{F-Pb}$ parameter, this is slightly modified from 0.490 to 0.520 eV for 300, 400, 500 and 600 K, and modified from 0.490 to 0.510 eV for 700–900 K [16], performing a new simulation set with the previous conditions. For each temperature, the $\rho_{F-Pb}$ is obtained to adjust the experimental lattice potential.
It is found that as the temperature increases, the parameter $\rho_{\text{Pb}-\text{F}}$ decreases and therefore the potential suffers a decrease, causing the fluoride and lead ions to experience a lower repulsion force with each other. While it might be expected that the repulsion between the ions would be greater because the crystal expands with temperature, this behavior arises as compensation for the classical thermal energy that is imparted to each ion. Likewise, a decrease in potential energy allows fluorine ions to move more easily between available sites in the lattice.

Figure 2 shows the behavior of the potential of the Pb–F interaction, in the region of low (Figure 2a) and high (Figure 2b) temperatures. It is inferred that the proposed modification affects the potential in the order of tenths of eV by varying the thermal bath temperature by ± 100 K, corroborating that the Eq. (8) agrees with the Eq. (3).

3.2. Validation of interatomic potential

3.2.1. Structural stability

To verify that the proposed potential does not lead to the study material undergoing a structural phase transition, or an undesired disorder in any of its sublattice with the rise of the temperature, the radial distribution functions $g(r)$ is calculated. This function allows knowing how the ions are distributed from the atoms of the material with respect to a type of ion in a spherical shell (refer to Figure 3). The $g(r)$ has a first well-defined maximum attributed to the first neighbors, and as the temperature increases, it loses its shape, decreasing their intensity and increasing their width, because the ions experience greater mobility as the temperature increases of the system. However, unlike lead ions (Figure 3c), the $g(r)$ of fluorine ions (Figure 3a) changes to a radial distribution function, similar in shape to liquid materials. This behavior partly explains the superionic conductivity in this kind of material, where the sublattice of the mobile ions, has a quasi-liquid behavior due to its high mobility and diffusion, which can be corroborated in Figure 4, in which the ions of lead remain in their network sites, while those of fluorine begin to diffuse gradually through the crystal as the temperature increases. From Figure 3b, a mean distance of 2.42 Å between the lead and fluorine ions at room temperature was calculated. This distance turned out to be slightly greater than that reported by Silva et al. [17], using temperature-independent potentials, but lower than that experimentally

$$ \rho_{\text{Pb}-\text{F}}(T) = \begin{cases} 0.5089\text{Å} - 3.34 \times 10^{-6}T\text{Å/}K, & \text{if } 300K \leq T \leq 700K \\ 0.4654\text{Å} + 1.1360 \times 10^{-4}T\text{Å/}K - 8.1603 \times 10^{-8}T^2\text{Å}^2/K^2, & \text{if } 700K < T \leq 920K \end{cases} $$

(8)
Figure 3. Function of radial distribution of $\beta$-PbF$_2$ at different temperatures, for (a) F–F, (b) Pb–F, (c) Pb–Pb.

Figure 4. Coefficient of diffusion against temperature for ionic fluorine and lead spices.

Figure 5. Dependence of the lattice parameter with temperature.
3.2.2. Thermal expansion coefficient

Figure 5, shows the variation of the lattice parameter when the temperature of the system, is increased and compared with the results obtained experimentally by Goff et al. [18], (in red circles), against those calculated in this paper (in cyan diamonds); when using the modification proposed in Eq. (8). A high coincidence with that reported experimentally is shown, even fitting to the anomaly present near 700 K, so the choice of two regions shows to be an appropriate process.

Once the dependency of the lattice parameter \( a_0 / \alpha (\text{K}) \), with the temperature, is found, it is possible to find the linear thermal expansion coefficient \( \alpha \) (see Eq. (9)) as [19]:

\[
\alpha = \frac{1}{a_0} \frac{\partial a_0}{\partial (293 K)} \tag{9}
\]

A sixth-grade polynomial fit for the data obtained in this work was done, and its gradient was calculated (see Figure 6) where it is compared with experimental data of the linear thermal expansion coefficient obtained by Roberts and White [20]. It is interesting to note that the calculated curve has a maximum close to 670 K while the experimental curve around 690 K. However, both have the same line shape and order of magnitude, with an error slightly lower than 30\% in area, in the range of 350–800 K. Beyond 800 K, the data deviates from experimental behavior. Also, it is the first time that this type of graph has been made by MD calculations for \( \beta \)-PbF\(_2\).

3.2.3. Heat capacity

The heat capacity \( C_p \) of any material is defined as the amount of heat per unit of mass required so that its temperature raises 1 K [21]. For the molecular dynamic case, \( C_p \) is calculate by:

\[
C_p = \frac{1}{n} \left[ \frac{\partial H}{\partial T} \right]_{\text{NpT}} \tag{10}
\]

Being \( n \) the number of moles, \( H \) the enthalpy computed in the NpT ensemble of the system, and \( T \) is the crystal temperature. The \( C_p \) curve calculated from Eq. (10), from 300 K to 800 K is plotted in Figure 7. The \( C_p \) increases in a wide region around 700 K, near the superionic transition, predicting a phase transformation to the superionic state due to an increase in internal energy, related to the mobility of mobile ions. In previous study [11], the \( C_p \) values employing only the Walker’s potential showing a higher value \((80.49 \text{ J K}^{-1} \text{ mol}^{-1})\) than the experimental one \((76.81 \pm 0.51 \text{ J K}^{-1} \text{ mol}^{-1} \text{ at } 400 \text{ K})\) and above 700 K a chaotic behavior arise, without exhibiting a phase transition. In this work, a lower value \( C_p \) than the experimental is calculated \((52.3 \text{ J mol}^{-1} \text{ K}^{-1} \text{ at } 500 \text{ K} \text{ against } 79.87 \text{ J mol}^{-1} \text{ K}^{-1} \text{ measured by Volodich et al., } [22])\), and a phase transition is predicted. The difference between the values (experimental and theoretical) is the non-existence of a nominal value in the literature for temperatures above room temperature. However, a similarity in the shape of the lines can be appreciated. From temperatures above 800 K, a difficulty arises in fitting the enthalpy data.

4. Conclusions

In summary, the potential proposed by Walker et al. [15], was modified, to describe the structural properties of \( \beta \)-PbF\(_2\) as a temperature function, by making this equation parametrically dependent on temperature. The fitting constant \( \rho_{\beta-PbF_2} \) was replaced by a linear function \( \rho_{\beta-PbF_2}(T) \). This parameter was fitted with experimental data of the lattice parameter versus temperature. It was found that two adjustment regions naturally emerged, the first one from 300 to 700 K and another from 700 to 900 K. The first is characterized by having a linear behavior, while the second exhibits a second-order polynomial form. From these functions, calculations by molecular dynamics were made, and it is possible to conclude that there exists a high coincidence between the data of the evolution of the lattice parameter with the temperature and those calculated with our proposed potential, it even fits the anomaly present in the transition temperature towards the superionic state. From the \( g(r) \) function obtained, it is possible to conclude that the proposed potential does not involve a structural phase transition, and the ions distance is almost equal to the experimental one. The \( C_p \) curve is firstly for this material calculated from 300 to 800 K, exhibiting a growing line shape, until a maximum; then, around it, a phase transition towards the superionic state is predicted.

Declarations

Author contribution statement

López J.D.; Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Contributed materials, analysis tools or data; Wrote the paper.

Diosa J.E., H. Correa: Conceived and designed the experiments; Analyzed and interpreted the data.
García G.: Conceived and designed the experiments; Contributed materials, analysis tools or data.
Mosquera E.: Analyzed and interpreted the data; Wrote the paper.

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Data availability statement
Data associated with this study has been deposited at Data in Brief - Volume 28, February 2020, 104865 (https://doi.org/10.1016/j.dib.2019.104865).

Declaration of interests statement
The authors declare no conflict of interest.

Additional information
No additional information is available for this paper.

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