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Ion transport mechanism in anhydrous lithium thiocyanate LiSCN part III: charge carrier interactions in the premelting regime

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In lithium thiocyanate Li(SCN), the temperature regime below the melting point (274 °C) is characterized by excess conductivities over the usual Arrhenius behavior (premelting regime). Here, the Schottky defect pair concentration is high, and the point defect chemistry can no longer be considered as dilute. Coulomb interactions of Schottky pairs are expected to occur lowering the formation energy of new carriers and hence leading avalanche-like to a transition into a fully defective superionic state. The respective non-linear behavior is investigated using the cube-root law approach characterized by a defect interaction parameter J , which is a measure of the effective defect-lattice energy. In the case of Li(SCN), the rather pronounced volume expansion is to be included in the model. A literature comparison with other materials emphasizes to what degree defect formation as well as defect interactions depend not only on the dominant mobile defect, but also on the respective sublattice. Overall, a quantitative description of the defect chemistry of Li(SCN) in the premelting regime is derived.

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Introduction

It was shown that anhydrous Li(SCN) is Schottky disordered with lithium vacancies V_{Li} as mobile charge carriers.¹ Their mobility is affected by slow reorientation processes in the $(SCN)^-$ anion sublattice.² These findings explain ionic conductivities and their dependencies on doping and (impedance) frequency in the temperature range from 25 to 250 °C, provided that also trapping effects of the dopants are taken account of. In this temperature range the defect concentrations are still relatively low and the system can be regarded as dilute. Here, we investigate the defect chemistry above 250 °C close to the melting point (m.p.) of Li(SCN) in the so-called premelting regime, as well as beyond the melting point.

In literature, one can find many examples of anomalous (nonlinear) behavior close to a transition or melting point for various properties (conductivity,^{3–10} volume,^{11,12} strain,¹² spin-lattice relaxation,^{13,14} and others^{15,16}). In regard to conductivity, this is manifested as a nonlinear increase in the Arrhenius behavior, *i.e.* an excess conductivity. For AgBr,³ Schmalzried

attributed the excess conductivity to an increase of mobile defect concentration and mobility caused by the volume thermal expansion. Later investigations, however, indicated that volume changes cannot fully explain the increase in conductivity.⁴ Since defect concentrations are high near a transition, dilute theories (*e.g.* Debye–Hückel) are not well suited to explain the interactions of defects in this regime.

Therefore, Hainovsky and Maier derived an approach which describes this interaction by a Madelung energy of oppositely charged defects. This leads to a Coulombic stabilization of the system expressed by an interaction parameter J .^{4,17,18} This is applicable in particular for high defect concentrations, and results in a cube-root law (the excess free enthalpy scales with the cube-root of defect concentration). Since the free enthalpy of defect formation decreases, it results in an avalanche-like defect formation termed superionic transition. This transition occurs in a certain temperature range just below the actual phase transition (structural or melting). Within the cube-root model, a critical temperature can be defined at which the steepest increase of defect concentration occurs.⁴ Phenomenologically, a related onset temperature of the superionic transition can be determined from linear extrapolations of the low and high temperature regimes in the conductivity Arrhenius plot.

This model of defect interactions allows one to clearly distinguish the superionic state from the defective state, as illustrated in Fig. 1. A crystal can be defect-free at zero absolute temperature. Above 0 K the defect concentrations increase with

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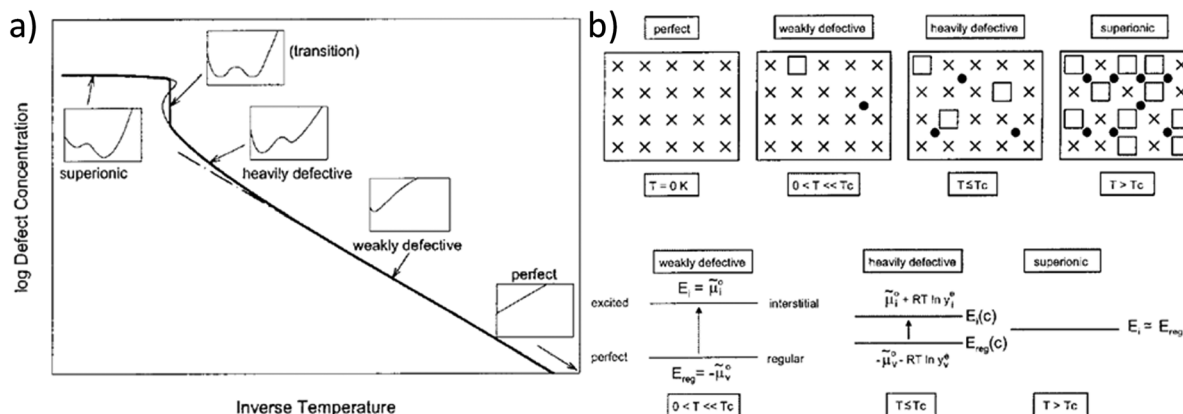


Fig. 1 Schematic representation of the superionic transition. (a) Schematic plot of defect concentration as a function of temperature. The insets for different regimes (perfect, weakly defective, heavily defective, transition, and superionic) give the respective curves of free enthalpy as function of defect concentration. (b) Crystallographic pictures and energy level diagrams for the different regimes. Reproduced from ref. 18 with permission of John Wiley & Sons.

temperature but the absolute defect concentrations remain small. This weakly defective state is characterized by a high defect formation enthalpy (energy gap between perfect and excited state, *e.g.* regular and interstitial site). This energy requirement decreases with increasing defect concentration (heavily defective) owing to the mutual Coulombic stabilization of the charged ionic defects. In the curve of defect formation free energy *versus* defect concentration (insets in Fig. 1a) a second minimum at high defect concentration develops and becomes increasingly deeper. Finally, in the superionic state the defect formation energy vanishes. The term “superionic dissociation” within an MX crystal designates the dissociation from the Madelung-bonded perfect MX lattice, irrespective of the fact that the “lattice molecules” MX are dissociated and ionized (M^+ and X^-). This defect interaction model predicts a transition to a superionic state at high temperatures for any crystal, if the crystal structure can be maintained. Then a criterion for the order of the transition could be derived.⁴ In most cases the transition is preceded by a phase transition into a different structure (*e.g.* α -AgI) that can better accommodate the high defect concentration (molten sublattice), or into a completely molten state (*e.g.* AgCl). The quantitative agreement of the model with measured data is best when the transition is of higher order (*e.g.* in PbF_2).

So far, very little is known about the premelting regime and melting of Li(SCN) at 274 °C without preceding phase transition. In contrast, K(SCN) has been extensively investigated by Schranz¹¹ and Plester *et al.*,¹⁰ and compared to related systems by Fuith.¹² K(SCN) undergoes a first order structural phase transition at 142 °C, which is characterized by an upward bending in the volume right before the transition.¹¹ For the nonlinear volume and conductivity changes in the premelting regime of K(SCN) (m.p. 175 °C) Plester *et al.* concluded that the mere volume thermal expansion is insufficient to explain the increased defect formation, and suggested an additional mechanism of defect stabilization by Coulomb interactions.¹⁰

Since thiocyanates exhibit significant thermal volume expansion due to the anisotropic motion of the $(SCN)^-$ anion,^{10,19} in this work

the cube-root law approach is employed to describe transport in the premelting regime of Li(SCN) integrating volume changes into the model. The ionic conductivity of Li(SCN) is measured by electrochemical impedance spectroscopy (EIS) close to and beyond its melting point. The data are fitted with the cube-root law approach in two ways: first in the conventional manner with a constant Coulomb interaction parameter J , and second by including the large volume expansion of Li(SCN) at high temperatures, resulting in a temperature dependence of J . The present application of the cube-root law to a Schottky defective material is compared to the Frenkel defective silver halides, and a general comparison of superionic transition temperatures of various material systems is made. The thermodynamic and kinetic data of Li(SCN) are compiled and compared to other materials to complete this extensive trilogy study (Parts I¹ to III) on anhydrous Li(SCN).

Results and discussion

The previously reported ion transport results of doped and undoped Li(SCN) in Part I¹ and II² were collected under conditions of low defect concentrations, *i.e.* a dilute defective system, where Coulomb defect–defect interactions between the native carriers can be neglected. However, near the melting point the material transforms into a concentrated defective system. In case of Frenkel defects, the cation sublattice becomes completely disordered (also referred to as “molten sublattice”) and the material becomes superionic.^{4,20} In the case of Schottky defects (in contrast to Frenkel disorder) both cation and anion sublattices are affected. This means that the defect concentrations can only increase until the defect lattice shows the same occupancy as the regular lattice (1/2 of all available lattice sites) before the material melts.

Li(SCN) exhibits a pronounced lattice expansion, which is a general feature for thiocyanates due to the local motion of the $(SCN)^-$ anions.^{10,19} Fig. 2 shows that for Li(SCN) the expansion is anisotropic (as observed for other thiocyanates),^{11,19} being strongest in the b direction. The volumetric expansion



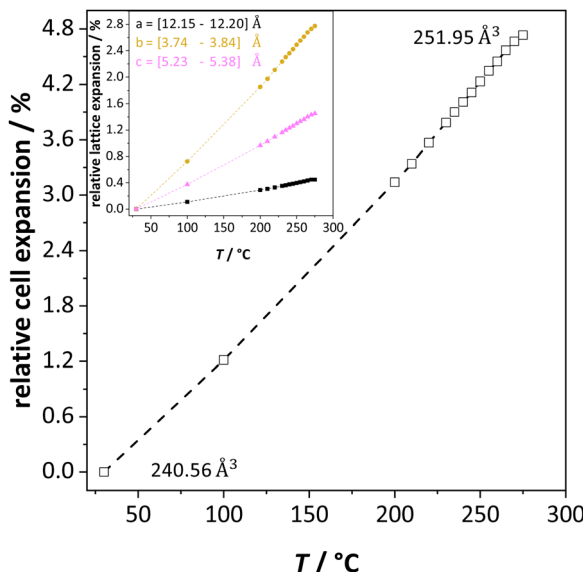


Fig. 2 Unit cell volume expansion of undoped Li(SCN) as a function of temperature. The insert shows the lattice expansion in directions *a* (black), *b* (orange) and *c* (magenta), respectively.

approaches 5% close to the melting point. This expansion is moderately larger than for LiI (4% from 25 °C to 270 °C).²¹ Nevertheless, it approximately follows a linear relation without anomaly up to the melting point.

In the following we will consider the impact of the relative cell volume expansion on the superionic phase transition in Li(SCN), and for this purpose ignore the anisotropy of the

lattice expansion. The volume expansion of Li(SCN) can be extracted from the data in Fig. 2:

$$\Delta V(T) = 4.60 \times 10^{-26} \text{ cm}^3 \text{ K}^{-1} \cdot T - 2.27 \times 10^{-22} \text{ cm}^3 \quad (1)$$

To investigate the defect chemistry of Li(SCN) within and above the premelting regime, impedance measurements were conducted near the phase transition as well as of molten Li(SCN). Investigations of molten thiocyanates had already shown that the (SCN)⁻ anions remain intact.²² Fig. 3a clearly shows that when approaching the melting point, the conductivity increases more strongly with temperature than expected from the Arrhenius relation. Conductivity data in this premelting regime were fitted with the cube-root law approach (details about the mathematical procedure are given in the Experimental).^{4,5,17,18,23} Although the concentrations of both lithium (V'_{Li}) and thiocyanate (V^*_{SCN}) vacancies are substantial in this premelting regime, the employed model considers only the contribution of V'_{Li} to the ionic conductivity. The mobility of V^*_{SCN} is too low to be significant. It would require a jump of the comparably large (SCN)⁻ anion, which is considered relevant only in the molten state. Since the phase transition is of first order, the transition occurs discontinuously, and a fit can only be conducted in the premelting regime (up to ≈ 270 °C, cf. Experimental for more information). In a first approach, the possible impact of the volume expansion was not included in the fitting, and the Coulomb interaction parameter J was taken to be temperature independent (Fig. 3, dark blue lines). With this approach the data in the premelting regime of solid Li(SCN) (Fig. 3a, full squares) could be fitted up to 261 °C (Fig. 3a, dark blue solid line).

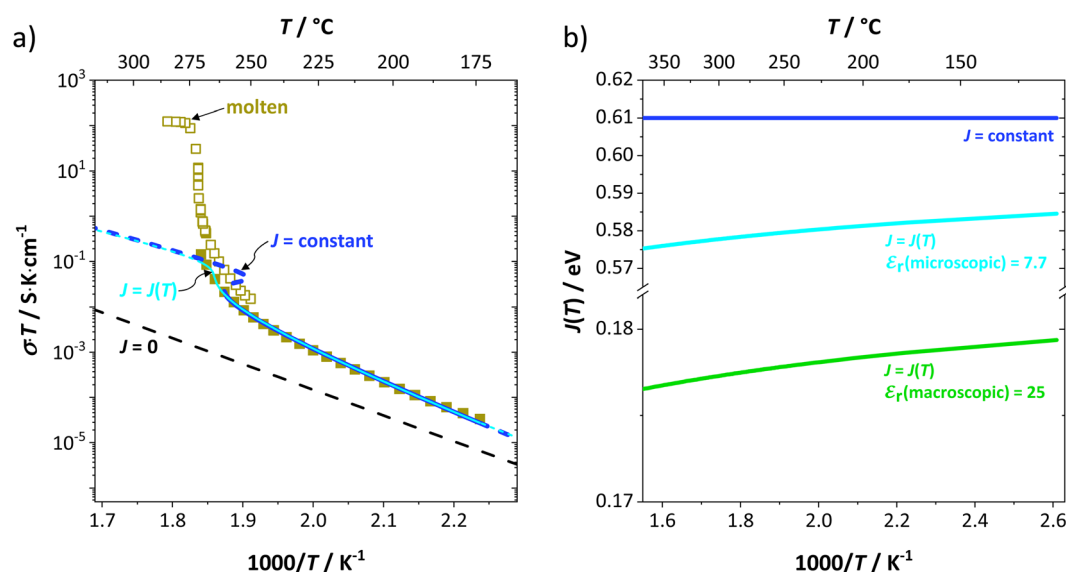


Fig. 3 (a) Measured conductivities of undoped Li(SCN) in the premelting regime (full squares, only every 4th data point is shown) and upon melting (open squares, only every 50th data point is shown). All lines correspond to non-linear fits of the premelting regime with $J = 0$ (black dashed line), $J = \text{constant}$ (dark blue solid and dashed line), and $J = J(T)$ (light blue solid and dashed line). For $J = \text{constant}$ the volume change was neglected, while for $J = J(T)$ the volume change was included using the Madelung approach with the microscopic dielectric constant $\epsilon_r(\text{microscopic})$. (b) Coulomb interaction parameter J shown for three different cases: (i) temperature independent $J = \text{constant}$ (dark blue solid line); temperature dependent $J = J(T)$ (green solid line), calculated with the Madelung approach using the macroscopic dielectric constant $\epsilon_r(\text{macroscopic})$, (ii) temperature dependent $J = J(T)$ (bright blue solid line) calculated with the Madelung approach using the microscopic dielectric constant $\epsilon_r(\text{microscopic})$.



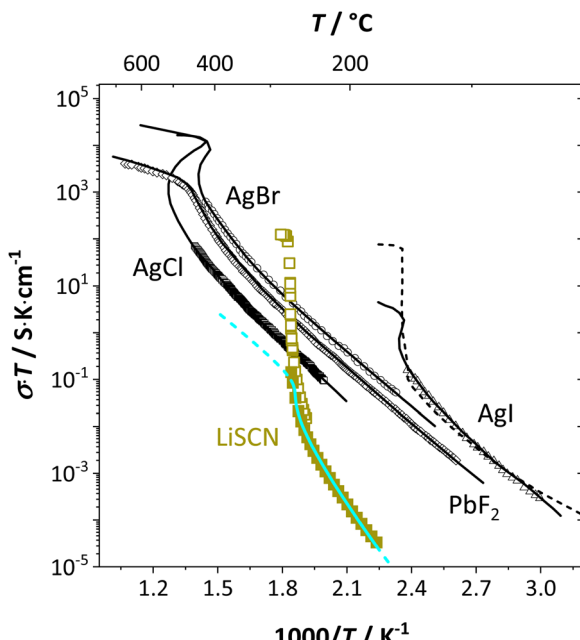


Fig. 4 Comparison of first and second order superionic phase transitions in different materials; brown squares and bright blue line for Li(SCN) (this study, cf. also Fig. 3a), black symbols and solid lines for PbF₂, AgCl, AgBr, AgI,⁴ and dashed line for AgI.²⁰

The resulting fitting parameters were used to simulate data beyond 261 °C (Fig. 3a, dark blue dashed line). However, in view of the remarkable volume effects, this approach was not able to successfully describe the complete premelting regime in Li(SCN).

In the second approach the volume expansion was included into the calculations, which makes the Coulomb interaction parameter temperature dependent, and $J = J(T)$. $J(T)$ describes Coulomb interactions and is related to the Madelung energy of the defect lattice. The Madelung energy is proportional to the reciprocal lattice constant, thus the inclusion of eqn (1) into the original cube-root model introduces a temperature dependence of J (i.e. $J(T) \propto 1/\text{lattice constant}(T)$). However, the calculated values for $J(T)$ from eqn (12) in the Experimental section are far too low to describe the Coulomb interaction in Li(SCN) (Fig. 3b, green solid line) when the high macroscopic dielectric constant of 25 for Li(SCN) is used (which reflects the strong polarizability of the (SCN)⁻ anion). Since the cube-root law approach is essentially an effective Madelung model with the “mean-field approach”, the overestimation of the (SCN)⁻ anion polarizability has to be corrected by employing a smaller, microscopic dielectric constant in the calculations (comparable to a system with a less polarizable anion, e.g. like LiF with $\epsilon_r \approx 9$).²⁴ In contrast to the macroscopic dielectric constant, the microscopic one describes the interaction of charged defects (V'_{Li} to V'_{SCN}) at atomistic distances. The macroscopic dielectric behavior may contain significant contributions from longer-range collective relaxations in a certain volume element (e.g. the octahedra in BaTiO₃). If the distance between interacting defects is reduced below such length scales (i.e. shrinks to atomistic distances), the dielectric constant is lowered to

a “microscopic” value. Such a rationale was also invoked e.g. for PbF₂ and SrTiO₃.^{23,25} A decrease of permittivity as a consequence of effects such as local distortions, proximity to charged defects, or decreasing grain size is well known and was investigated in detail e.g. for SrTiO₃.^{26,27} Therefore, ϵ_r (microscopic) was used as a parameter in the refinement with reasonable boundaries and $J(T)$ directly calculated to fit the data (Fig. 3a and b, light blue solid lines). With this approach considering the impact of the volume expansion the entire premelting regime up to 270 °C could be fitted.

From the data in Fig. 3a (full and open squares) it is obvious that melting of Li(SCN) occurs as a first order transition, as expected. This is also reflected by the simulated data above 261 °C in Fig. 3a (dark blue dashed line) using $J = \text{constant}$, which yields a similar swerve as obtained for AgCl and AgBr in Fig. 4. Interestingly, including the volume expansion changes the behavior of the simulated data, and it resembles rather a second order or diffuse phase transition (cf. PbF₂ in Fig. 4). This means that if Li(SCN) did not melt before, the occurring transition would be of second order when volume changes are considered. Overall, the superionic transition of Li(SCN) in the premelting regime is well described with the cube-root law (Fig. 3a).

The Coulomb interaction parameter J between the intrinsic $V'_{\text{Li}}/V'_{\text{SCN}}$ defects in Li(SCN) has a value of ~0.6 eV in the premelting regime. This is slightly higher than for β -AgI and lower than for AgBr.⁴ In the row of the silver halides (all with Frenkel disorder), J scales inversely with the size of the anion and becomes smaller when the lattice parameter is increased. Since Li⁺ is smaller than Ag⁺ but (SCN)⁻ is larger than I⁻, the resulting defect interactions seem to be comparable. The superionic transitions of Li(SCN) and β -AgI in Fig. 4 are rather similar in appearance. However, Li(SCN) is Schottky disordered, which means that in order to become superionic, both cation and anion sublattices must have a defect concentration similar to the number of lattice sites, and at least one must have a high mobility. As discussed in Part II,² the mobilities of cation and anion can be (and indeed often are) correlated, and in case of a Schottky disordered material it is therefore more likely that the completely molten state is more stable. In contrast, in Frenkel disordered β -AgI a stable situation can be reached in which only the cation lattice is molten (transition to α -phase).

The cube-root law application in Fig. 3a and 4 nicely describes the conductivity data right up to the transition point, after which a “jump” occurs. Here, the model cannot predict the behavior anymore, as mobility effects are not included. In contrast to the silver halides, for Li(SCN) the increase in volume had to be incorporated in the model. Large volume increases are often seen for systems with complex anions,^{12,28,29} and have also been observed in other thiocyanates.^{11,19}

One can find examples of superionic phase transitions for various ion conductors in literature, as shown in Fig. 5. This is not surprising, as every ionic crystal will enter some sort of regime where its defect concentration increases excessively,



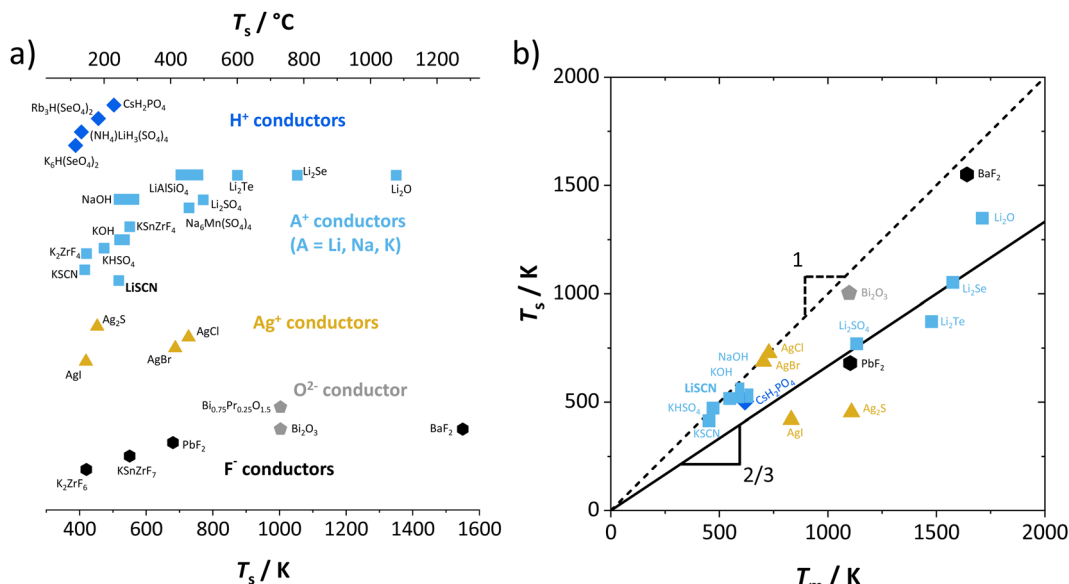


Fig. 5 Comparison of superionic phase transition temperatures for different materials with mobile ions such as F^- ,^{5,9} O^{2-} ,^{6,30} Ag^+ ,^{5,31} Li^+ ,^{7,15,32} Na^+ ,^{8,33} K^+ ,^{8,9,11,16} and H^+ .^{14,34-36} (a) Superionic transition temperatures T_s for various ionic conductors; (b) correlation of T_s for selected ion conductors from (a) with the respective melting temperatures T_m . The values of T_s and T_m were either directly taken from literature, or as phenomenological onset temperatures from respective data (intersection point from linear fits before and after transition (cf. Fig. 7), estimated error of ~ 10 K).

unless a first order transition occurs “earlier” (*i.e.* at lower temperature). A general trend for the temperature range of the superionic transition is difficult to discern. Overall, “soft” ionic systems with large polarizable ions such as Ag^+ , or with flexible hydrogen bonds in proton conductors tend to have lower transition temperatures.

Regarding the onset temperature of the superionic transition and the melting point (Fig. 5b, also *cf.* Fig. 3 in ref. 5), one can generally expect that for a Schottky defective material both temperatures are rather close. This is indicated in Fig. 5b by the dashed line with a slope of 1. As mentioned above, the necessary condition for a superionic transition is difficult to achieve for Schottky defects without the melt taking over as most stable state. Instead, in (anti-)Frenkel defective materials high defect concentrations do not necessarily mean a collapse of the lattice, and a superionic transition can occur before the melting point (note that melting of a Frenkel disordered material occurs if both sublattices being molten is the more stable state than only one

sublattice). The onset of the superionic transition often occurs at approximately 2/3 of the melting point (Fig. 5b, solid line with slope of 2/3). This is reminiscent of the Tammann rule (well known in solid state chemistry) stating that a material becomes reactive at roughly 2/3 of its melting temperature. Within the cube-root model, this can be rationalized from the fact that in this temperature range the defect formation approaches the superionic transition.⁵ The Tammann rule should in particular apply for Schottky disordered materials, since there both sublattices are involved in defect formation. The investigation of the defect chemistry in the premelting regime, *i.e.* superionic transition, might therefore also be indicative of the intrinsically formed defect types, which can help to identify the mobile defect if other methods such as doping are difficult to perform.

Finally, we can compare the thermodynamic and kinetic data of Li(SCN) from the experimental study in Part I¹ to the obtained values from the cube-root law fitting, and to literature data (Table 1). The Li(SCN) data sets agree reasonably well with

Table 1 Comparison of thermodynamic and kinetic data in the premelting regime of various materials using the cube-root law approach. ΔH° and ΔS° refer to the respective defect formation reactions (defect formation enthalpy and entropy), and $\Delta_m H_j$ and ν_j to defect migration of the defect j (migration enthalpy and pre-exponential factor). For the calculation of the parameters γ and γ_{crit} from which the order of the phase transition can be deduced, *cf.* ref. 4.

Compound	Type of defects		$\Delta H^\circ/\text{eV}$	$\Delta S^\circ/k_B$	$\Delta_m H_j/\text{eV}$	$\log_{10}[\nu_j/(\text{cm}^2 \text{ K s}^{-1} \text{ V}^{-1})]$	J/eV	$\gamma = J/\Delta H^\circ$	γ_{crit}	Ref.
PbF ₂	anti-Frenkel	Cube-root	1.08	8.48	0.18	2.94	0.75	0.69	0.78	4
AgCl	Frenkel	Cube-root	1.48	9.73	0.05	2.48	1.03	0.70	0.58	4
AgBr	Frenkel	Cube-root	1.15	7.67	0.15	3.22	0.80	0.70	0.65	4
β -AgI	Frenkel	Cube-root	0.82	11.77	0.24	1.60	0.50	0.61	0.58	4
Bi _{0.75} Pr _{0.25} O _{1.5}	anti-Frenkel	Cube-root	1.51	5.78	0.90	2.01	1.78	1.18	1.34	6
					0.48					
Li(SCN)	Schottky	Long-range hopping	0.6 ± 0.3	5 ± 2	0.89 ± 0.08	5.1 ± 0.4				1
		Local hopping			0.6 ± 0.1	2 ± 1				2
		Cube-root J	0.6	3.0	0.8	3.5	0.61	1.02	0.96	This work
		Cube-root $J(T)$	0.8	7.9	0.7	2.4	0.57–0.59	0.72–0.73	0.70	This work

each other within the experimental uncertainties. The largest deviations are observed for the entropy terms, which reflects the error-proneness of this quantity. The comparison of the defect formation enthalpy ΔH° and Coulomb defect–defect interaction parameter J reflects the types of defects, and, more importantly, the overall chemistry of the materials, which also involves the other sublattice. The most important factor is the ion charge, which explains the difference between the oxygen vacancy conductor $\text{Bi}_{0.75}\text{Pr}_{0.25}\text{O}_{1.5}$ and the other materials. The respective sublattice composition is determining for defect–defect interactions, evident by comparing the silver halides for which J decreases with increasing anion size. However, these considerations only apply to the defect formation thermodynamics. Regarding $\text{Li}(\text{SCN})$, the values of ΔH° and J suggest a similar defect chemistry to the silver halides (despite being Schottky and not Frenkel disordered). However, the enthalpy of migration for long-range ion transport in $\text{Li}(\text{SCN})$ is much higher compared to the silver halides, and almost the same as for $\text{V}_\text{O}^{\bullet\bullet}$ migration. This strong mobility suppression was addressed in Part II,² and reflects the occurrence of slow reorientational processes of $(\text{SCN})^-$ during cation jumps. This is the origin of the very different transport properties of $\text{Li}(\text{SCN})$. Table 1 nicely shows that the cube-root law approach can be applied to (anti-)Frenkel as well as Schottky defective materials, yet some additional, material specific considerations might have to be included (e.g. anisotropic ion transport in $\text{Bi}_{0.75}\text{Pr}_{0.25}\text{O}_{1.5}$ or lattice expansion in $\text{Li}(\text{SCN})$). $\text{Li}(\text{SCN})$ demonstrates how important specific effects can be for ion transport, *i.e.* the very different Li–N and Li–S interaction strengths of an anisotropic $(\text{SCN})^-$ anion.

Conclusion

The cube-root law for quantifying Coulomb defect–defect interactions in a concentrated system was successfully applied to the superionic transition in the premelting regime of anhydrous $\text{Li}(\text{SCN})$ to describe the non-linear conductivity increase. Two fitting models were employed; in one the Coulomb interaction parameter J is constant, while in the other it is temperature dependent to account for the substantial lattice volume expansion in $\text{Li}(\text{SCN})$. The inclusion of volume expansion enabled a smoother and more stable data fit without increasing the number of fitting parameters. The inclusion of volume changes, however, was not able to account for the complete conductivity increase upon melting, as additional mobility effects are not included. Together with literature examples, these investigations show that the cube-root law approach is very useful to better understand the behavior of mobile defects, phase transitions, and even general material properties such as reactivity at elevated temperatures and impact of lattice expansion.

Experimental

General

Synthesis, sample preparation, measurement conditions and set-ups were reported in Part I¹ of this series on anhydrous $\text{Li}(\text{SCN})$.

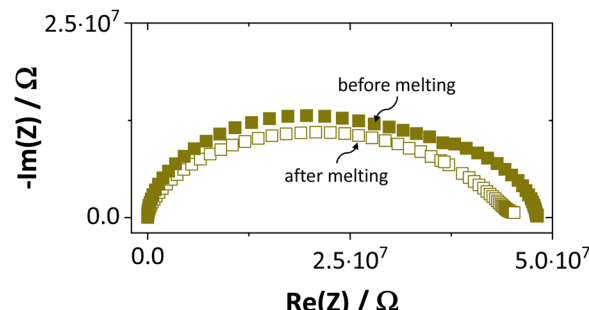


Fig. 6 Exemplary impedance spectra of anhydrous $\text{Li}(\text{SCN})$ at 134 °C before and after melting.

Temperature dependent X-ray diffraction measurements on $\text{Li}(\text{SCN})$ were performed according to the procedure described in the Supporting Information of Part I.¹ Regarding electrochemical impedance spectroscopy, some additional remarks have to be made. At temperatures close to the melting point (premelting regime) of $\text{Li}(\text{SCN})$, the EIS measurement conditions were changed compared to the ones reported in Part I. Measurements were taken continuously with 1 K steps without equilibration for 24–30 times. Measurements above the m.p. could not be conducted under high vacuum, since the material would start to sublime. Therefore, both the drying as well as EIS measurements were conducted under constant flow of dry Ar or N_2 gas (typically 50–100 sccm). Due to the phase transition (melting point), a different measurement cell geometry was used for these measurements. $\text{Li}(\text{SCN})$ powder was filled into a fused silica crucible equipped with Pt electrodes to measure the impedance directly from the molten or resolidified sample. The cell constant was determined with aqueous KCl solutions. For these measurements at high temperatures, the stray impedance of the set-up (measured separately by short-circuiting the electrodes) interfered, and therefore had to be pointwise subtracted. Exemplary impedance spectra before and after melting are shown in Fig. 6.

The onset temperature of a superionic transition was determined by linearly fitting the conductivity data in the Arrhenius plot before and after the transition, as shown in Fig. 7. Since the fitting range is set by visual inspection, values obtained with this method can vary to some degree (estimated error up to ~ 10 K). However, this uncertainty is negligible relative to the temperature range considered for the materials comparison in Fig. 5.

Cube-root law model fitting

Conductivity data of $\text{Li}(\text{SCN})$ between 174 and 270 °C (premelting regime: 242–270 °C) were fitted with the cube-root law described in literature.^{4–6,17,18,23} The fitting was conducted with Matlab (Mathworks, Version R2017a) using the lsqnonlin-function. A function handle was created in which the implicit equation for defect concentration was solved iteratively with the vpasolve-function at every temperature step. All parameters had to have reasonable boundaries for the fit to converge. The fits had a low residual sum of squares in the range of ~ 0.1 or smaller. The impact of lattice expansion in the premelting

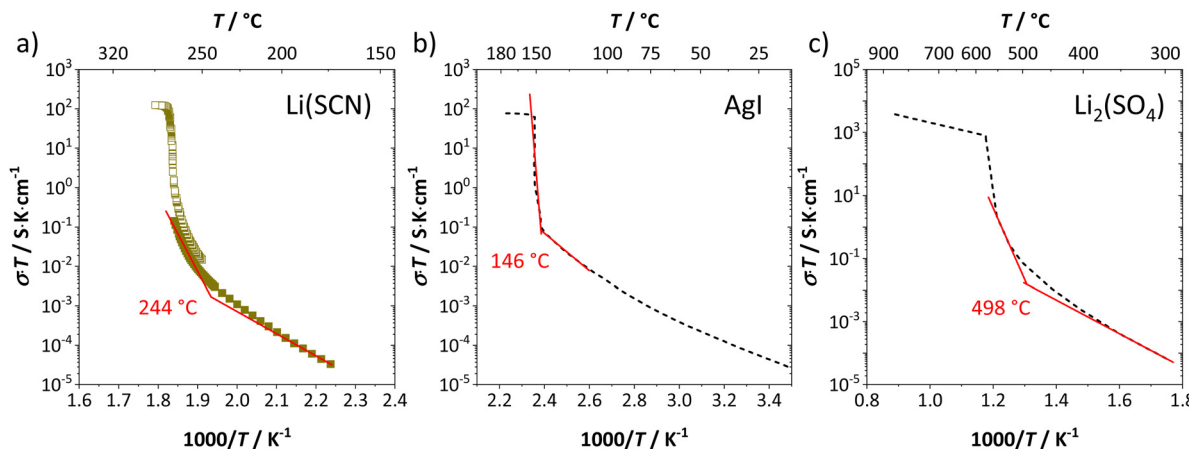


Fig. 7 Exemplary procedure to determine the superionic transition of (a) Li(SCN), (b) AgI,^{4,20} and (c) Li₂(SO₄).^{7,37} Symbols and black dashed lines are measured data, and red solid lines are linear fits before and after the transition. The shown temperatures are the obtained onset temperatures of the superionic transition.

regime for defect–defect interactions was considered and compared to fitting with constant J , as discussed in the main text.

Li(SCN) is a Schottky defective material. Accordingly, in the intrinsic regime we have:



and

$$K_{\text{S}} = [\text{V}'_{\text{Li}}][\text{V}^{\bullet}_{\text{SCN}}] = N_{\text{Li}}N_{\text{SCN}} \cdot \exp\left(\frac{\Delta_{\text{S}}S^{\circ}}{k_{\text{B}}}\right) \exp\left(-\frac{\Delta_{\text{S}}H^{\circ}}{k_{\text{B}}T}\right) \quad (3)$$

where K_{S} is the Schottky equilibrium constant for the regime before premelting, $[\text{V}'_{\text{Li}}]$ and $[\text{V}^{\bullet}_{\text{SCN}}]$ the respective vacancy concentrations, $N_{\text{Li}} = N_{\text{SCN}}$ are the number of available sites in the lattice, $\Delta_{\text{S}}S^{\circ}$ and $\Delta_{\text{S}}H^{\circ}$ the standard Schottky entropy and enthalpy of formation, k_{B} the Boltzmann constant, and T the temperature.

One can redefine N_{Li} into $N_{\text{Li}} = g_{\text{Li}} \cdot \alpha_{\text{Li}}$, in which g_{Li} is the degeneracy of Li sites and α_{Li} the number of occupiable Li sites. If one defines g_{Li} and α_{Li} per unit cell ($Z = 4$),³⁸ $N_{\text{Li}} = 1.4$, and if one defines it per lattice site, $N_{\text{Li}} = 1.1$. In the linear regime ($T \leq 242$ °C) the system is considered dilute and Maxwell–Boltzmann statistics apply. However, in the (non-linear) premelting regime with 242 °C $\leq T \leq 270$ °C the system becomes concentrated and Fermi–Dirac statistics have to be used (within the Brouwer approximation, $[\text{V}'_{\text{Li}}] = [\text{V}^{\bullet}_{\text{SCN}}]$):^{4,18}

$$N_{\text{Li}} = g_{\text{Li}} \left(\alpha_{\text{Li}} - \sqrt{K_{\text{S}}} \right) \quad (4)$$

and $\sqrt{K_{\text{S}}}$ can be regarded as the equilibrium defect concentration, $\sqrt{K_{\text{S}}} \equiv c_{\text{E}}$.

In a concentrated (superionic) defective state Coulomb interactions between oppositely charged defects become significant. Therefore, a Coulomb interaction term $\mu_{\text{int}}(c_{\text{E}})$ ($\mu_{\text{int}}(c_{\text{E}}) < 0$, since the Coulomb interaction is stabilizing) is introduced; $\Delta_{\text{S}}G^{\circ} = \Delta G^{\circ} + \mu_{\text{int}}(c_{\text{E}})$. It was suggested that this Coulomb interaction follows a cube-root law dependence yielding:⁴

$$\mu_{\text{int}}(c_{\text{E}}) = -Jc_{\text{E}}^{1/3} \quad (5)$$

in which J is the Coulomb interaction parameter.

The final equation, which was used to iteratively calculate the defect concentrations of Li(SCN) at respective temperatures in Matlab with vpasolve (c_{E} as symbolic variable) can be expressed as:

$$\frac{c_{\text{E}}^2}{g_{\text{Li}}^2(\alpha_{\text{Li}} - c_{\text{E}})^2} = \exp\left(-\frac{(\Delta G^{\circ} - Jc_{\text{E}}^{1/3})}{k_{\text{B}}T}\right) \quad (6)$$

The resulting defect concentrations are dimensionless (number of defects per lattice site) and were consequently converted into units of cm^{-3} with $c'_{\text{E}} = c_{\text{E}} \cdot (4/V_{\text{UC}})$, where V_{UC} is the unit cell volume of Li(SCN) ($V_{\text{UC}} = 240.56 \times 10^{-24} \text{ cm}^3$).³⁸

The mobilities were calculated according to:

$$u_{\text{V}'_{\text{Li}}} T = 10^{\log_{10}\left(\nu_{\text{V}'_{\text{Li}}}\right)} \cdot \exp\left(-\frac{\Delta_{\text{m}}H_{\text{V}'_{\text{Li}}}}{k_{\text{B}}T}\right) \quad (7)$$

and

$$\nu_{\text{V}'_{\text{Li}}} = \frac{r_{\text{Li}}^2 e}{Nk_{\text{B}}} \nu_0 \cdot \exp\left(\frac{\Delta_{\text{m}}S_{\text{V}'_{\text{Li}}}}{k_{\text{B}}}\right) \quad (8)$$

with lithium vacancies being the dominant mobile defects. In eqn (7) and (8) $u_{\text{V}'_{\text{Li}}}$ is the mobility, $\nu_{\text{V}'_{\text{Li}}}$ the pre-exponential factor, $\Delta_{\text{m}}H_{\text{V}'_{\text{Li}}}$ and $\Delta_{\text{m}}S_{\text{V}'_{\text{Li}}}$ are the enthalpy and entropy of migration, $r_{\text{V}'_{\text{Li}}}$ and N are the distance to and the number of available neighboring sites, e is the charge of an electron and ν_0 is the jump attempt frequency of lithium vacancies.³⁹ The correction term for mobility in concentrated systems proposed in ref. 4 did not lead to improvements in the fitting and was therefore neglected. Eqn (6) and (7) were combined to calculate



the conductivity $\sigma_{V'_{\text{Li}}}$ according to:

$$\sigma_{V'_{\text{Li}}} T = z_{V'_{\text{Li}}} e \cdot u_{V'_{\text{Li}}} T \cdot c'_{\text{E}} \quad (9)$$

where $z_{V'_{\text{Li}}}$ is the valence of V'_{Li} (equal to 1 for Li^+).

With the above set of equations, the cube-root model can be used to fit conductivity data as a function of $1000/T$ using a total of 5 parameters: $\Delta_{\text{S}}S^\circ$, $\Delta_{\text{S}}H^\circ$, J , $\log_{10}(\nu_{V'_{\text{Li}}})$ and $\Delta_{\text{m}}H_{V'_{\text{Li}}}$. The described approach (J is a constant parameter) was capable of fitting conductivity data up to 261 °C.

However, at higher temperatures the model failed to describe the data. A possible reason for this failure was the neglect of volume expansion in $\text{Li}(\text{SCN})$ in the premelting regime. The volume expansion $\Delta V(T)$ was derived from *in situ* XRPD data yielding:

$$\Delta V(T) = 4.60 \times 10^{-26} \text{ cm}^3 \text{ K}^{-1} \cdot T - 2.27 \times 10^{-22} \text{ cm}^3 \quad (10)$$

In this case, J is no longer a parameter and obtained from fitting the data, but instead directly calculated with material specific physical quantities related to the Madelung energy. To include the volume expansion in the fitting, the following approach was used:^{5,18}

$$J = \frac{4}{3\epsilon_r} \cdot U_{\text{M}} \cdot \frac{f_{\text{d}}}{f} = \frac{e}{3\pi\epsilon_0\epsilon_r} \cdot f_{\text{d}} \cdot \frac{1}{\varrho} \quad (11)$$

Here U_{M} is the Madelung energy, f_{d} is the effective Madelung constant of the defect superlattice (~ 0.73),⁵ f is the Madelung constant, ϵ_0 and ϵ_r are the electrical permittivities (dielectric constants) of free space and $\text{Li}(\text{SCN})$, and ϱ is the distance between nearest neighbors. Two simplifications are introduced: (i) the volume expansion is taken to be isotropic, and (ii) the orthorhombic lattice of $\text{Li}(\text{SCN})$ is considered to be pseudo-cubic for Coulomb interactions between neighbors, which means $\varrho_{\text{pc}}(T) = \sqrt[3]{\Delta V(T)}/2$. The impedance measured dielectric constant (~ 25 in the premelting regime) is a macroscopic magnitude ($\epsilon_r(\text{macroscopic}) = \epsilon_r^{\text{ma}}$), however, for defect-defect interactions the microscopic dielectric constant ($\epsilon_r(\text{microscopic}) = \epsilon_r^{\text{mi}}$) has to be considered.^{23,25} Finally, we derive the following equation:

$$J(T) = \frac{e}{3\pi\epsilon_0\epsilon_r^{\text{mi}}} \cdot f_{\text{d}} \cdot \frac{1}{\varrho_{\text{pc}}(T)} \quad (12)$$

to calculate the Coulomb interaction parameter while including the impact of volume expansion with temperature. In this approach J becomes a temperature dependent variable $J(T)$ and the fifth parameter is now ϵ_r^{mi} . With the inclusion of $\Delta V(T)$ the fitting could be performed successfully up to 270 °C.

Author contributions

All authors have contributed to the experimental results, calculations and writing of the manuscript, and have given their approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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