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Entropy and Isokinetic Temperature in Fast Ion Transport

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Ion transport in crystalline solids is an essential process for many electrochemical energy converters such as solid-state batteries and fuel cells. Empirical data have shown that ion transport in crystal lattices obeys the Meyer-Neldel Rule (MNR). For similar, closely related materials, when the material properties are changed by doping or by strain, the measured ionic conductivities showing different activation energies intersect on the Arrhenius plot, at an isokinetic temperature. Therefore, the isokinetic temperature is a critical parameter for improving the ionic conductivity. However, a comprehensive understanding of the fundamental mechanism of MNR in ion transport is lacking. Here the physical significance and applicability of MNR is discussed, that is, of activation entropy-enthalpy compensation, in crystalline fast ionic conductors, and the methods for determining the isokinetic temperature. Lattice vibrations provide the excitation energy for the ions to overcome the activation barrier. The multi-excitation entropy model suggests that isokinetic temperature can be tuned by modulating the excitation phonon frequency. The relationship between isokinetic temperature and isokinetic prefactor can provide information concerning conductivity mechanisms. The need to effectively determine the isokinetic temperature for accelerating the design of new fast ionic conductors with high conductivity is highlighted.

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DOI: 10.1002/advs.202305065

1. Introduction

The search for materials with high ionic mobility and diffusivity, so-called fast ionic conductors (FICs), remains an ongoing quest. FICs are essential components for electrochemical energy conversion devices, such as solid oxide fuel cells, electrolyzers, and solid-state batteries.[1-3] The commercialization of these devices could constitute important aspects of the development of green energy. All of these applications require high ionic conductivity. To achieve these objectives, it is crucial to fully understand the peculiarities in ion transport mechanisms. The temperature dependence of ionic conductivity in solids often follows the Arrhenius law, which is characterized by constant activation energy and prefactor,[4] written as:

$$\sigma = \frac{\sigma_0}{T} \exp\left(-\frac{E_a}{k_B T}\right) \tag{1}$$

where σ is the conductivity, T is the temperature, σ_0 is the prefactor, E_a is the activation energy, and k_B is the Boltzmann constant.

It is frequently assumed that decreasing the activation energy (lowering the activation barrier) of ionic conductors will improve their conductivity. However, the improvement in ionic conductivity due to the reduced activation energy may be less than expected, because the prefactor is also lower, and compensates the decrease in activation energy. In some cases, the ionic conductivity is even lower when the activation energy is reduced.^[5] Since the 1920s, numerous examples of such compensation have been found. This compensation effect in the prefactor is found to be proportional to E_a , or to ΔH , the activation enthalpy. [6-11] This results in the intersection of the Arrhenius plots for different activation energies of related samples at an isokinetic temperature. This observation is sufficiently common to be called^[10] the compensation law (compensation effect^[12]), the isokinetic law, and the MNR, for the authors who reported its observation in disordered solids.[8] In particular, such an effect is frequently observed in electronic and ion conduction and in atomic and ionic diffusion. $^{[13-19]}$ Figure 1a shows an example Arrhenius plot for conductivity of garnet-structured lithium ionic conductors $\text{Li}_6\text{MLa}_2\text{Ta}_2\text{O}_{12}$ (M = Ba, Ca, Sr, and $\text{Sr}_{0.5}\text{Ba}_{0.5}$) with variations of dopant or dopant concentration. In terms of Equation (1), with logarithm applied, we may write

$$ln\sigma_0 = ln\sigma_{00} + \frac{E_a}{k_B T_{iso}} \tag{2}$$

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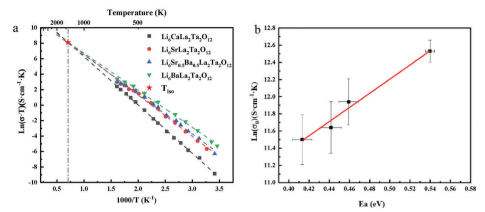


Figure 1. Arrhenius plot a) and Meyer-Neldel plot b) of garnet Li₆MLa₂Ta₂O₁₂ (M = Ba, Ca, Sr, and Sr_{0.5}Ba_{0.5}) lithium ionic conductors.^[20] The T_{iso} is shown as asterisk and the grey dash dot line indicates the value of T_{iso} . The solid line in (b) is the least square fit line according to Equation (2).

where σ_{00} is the isokinetic prefactor, which is determined by parameters related to elementary ion hopping in FICs, and is discussed in detail in Section 6. T_{iso} is the isokinetic temperature, and $k_B T_{iso}$ is also known as Meyer-Neldel energy, sometimes denoted as Δ_0 .

The activation energy and prefactor are determined from the slope and intercept of the Arrhenius plot, as shown in Figure 1a. The point of isokinetic temperature is also shown in Figure 1a. By fitting the prefactor as a function of activation energy using Equation (2), that is, the Meyer-Neldel plot (M-N plot), as shown for the garnet family $\text{Li}_6\text{MLa}_2\text{Ta}_2\text{O}_{12}$ in Figure 1b, T_{iso} is obtained from the inverse slope of the linear least square fit to the data, as discussed below.

The MNR indicates that when the activation energy changes, the prefactor changes accordingly, compensating for the change in activation energy, reducing the improvement in conductivity. Therefore, researchers have been trying to break the limitation of the MNR^[21] or look for materials which do not show a compensation effect.^[22] However, since the ionic conductivity is affected by multiple factors related to the material structure, a comprehensive understanding of the MNR in ion transport has been challenging.

Here, we review the contribution of entropy and lattice dynamics to the ionic conductivity within the framework of the MNR. After briefly introduce the physical meaning of the MNR and the condition under which this rule is applied, we then discuss the material parameters that determine the isokinetic prefactor based on the fundamental physics of ion transport, and the effort to apply the MNR to improving the ionic conductivity by controlling the isokinetic temperature. Then we evaluate the approaches to determination of the isokinetic temperature, and the relationship between lattice vibration and isokinetic temperature. Finally, we investigate the origin of the relationship between isokinetic temperature and isokinetic prefactor and examine the mechanism of conductivity in FICs.

2. Activation Entropy and Isokinetic Temperature in MNR

It has been known since the work of Eyring $^{[23]}$ in the 1920s, that a more rigorous form of the Arrhenius equation for any activated

process is given by

$$X = X_0 \exp\left(\frac{-\Delta G}{k_{\rm R}T}\right) \tag{3}$$

where the free energy of activation, ΔG , is given by

$$\Delta G = \Delta H - T \cdot \Delta S \tag{4}$$

In Equation (4), ΔH and ΔS are the activation enthalpy and entropy. It has been observed that the activation enthalpy and entropy of chemical reactions often show a linear relation, the slope of which has been defined as the isokinetic temperature. But caution must be practiced to not over-interpret the outcome of such mathematical construction. [24–26] The activation entropy, ΔS , for movement of a carrier, electronic or ionic, called the migration entropy, is expressed as [27,28]

$$\Delta S = k_B ln \left(\frac{\prod_{i=1}^{3N} \nu_i^I}{\prod_{i=1}^{3N-1} \nu_i^S} \right)$$
 (5)

where v_i^I and v_i^S are the vibrational frequencies of the initial state and the transition state of the ion hopping process, respectively. Equation (5) shows that ΔS depends only upon the ratio of vibrational distribution functions of the initial and transition states.

It is now generally recognized^[10] that MNR occurs in closely related systems. It has recently been shown^[29] that it is obeyed when the free energy of the samples considered is a linear function of T and of one other physical variable, such as in a structure-property relationship. We take this to be the condition for "closely related systems". Further, $\frac{1}{T_{\rm iso}} > 0$ when ΔH is large compared to $k_B T$ and to the excitation energies of the system under study, and that it is an entropic effect. That is, if the process obeys MNR, there is a contribution to ΔS that compensates ΔH , and is given by:

$$\Delta S_M = \frac{\Delta H}{T_{iso}} \tag{6}$$

 ΔS_M has been called multi-excitation entropy. The multi-excitation entropy model^[10] (MEE model) suggests that, when

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the thermal energy is too small to provide the excitation energy for ion transport, ΔS_M is associated with the collected excitations which provide the energy needed in order to overcome the barrier.

For a given ΔH , at low T, X is larger than it would be if such compensation did not exist. At T_{iso} , X is independent of ΔH . If the data of Figure 1a, are extrapolated to $\frac{1}{T}=0$, we may obtain the log or ln of the experimental prefactors. If the latter is plotted as a function of E_a , the inverse slope of the best linear fit yields T_{iso} as in Figure 1b, the M-N plot. As discussed in Section 5, this is generally a more accurate determination of T_{iso} , than can be made from Figure 1a. Then, ΔS_M can be calculated using Equation (6). However, there may be a contribution to ΔS that is independent of ΔH , which we call the change of configurational entropy, ΔS_C , between the transition and initial states. Then

$$\Delta S = \Delta S_M + \Delta S_C \tag{7}$$

It is not generally feasible to accurately calculate ΔS_C from computational or experimental data. However, a comparison between Li-ion conductors LiTi₂(PS₄)₃ (LTPS) and Li₁₀GeP₂S₁₂ (LGPS) suggests that a smooth energy landscape can lead to a larger entropy of the transition state. Fortunately, we can frequently determine the sign of ΔS_C . In the great majority of experimental circumstances, ΔS_C is small, or positive. In a typical kinetic process, the disorder, that is, the configurational entropy, of the transition state is likely to be very little different from that of the initial state. The atoms or molecules in a chemical reaction, the mobile species in a diffusion or conduction process can move in many directions. Thus, at some T below T_{iso} , either ΔG becomes zero, so that Arrhenius no longer applies (the process is no longer activated), or the condition $k_B T \ll \Delta H$ is no longer valid, so that MNR no longer applies.

In a small fraction of experiments, most notably on fast ionic conduction, as discussed below, and on relaxation of polymeric glasses, [30] ΔS_C is negative. Then, ΔG is negative at T_{iso} , that is, the phenomenon continues to be activated, with higher ΔH yielding a more rapid process. If the controlling mechanism does not change, and $k_BT \ll \Delta H$ continues to be valid, this situation continues until a temperature, approximately:

$$T = \frac{\Delta H}{\Delta S} \tag{8}$$

is reached, so that ΔG becomes zero. For entropy to be lower in the transition than in the initial state, this motion must be constrained. This is precisely the case for the two notable examples of negative ΔS_C which we have cited. In glass-forming polymers, a molecule is normally intertwined with its neighbors. For two neighboring molecules to move with respect to each other, most likely they are in a particular relative position. In each of the relaxations of a polymer glass, there are distinct values of T_{iso} and of negative ΔS_C associated with the molecular segments which move. [30] For the relaxation at the highest temperature, α relaxation, for example, large portions of neighboring molecules must be close to parallel.

It is now well established that FICs, such as perovskite-type oxides, behave as they do because of particular paths in their crystal structure, [31] through which the ions can move readily. The price

of this logistic advantage is their confinement, and the associated low entropy. Their conductivities above T_{iso} may behave in a way which is quite different from that of typical semiconducting or insulating materials, whose carrier densities are activated, and carrier mobilities decrease slowly with T. In FICs, both maybe activated.

3. Applicability of MNR in Solid State Ionics

We now examine the fundamental physics of ion transport, in order to understand the effect of MNR, and especially, of $T_{iso,}$ on ionic conductivity at T. The conductivity, σ , is given by:

$$\sigma = cq\mu \tag{9}$$

where c is the mobile ion concentration, q is the ionic charge, and μ is the ionic mobility.

The ionic mobility μ is related to the macroscopic long-range diffusion coefficient D_{σ} by the Nernst-Einstein equation:^[32]

$$\mu = \frac{qD_{\sigma}}{k_{\rm B}T} \tag{10}$$

The classical model of ion diffusion in solids considers that the ion hopping events are random, that is, the ion hopping direction is independent of the previous hopping direction. This model applies when the ion concentration is low. [33] When ion transport is uncorrelated and independent, one can treat D_{σ} as the random diffusion coefficient D_{σ} :

$$D_r = \frac{a^2 v}{b} \tag{11}$$

where a is the ion jump distance, v is the ion jump frequency for successful jumps which leads to macroscopic diffusion, b is a geometric factor, for 1D, 2D, or 3D diffusion, b is 2, 4, or 6 respectively.

The ion jump frequency v is described by:

$$v = v_0 \exp\left(-\frac{\Delta G}{k_B T}\right) = v_0 \exp\left(\frac{\Delta S}{k_B}\right) \exp\left(-\frac{\Delta H}{k_B T}\right)$$
(12)

where v_0 is the attempt frequency which includes both the successful jumps and unsuccessful jumps.

According to Equation (9) to Equation (12), the complexity of temperature-dependent ionic conductivity σ can be mathematically illustrated by:[34]

$$\sigma(T) = \frac{1}{b} \frac{cq^2}{k_B T} a^2 v_0 \exp\left(\frac{\Delta S}{k_B}\right) \exp\left(-\frac{\Delta H}{k_B T}\right)$$
(13)

where the enthalpy ΔH is also called activation energy (denoted as E_a). Comparing Equation (1), and Equation (13), the prefactor is written as:^[5]

$$\sigma_0 = \frac{1}{b} \frac{cq^2}{k_B} a^2 v_0 \exp\left(\frac{\Delta S}{k_B}\right) \tag{14}$$

From Equation (14), one may see that, in addition to the entropy, the prefactor is affected by many parameters, including

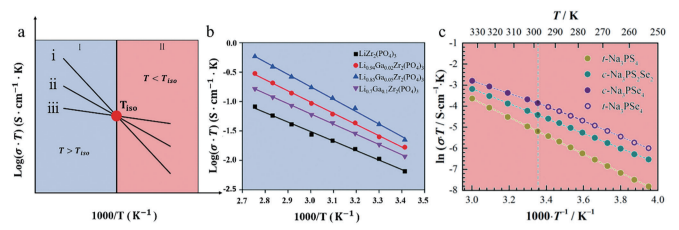


Figure 2. a) The role of T_{iso} in ionic conductivity. Region I (blue) and Region II (pink) represent the cases of measurement temperature larger and smaller than T_{iso} . The lines i, ii, and iii represent three conductivity lines with activation energies and prefactors decreasing sequentially. b) the lithium-ion conductivity of Li_{1.3x}Ga_xZr₂(PO₄)₃ (x = 0, 0.02, 0.05, 0.1). Reproduced with permission. Copyright 2021, The Royal Society of Chemistry. c) The sodium ionic conductivity of Na₃PS_{4-x}Se_x (x = 0, 2, 4). Reproduced with permission. Dopyright 2018, American Chemical Society.

the mobile ion concentration, ion jump distance, and attempt frequency. Therefore, it is worth noting that MNR applies only to "closely related systems", when the material candidates have similar composition and structure, so that other parameters do not differ much between different samples or for variable measurement parameters, as discussed in the literature on chemical reactions [24,25,35] and in the previous and following sections. However, when the ion concentration changes, the defect formation energy may also vary, causing changes in the activation energy. The variation in ion concentration adds difficulties to the determination of T_{iso} , and may be one reason for the scattering of data. Therefore, we propose that T_{iso} can be more rigorously determined using the diffusion coefficients instead of ionic conductivities, when the influence of concentration is excluded.

4. Improving the Ionic Conductivity According to the MNR

Reducing the activation energy has been considered to be an effective method for improving ionic conductivity. However, the MNR shows that lower activation energy is not always related to high ionic conductivity. Here, we consider how activation energy determines the ionic conductivity according to MNR.

Because of the entropy-enthalpy compensation suggested by the MNR, we can use E_a to replace ΔS_M to formally describe the ionic conductivity. That is, combining Equation (2), Equation (6), Equation (7), and Equation (13), the conductivity becomes:

$$\sigma(T) = \frac{1}{b} \frac{cq^2}{k_B T} a^2 v_0 \exp\left(\frac{\Delta S_C}{k_B}\right) \exp\left[\left(\frac{E_a}{k_B}\right) \left(\frac{T - T_{iso}}{T_{iso}T}\right)\right]$$
(15)

where T is the measurement temperature, or the operating temperature. Equation (15) suggests that the argument $(\frac{E_a}{k_B})(\frac{T-T_{iso}}{T_{iso}})$ in the exponent may be considered to be an indicator for modulating the material conductivity. If $T > T_{iso}$, the measured conductivity falls in Region I in **Figure 2**a. That is, the ionic conductivity is

higher when E_a is larger. When $T < T_{iso}$ and the measured conductivity is located in Region II in Figure 2b, it is necessary to decrease E_a to improve the ionic conductivity. When $T = T_{iso}$, the conductivity is independent of E_a .

In Figure 2 we show examples of the behavior described above. In Figure 2b, the conductivity of $\mathrm{Na_3PS_{4-x}Se_x}$ decreases with the increase of E_a , whereas that of $\mathrm{Li_{1-3x}Ga_xZr_2(PO_4)_3}$ (LGZP) increases with the increase of E_a in Figure 2c. The T_{iso} of LGZP^[36] and $\mathrm{Na_3PS_{4-x}Se_x}^{[37]}$ are presented in **Table 1**. According to the above discussion, the conductivity of LGZP and $\mathrm{Na_3PS_{4-x}Se_x}$ should fall in Region I and Region II of Figure 2a, respectively. This is confirmed by the measured conductivity, as shown in Figure 2b,c. The proton conductivity of $\mathrm{BaZr_{0.9}Y_{0.1}O_3}$ under high pressure $\mathrm{[^{38]}}$ shows the same behavior as in the $\mathrm{Na\text{-}ion}$ conductivity of $\mathrm{Na_3PS_{4-x}Se_x}$. More data from various lithium-ion and proton conductors are presented in Tables S1 and S2 (Supporting Information). For most of the materials investigated, T_{iso} is higher than the measurement temperature.

The activation energy can be affected by multiple factors including crystal symmetry, defects, and lattice softness.^[34] For related material systems, the activation energy is lower when the crystal structure exhibits higher symmetry and less disorder. A softer lattice is generally considered to be related to lower activation energy.^[21] In practice, the activation energy can be tuned by doping,^[39] applying strain,^[40] by modifying the vibration frequency,^[21] and the density of grain boundaries.^[41]

Notably, interesting results have been found in recent investigations whose objective was to reduce the activation energy. For

Table 1. The measurement temperature T, isokinetic temperature T_{iso} , and T vs. T_{iso} of $\text{Li}_{1.3x}\text{Ga}_x\text{Zr}_2(\text{PO}_4)_3^{[37]}$ and $\text{Na}_3\text{PS}_{4-x}\text{Se}_x$ (x=0,2,4). [36]

Material	Operating temperature T [K]	T_{iso} [K]	T vs. T _{iso}
$Li_{1-3x}Ga_xZr_2(PO_4)_3$	293–363	224.9	$T > T_{iso}$
$Na_3PS_{4-x}Se_x$ (x=0, 2, 4)	250–330	396.7	$T < T_{iso}$



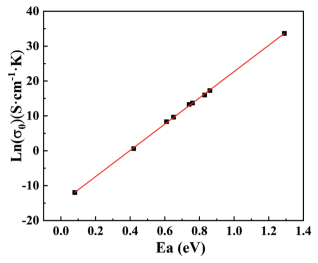


Figure 3. Meyer-Neldel plot of $\rm Li_{6.4}La_3Zr_{1.4}Ta_{0.6}O_{12}$ under variable pressure. [42]

instance, in the materials that manifest mobile ion disordering, the frustration in the LTPS framework enlarges the ionic jump distances compared to LGPS.^[22] As a result, LTPS exhibits a larger prefactor but a lower activation energy, which deviates from the MNR. Thus, the disorder in the material offers an alternative approach to significantly increasing the ionic conductivity.

5. Determination and Physical Significance of Isokinetic Temperature in Fast Ion Conductors: The Role of Lattice Vibrations

In Section 3, we have suggested that isokinetic temperature is affected by the material structure, lattice parameters, and composition. All these factors vary when the dopant concentration changes, thus makes the determination of isokinetic temperature challenging. Most often, the investigation of MNR of a property of a family of similar materials involves the preparation of a material by different techniques, or of similar composition, for example, by element substitution and doping. [17,39] Conduction in FICs may also be investigated using a single sample by imposing strain in the material (e.g., with pressure [18]). For instance, **Figure 3** shows a M-N plot demonstrating the strain-induced variation in E_a and σ_0 under high compressive strain. [42] Then T_{iso} is calculated from Equation (2) according to experimental data.

It is evident from Figure 3 that under these circumstances, of a single material under varying pressure, MNR is rigorously obeyed, as predicted by Sapunov. [35] In contrast, in Figure 1b, in which garnets containing different ions are compared, the criterion of "closely related systems" is not completely satisfied. When tuning the material composition, the adjustment in lattice parameters leads to changes in migration entropy. We cannot use MNR to determine the properties of these materials. However, it can still be used as a rule of thumb to suggest that such systems may be expected to exhibit similar values of $T_{\rm iso}$. [17,42,43]

A recent molecular dynamics (MD) study of MNR in atomic diffusion in simple metals^[44] strongly suggests that the phe-

nomenon is quite complicated, with substantial changes in vibrational densities of states between the initial and the transition state during diffusion, and that this provides the entropy for the compensation. Density functional theory and other first principles computational methods can only obtain the entropy, S, and activation energy E_a , of the initial state, but not ΔS_M . [34,42] Therefore, isokinetic temperature cannot be calculated from such theories, for now. To date, there have been no MD studies of MNR in ion conduction. Thus, we rely upon simple models or experiments to clarify the situation.

Within the simple, phenomenological, MEE model, [10,45] inspired by microscopic models for particular processes, a prediction for $T_{\rm iso}$ has been proposed. It assumes that one excitation of the system is most strongly coupled to that process which takes place, when ΔH is large, a number of these excitations must be accumulated to overcome the barrier. This concentration results in ΔS_M . [10] For optical phonons it is proposed that:

$$k_B T_{iso} = \frac{h \nu}{l n \kappa} \tag{16}$$

where h is Planck's constant, $h\nu$ is the excitation energy, and κ is a coupling constant. It has been suggested that $ln\kappa$ for polaronic materials, particularly ionic polarons, is related to the characteristic phonon occupation, [46] that is, the phonon number in a specific excitation state of vibrational mode.

The MEE model prediction^[10] for T_{iso} of a material, in which the only excitations are acoustic phonons, is different from Equation (16). In that case, it does not predict a particular frequency within the broad spectrum, for a situation such as that considered in ref.[44]. In experiments concerning a number of phenomena, it has been possible to identify excitations which satisfy Equation (16), assuming^[10] that $ln\kappa$ ranges between 0.5 and 2. These include studies of chemical reactions,^[47] where the excitations are those of molecular vibrations; relaxations,^[48] where they are the energies of the relaxing entities, conduction of FICs,^[17,18,43] where they are optical phonons.

It is still not clear whether Equation (16) is widely applicable. For example, in Y-doped BaMO3 (M = Zr or Ce) proton conductors, the relation between isokinetic temperature and average M-O stretch vibration follows Equation (16), as shown in Figure 4a.[17] In contrast, in lithium superionic conductors (LISICON) and olivine compounds, the phonon band center decreases with the increase of Meyer-Neldel energy, as illustrated in Figure 4b.[49] It is possible that the excitations responsible for T_{iso} are not Li phonons in these lithium conductors. However, recent work on garnet-type lithium conductors using isotope substitution have demonstrated that lower lithium vibration frequency corresponds to higher ionic conductivity.^[50] Clearly, further investigation as to whether Equation (16) is universally applicable to FICs is needed. If it is, it is important to identify the phonons which contribute the excitation that determines the isokinetic temperature. One promising experimental method to perform this task could be nuclear resonant vibration spectroscopy (NRVS).[51] New developments in synchrotron based xray methods permit the determination of element-specific vibration spectra, provided the element is available as a Mössbauer active isotope.[51]

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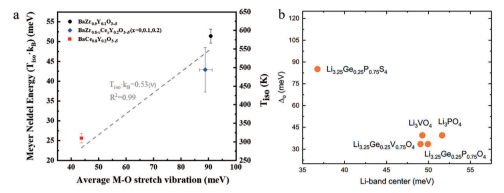


Figure 4. a) The relationship between Meyer–Neldel energy or isokinetic temperature and the average M-O stretch vibration $\langle \nu \rangle$ for BaZr_{0.9}Y_{0.1}O_{3-δ}, BaZr_{0.8}-xCe_xY_{0.2}O_{3-δ} (x=0,0.1,0.2), and BaCe_{0.8}Y_{0.2}O_{3-δ}. Reproduced with permission. [17] Copyright 2021, Wiley-VCH. b) The relationship between Meyer–Neldel energy (Δ_0) and Li-band center for Li_{3.25}Ge_{0.25}P_{0.75}S₄, Li_{3.25}Ge_{0.25}V_{0.75}O₄, Li₃VO₄, Li₃PO₄, and Li_{0.325}Ge_{0.25}P_{0.75}O₄. Reproduced with permission. [49] Copyright 2018, American Chemical Society.

6. Relationship between Isokinetic Temperature and Isokinetic Prefactor

Over the past several decades, it has been observed^[10] that for some families of electronic and ionic conductors obeying MNR, there is a correlation, that is, an approximate relationship between the isokinetic prefactor, σ_{00} and the isokinetic temperature T_{iot} , σ_{00} , σ_{00} , and σ_{00} are the isokinetic temperature σ_{00} , σ_{00} , and σ_{00} are the isokinetic temperature σ_{00} , σ_{00} , and σ_{00} are the isokinetic temperature σ_{00} , σ_{00} , σ_{00} , and σ_{00} are the isokinetic temperature σ_{00} , σ_{00} , σ_{00} , σ_{00} , and σ_{00} , σ_{00} ,

$$\ln \sigma_{00} = \ln \sigma'_{00} - \frac{E_{\rm n}}{k_{\rm R} T_{\rm iso}} \tag{17}$$

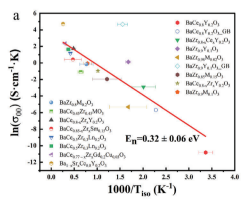
where σ'_{00} and E_n are empirical constants, discussed below. From Equation (16) we see that T_{iso} is inversely proportional to κ , the coupling to excitations. On this basis, it has been proposed^[10,43] that positive E_n corresponds to polaronic conduction, while negative E_n corresponds to trap-limited conduction. This has led to the identification of polaronic conduction in numerous materials. These include electronic conduction in chalcogenide glasses, ^[54] proton conductivity in minerals, ^[19] and ionic conductivity in perovskite-type oxides. ^[17] **Figure 5** shows other examples from the literature. ^[55–65]

Figure 5a shows the isokinetic prefactor σ_{00} as a function of the reciprocal of isokinetic temperature T_{iso} for some perovskite-type proton conductors. Neutron scattering experiments have also confirmed that for perovskite proton conductors, the proton jump time follows a polaron model. [66,67] Figure 5b shows a similar relationship for the LGPS family Li-ion conductors with variable compositions. Note that the data points scatter because the criterion of closely related system is not completely satisfied, for instance, the mobile ion concentration, attempt frequency, or other parameters change. Now, let us consider the significance of Equation (17) and the physical meaning of σ'_{00} and E_n .

Applying Equation (2), Equation (6), and Equation (7) to Equation (14), yields

$$\sigma_{00} = \frac{1}{b} \frac{cq^2}{k_B} a^2 v_0 \exp\left(\frac{\Delta S_C}{k_B}\right)$$
 (18)

Therefore, in random and uncorrelated ion hopping processes, the isokinetic parameter, σ_{00} , is determined by the ion concentration, jump distance, attempt frequency, and ΔS_C . As discussed in Section 2, ΔS_C is positive or small for most properties of most



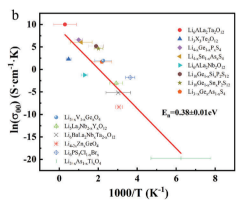
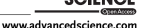


Figure 5. Correlation between isokinetic prefactor σ_{00} and isokinetic temperature T_{iso} for various compositions of perovskite-type proton conductors a), and lithium-ion conductors b). [55-65] The solid lines are the fit according to Equation (17). In (a), the solid, hollow, and semi-solid symbols correspond to the grain, grain boundary, and total conductivity. (All the relevant data obtained from literature which is found in Tables S1 and S2, Supporting Information).

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materials. However, as shown in Figure 2, it is negative and non-negligible for conductivity of FICs. As we may see, σ_{00} depends exponentially upon ΔS_C . There is no reason to expect the linear terms in Equation (8) to exhibit large changes from one member of a family to another. This leads us to suggest that, for FICs:

$$\sigma'_{00} = \frac{1}{h} \frac{cq^2}{k_B} a^2 v_0 \tag{19}$$

and E_n is expressed by:

$$E_n = -T_{iso}\Delta S_C \tag{20}$$

Therefore, E_n is related to the change of configurational entropy between initial and translational states, ΔS_C . It has been suggested that the value of E_a compared to E_n can determine the direction to modulate the isokinetic temperature in order to tune the ionic conductivity. [17] and thus E_n is called the critical energy of materials. [17]

Nevertheless, the extent to which the model we have proposed here may apply to other materials, including isotropic ionic conductors, is not evident. As pointed out in Section 3, most properties which obey MNR exhibit positive or very small ΔS_{C_n} which should then not determine the sign of E_{n_n} . However, in Figure 5, the calculated E_n according to Equation (17) is 0.32 ± 0.06 eV for perovskite-type proton conductors and 0.38 ± 0.01 eV for Li-ion conductors, respectively, indicating that E_n differs among different types of ionic conductors and can also be large. This also requires further investigation.

7. Conclusion

It has been shown that MNR is the result of entropy-enthalpy compensation, and it has been proposed that multi-excitation entropy, MEE compensates the lack of thermal energy to overcome the energy barrier to ion transport in FICs. Within this framework, the dependence of Arrhenius prefactor on the activation energy may be applied to closely related systems, in which the material parameters, including mobile ion concentration, ion jump distance, and attempt frequency, do not differ greatly. Applying these constraints, we have investigated the roles of entropy, of isokinetic temperature, T_{iso} , and the contribution of lattice dynamics in the ion transport processes in FICs.

The value of operating temperature T compared to T_{iso} determines the strategy for improving the conductivity by tuning the activation energy. When $T < T_{iso}$, the conductivity can be improved by decreasing the activation energy. When $T > T_{iso}$, it can be improved by increasing it. Since MNR applies to closely related systems, caution must be taken to determine its value. T_{iso} may be determined experimentally, using an Arrhenius plot or a M-N plot, using Equation (2). The pressure-tuning method is an effective approach for measuring T_{iso} in FICs. Controlling its value remains challenging, due to the lack of theoretical models. The MEE model suggests that T_{iso} is proportional to the excitation phonon frequency, given by Equation (16). The general validity of this prediction, in particular its applicability to FICs continues to be investigated.

Finally, we have proposed that for FICs, the critical energy E_n , determined from the relationship between isokinetic prefactor

 σ_{00} and isokinetic temperature T_{iso} , is related to the configurational entropy change between the initial and transition sites. A positive E_n suggests that the charge carriers behave as polarons. The new understanding will add insights to develop new FICs.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

Acknowledgements

Q.C. and P.D. acknowledge support from the National Natural Science Foundation of China (Grant No. 52272227) and the Shanghai Natural Science Foundation (Grant No. 22ZR1428800). A.B. and Q.C. acknowledge support from the Swiss National Science Foundation (Grant No. 200021-188588). P.D. and Q.C. thank Shouhang Bo, Yanming Wang (Shanghai Jiao Tong University), and Donglin Han (Soochow University) for valuable suggestions.

Conflict of Interest

The authors declare no conflict of interest.

Keywords

fast ionic conductors, ion transport, lattice vibration, Meyer-Neldel rule, solid state ionics

Received: July 24, 2023 Revised: September 21, 2023 Published online: November 3, 2023

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