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# Ion transport mechanism in anhydrous lithium thiocyanate LiSCN Part I: ionic conductivity and defect chemistry†

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This work reports on the ion transport properties and defect chemistry in anhydrous lithium thiocyanate Li(SCN), which is a *pseudo-halide*  $\text{Li}^+$  cation conductor. An extensive doping study was conducted, employing magnesium, zinc and cobalt thiocyanate as donor dopants to systematically vary the conductivity and derive a defect model. The investigations are based on impedance measurements and supported by other analytical techniques such as X-ray powder diffraction (XRPD), infrared (IR) spectroscopy, and density functional theory (DFT) calculations. The material was identified as Schottky disordered with lithium vacancies being the majority mobile charge carriers. In the case of  $\text{Mg}^{2+}$  as dopant, defect association with lithium vacancies was observed at low temperatures. Despite a comparably low Schottky defect formation enthalpy of  $(0.6 \pm 0.3)$  eV, the unexpectedly high lithium vacancy migration enthalpy of  $(0.89 \pm 0.08)$  eV distinguishes Li(SCN) from the chemically related lithium halides. A detailed defect model of Li(SCN) is presented and respective thermodynamic and kinetic data are given. The thiocyanate anion ( $\text{SCN}^-$ ) has a significant impact on ion mobility due to its anisotropic structure and bifunctionality in forming both Li–N and Li–S bonds. More details about the impact on ion dynamics at local and global scale, and on the defect chemical analysis of the premelting regime at high temperatures are given in separate publications (Part II and Part III).

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## 1. Introduction

With the growing interest in solid state lithium ion batteries, solid lithium ion conductors are intensively studied, and significant progress has been achieved.<sup>1–5</sup> A common strategy is to create a cation lattice with a flat potential landscape (high defect concentration and low activation energy), *e.g.* by choosing a highly polarizable anion lattice.<sup>6</sup> A famous example is the  $\text{Li}_2\text{S}$ – $\text{P}_2\text{S}_5$  binary system, which contains several crystalline phases, such as  $\text{Li}_3(\text{PS}_4)$  and  $\text{Li}_2(\text{P}_2\text{S}_6)$  as well as glass ceramics.<sup>5</sup> These compounds are essentially *pseudo-binaries* with the general formula  $\text{Li}_x\text{A}$  ( $\text{A} = \text{P}_y\text{S}_{2y+2}$ ). In contrast to

normal binaries, the molecular (complex) anion has an internal structure and can coordinate to  $\text{Li}^+$  in different ways. The coordination becomes most complex when the cation can interact with different elements of the anion ligand, *e.g.* in  $(\text{CN})^-$ ,  $(\text{OCN})^-$ ,  $(\text{SCN})^-$ ,  $(\text{N}(\text{CN})_2)^-$  or  $(\text{C}(\text{CN})_3)^-$ , often with different strengths of interaction. This molecular coordination chemistry is well established in liquid systems, but poorly studied for solids. Therefore, the investigation of Li(SCN) was motivated by its unique cation – to – anion coordination chemistry, which results in peculiar ion transport characteristics. In this sense, Li(SCN) serves as a model material which provides new insight into the parameters which affect ionic conductivity.

In terms of defect chemistry, some of the better understood binary lithium systems include  $\text{LiH}$ ,<sup>7</sup>  $\text{Li}_3\text{N}$ ,<sup>8,9</sup> lithium chalcogenides,<sup>10,11</sup> and lithium halides.<sup>12–14</sup> LiI has become an established additive for performance enhancement in various battery systems,<sup>15–19</sup> and a number of fundamental studies helped to understand its role as an additive.<sup>20–25</sup> Lithium thiocyanate (often referred to as a *pseudo-halide*) is chemically and structurally similar to lithium iodide, since they both have a large polarizable anion,<sup>26,27</sup> are hygroscopic and form hydrates.<sup>28–31</sup> However, in contrast to LiI and other halides, little is known about the defect

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chemistry of Li(SCN). Previous ion transport studies predominantly focused on composites as potential battery electrolytes including liquid electrolytes,<sup>32,33</sup> polymer blends,<sup>34–37</sup> or inorganic salt eutectics.<sup>38,39</sup> Transport properties of pure Li(SCN) were (so far) investigated only by Poulsen.<sup>30</sup> The fact that Li(SCN) does not follow the trend in conductivities of lithium halides with anion size (suggesting that it would have the highest value in the row LiF–LiCl–LiBr–LiI–Li(SCN)) indicates the significance of its singular coordination chemistry, and motivates us to elucidate the origins.

In the present publication, we focus on ion transport and defect chemistry. Electrochemical measurements and doping experiments show Li(SCN) to be Schottky defective and lithium vacancies  $V'_{\text{Li}}$  to be the majority charge carriers. We will first discuss the doping experiments to identify the mobile carrier, then identify the defect-chemical regimes (Brouwer diagrams) describing the defect chemistry semi-quantitatively, and finally extract quantitative data with respect to formation and migration energies and even mobilities and concentrations. These findings contribute to a better understanding of the transport behavior in electrolytes with complex anions (bidentate ligands). All experimental and computational details are given in the ESI.† In Part II of this series of publications we discuss the frequency dependence of the ion conductivity in Li(SCN),<sup>40</sup> and Part III reports on the defect chemistry in the premelting regime.<sup>41</sup>

## 2. Results and discussion

### 2.1. Doping experiments

Anhydrous Li(SCN) is stable beyond its melting point at 274 °C and does not decompose up to at least 550 °C.<sup>42</sup> Compacted pellets of nominally pure (undoped) Li(SCN) were measured both by electrochemical impedance spectroscopy (EIS) and by

direct current (DC) experiments. The first task was to identify the ionic or electronic nature of the conductivity. From an electromotive force (EMF) measurement with Li and LiAl electrodes at 110 °C, a Li<sup>+</sup> transference number of  $t_{\text{Li}^+} \geq 0.99$  (ESI† Fig. S8b) was estimated. In this configuration, most probably a solid electrolyte interface (SEI) layer of thermodynamically stable compounds formed, which kinetically stabilizes the Li(SCN) – Li metal interface (thermodynamically, taking the values of the analogous sodium compounds as a reference,<sup>43</sup> one expects an exothermic reaction with the conversion of Li(SCN) and Li metal into Li<sub>2</sub>S and Li(CN)). This measurement shows that Li(SCN) is a Li<sup>+</sup> cation conducting material with negligible electronic contributions. Correspondingly, impedance spectra with Li<sup>+</sup> blocking ruthenium electrodes show a Warburg-type low frequency feature at elevated temperatures at which the respective frequencies fall into the measurement range (ESI† Fig. S9). Note that the measured conductivities of a previous study (ESI† Fig. S13a) are higher than the present values due to incomplete drying.<sup>42</sup>

Aliovalent bulk doping is a well-established method to identify the nature of mobile defects (vacancies or interstitials). However, one has to stay within the solubility limit of the dopant, as otherwise formed secondary phases might also affect the conductivity. Mg<sup>2+</sup>, Zn<sup>2+</sup>, and Co<sup>2+</sup> are suitable cations to investigate the solid solution behavior of Li<sup>+</sup> conductors, given their chemical and size similarity (cf. ESI† Fig. S6). ICP-OES analysis confirmed the overall dopant concentration to be close to the nominal values (ESI† Fig. S7). Owing to the low dopant concentrations and the size similarity, it is not surprising that there is no clear trend in the comparison of the unit cell volumes (ESI† Fig. S6b). Fig. 1 and Fig. S6a (ESI†) show that Mg<sup>2+</sup>, Zn<sup>2+</sup>, and most likely also Co<sup>2+</sup> are soluble in Li(SCN) up to about 3–5 mol%. The changes in conductivity – discussed

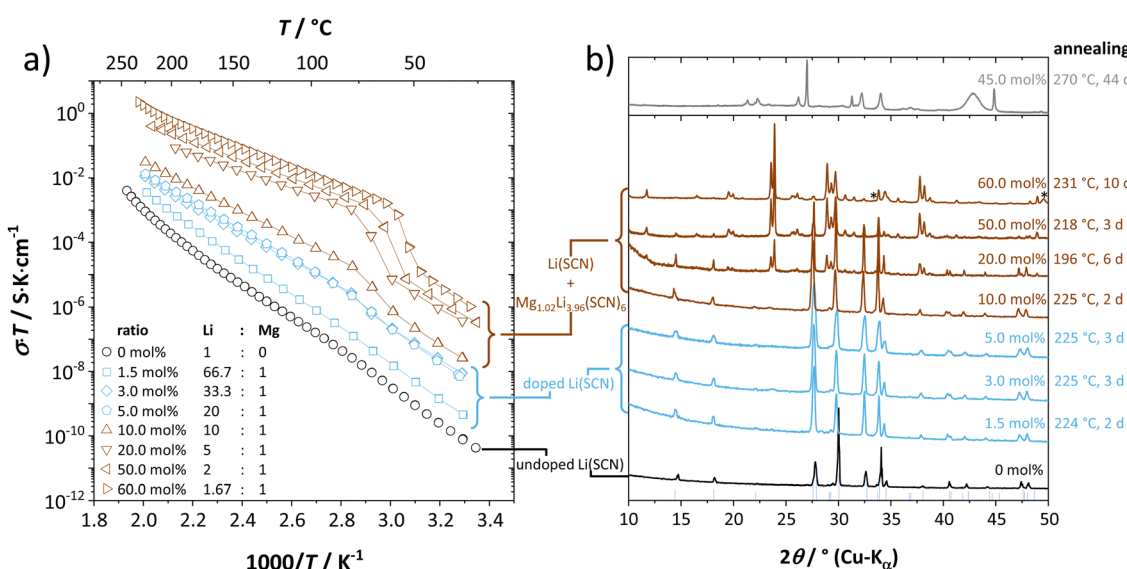
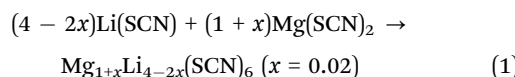


Fig. 1 Doping of Li(SCN) with Mg(SCN)<sub>2</sub>: nominally undoped Li(SCN) (black), Mg<sup>2+</sup>-doped Li(SCN) (blue) and two-phase samples of Li(SCN) and  $\text{Mg}_{1.02}\text{Li}_{3.96}(\text{SCN})_6$  (dark red). (a) Conductivities as a function of inverse temperature, and (b) XRD patterns. Reflections marked with an asterisk belong to an unknown side phase (most likely a decomposition product of Mg(SCN)<sub>2</sub>·4 H<sub>2</sub>O), and the blue bars at the bottom represent anhydrous Li(SCN) as a reference.<sup>31</sup>



in more detail in section 2.2 – indicate that  $Mg^{2+}$ ,  $Zn^{2+}$ , and  $Co^{2+}$  are indeed incorporated and active as dopants.

For  $Mg^{2+}$  concentrations  $\geq 10$  mol%, reflections of a new phase appear in the diffractograms (cf. Fig. 1b and ESI† Fig. S1). EIS and temperature dependent *in situ* XRPD<sup>42</sup> showed that this new phase has a structural transition between 47 and 62 °C into a high temperature modification with higher symmetry. The phase transition can also be recognized from the shape of the impedance spectra as well as the dielectric constant  $\epsilon_r$  (cf. ESI† Fig. S10). Due to synthetic difficulties and limitations of powder diffraction, an *ab initio* crystal structure solution was only possible of the high temperature modification, which was performed at 55 °C (longer synthesis times at high temperatures even lead to a highly disordered, different material; cf. Fig. 1b, grey pattern). The structure of lithium-magnesium thiocyanate indicates a variable lithium-to-magnesium ratio, and the final Rietveld<sup>44</sup> refinement yielded the following composition (cf. ESI† Fig. S1):



Apart from recently prepared tri-cationic cyanamides with the composition  $Li_2MSn_2(NCN)_6$  ( $M = Mg^{45}$  und  $Mn^{46}$ ), anhydrous thiocyanates, nitrides or cyanides of lithium and any divalent transition metal are unknown so far, which makes the structure of  $Mg_{1.02}Li_{3.96}(SCN)_6$  a *hitherto* unknown structure type (details in the ESI†). Mixed alkali and alkaline earth metal *pseudo*-halides are rare,<sup>47–49</sup> and even though  $Na_4Mg(SCN)_6$  has a very similar composition, its crystal structure is very different. The observation that the new phase  $Mg_{1.02}Li_{3.96}(SCN)_6$  has a much higher conductivity than even highly doped  $Li(SCN)$  is related to the coordination polyhedra of  $Mg^{2+}$  and  $Li^+$  in this structure (ESI† Fig. S3 and S4), and will be discussed later.

The impedance spectra and temperature dependence of  $\epsilon_r$  for both undoped and doped  $Li(SCN)$  (Fig. 2 and ESI† Fig. S12) are

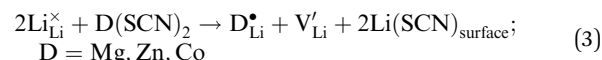
obviously more complex compared to other  $Li^+$  ion conductors.<sup>10,11,23</sup> This complexity is rooted in the frequency dependent conductivity of  $Li(SCN)$  and will be discussed in detail in Part II.<sup>40</sup> In the present work, only the DC resistance (low frequency intersection with the real axis) is of importance, thus only the low frequency part of the spectra was fitted with the circuit shown in Fig. 2.

## 2.2. Defect chemical model

In order to investigate the defect chemistry of single-phase  $Li(SCN)$ , dopant concentrations of 5 mol% were not exceeded. The ionic conductivity of undoped  $Li(SCN)$  shows two slope changes as indicated in Fig. 3a. Such behavior is common for many ion conductors (e.g.  $LiH$  and  $Li_2S$ ),<sup>7,10</sup> and is typically used as a basis for assigning three regimes: I intrinsic, II extrinsic, and III association. In the intrinsic regime the conductivity is dominated by the material's native defects. Since  $Li(SCN)$  is considered a *pseudo*-halide with a densely packed structure, one can expect the material to be Schottky defective:<sup>13,14</sup>



(Kröger-Vink notation). Since the material is a  $Li^+$  cation conductor, the dominant mobile defect should be lithium vacancies  $V'_{Li}$ . The transition to the extrinsic regime is marked by a more shallow slope in Fig. 3a. In the extrinsic regime the mobile carrier concentration is fixed by a dopant  $D_{Li}^{\bullet}$  (or possible donor-type impurity). The concentration of  $V'_{Li}$  can be increased by donor doping, e.g. with a bivalent cation of suitable ionic radius such as  $Mg^{2+}$ ,  $Zn^{2+}$  or  $Co^{2+}$  (Fig. 3b):



The conductivity increases with the concentration of all  $D^{2+}$  dopants (Fig. 3b and Fig. S14, ESI†), which shows that indeed  $V'_{Li}$  are the dominant mobile defects. Acceptor doping of the  $(SCN)^-$  anion with  $Li_2S$  and  $Li_2(SO_4)$  was attempted, yet despite

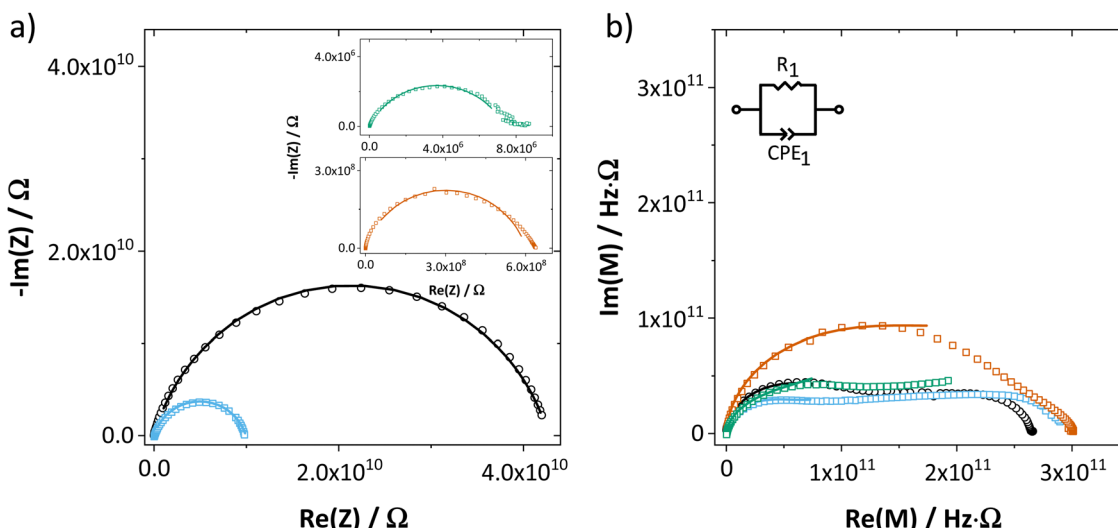


Fig. 2 (a) Impedance and (b) complex modulus of undoped  $Li(SCN)$  (black circles), as well as 1.5 mol% doped  $Li(SCN)$  (squares) using  $Mg^{2+}$  (blue),  $Zn^{2+}$  (red) and  $Co^{2+}$  (green) as dopant ( $T = 68$ –69 °C). The impedance spectra were fitted with the shown equivalent circuit.

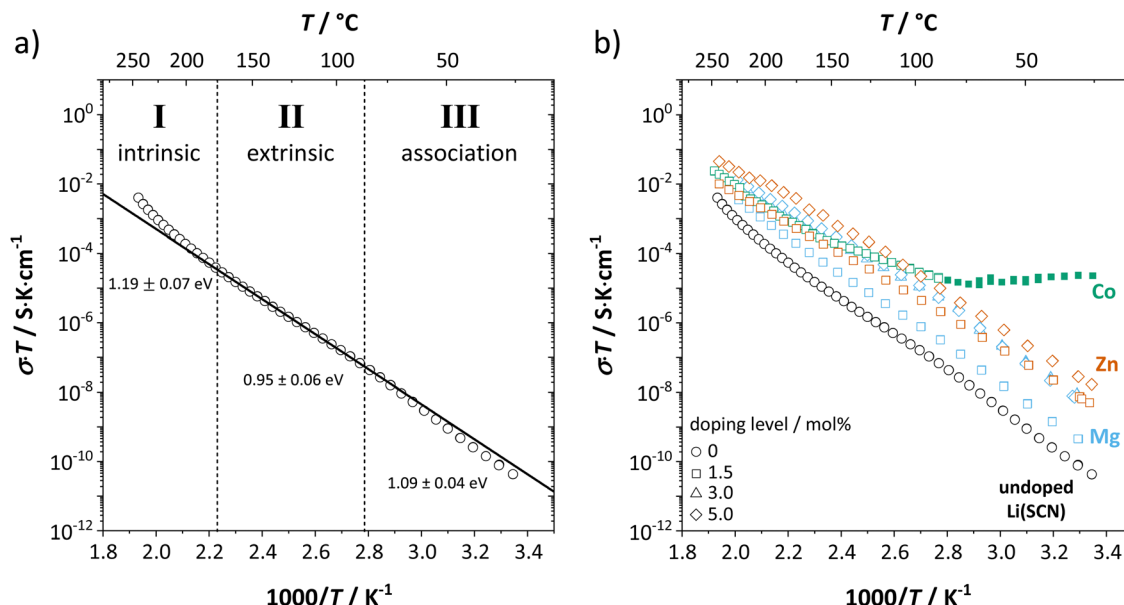
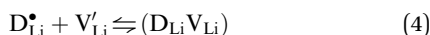


Fig. 3 (a) Assignment of defect chemical regimes in undoped Li(SCN). (b) Comparison of conductivities from undoped Li(SCN) (black), Mg<sup>2+</sup>-doped (blue), Zn<sup>2+</sup>-doped (red) and Co<sup>2+</sup>-doped Li(SCN) (green). The green full symbols correspond to the temperature range where for Co<sup>2+</sup>-doped Li(SCN) electronic conductivity prevails ( $-0.11 \text{ eV}$  apparent activation energy).

employing different preparation methods was unsuccessful, as indicated by the unchanged conductivities (ESI† Fig. S13b).

At lower temperatures defects tend to form associated species ( $D_{\text{Li}}V_{\text{Li}}$ ) according to:



Since mobile lithium vacancies are formed only by dissociation of these associates, the activation energy increases in regime III of Fig. 3a. While all donor dopants increased the conductivity, the ion transport behavior depends on the dopant element. The formation of associates was observed for both undoped and Mg<sup>2+</sup>-doped Li(SCN) (Fig. 3b), while Zn<sup>2+</sup>- and Co<sup>2+</sup>-doping showed a distinctively different behavior (intrinsic to extrinsic transition at  $180^\circ\text{C}$ , extrinsic to association transition at  $88^\circ\text{C}$ ). In the case of Zn<sup>2+</sup>, the measured conductivities do not show any significant changes in slope, which suggests that no associated species form and merely the extrinsic regime is observed. In contrast, the incorporation of Co<sup>2+</sup> in Li(SCN) does not only increase the concentration of  $V'_{\text{Li}}$ , but surprisingly also leads to electronic conductivity. Galvanostatic DC measurements (ESI† Fig. S11a) revealed a dominant electronic conduction below  $84^\circ\text{C}$ . Except for 1.5 mol% Mg<sup>2+</sup>-doped Li(SCN), the intrinsic regime was not observed for any other doped sample, as the transition to intrinsic is too close to the melting point (a small intrinsic regime would be expected for a Schottky defective material, since both the cation and anion lattice become increasingly disordered before finally the material melts).

The observed differences in transport behavior with specific dopants can be understood from the crystal structures and coordination chemistry of the respective thiocyanates (Fig. 4). Li<sup>+</sup> in Li(SCN) is coordinated to six (SCN)<sup>-</sup> anions, forming

octahedra with three Li–S and three Li–N bonds.<sup>31</sup> The Li–S bonds are rather weak, given the strong mismatch in polarizability according to the Pearson HSAB concept.<sup>50</sup> This results in a high tendency of Li(SCN) to form coordination compounds with oxygen-containing ligands, e.g. H<sub>2</sub>O or THF,<sup>51,52</sup> to replace Li–S with Li–O bonds. An even more extreme situation is observed for Mg<sup>2+</sup>, for which thiocyanate hydrates rather decompose upon drying than form Mg–S bonds.<sup>53</sup> The strong tendency of Mg<sup>2+</sup> in magnesium thiocyanates to coordinate with oxygen containing molecules (compared to e.g. Zn<sup>2+</sup> and Co<sup>2+</sup>)<sup>54</sup> suggests that this cation has a strong preference for electrostatic interactions with negatively charged species. This suggests that Mg<sub>Li</sub><sup>+</sup> defects can act as effective trapping sites for  $V'_{\text{Li}}$  forming associates.

This concept also explains the higher conductivity of the new Mg<sub>1.02</sub>Li<sub>3.96</sub>(SCN)<sub>6</sub> phase (Fig. 1a). Since Mg<sup>2+</sup> has a greater disfavor for Mg–S bonds than Li<sup>+</sup>, the (SCN)<sup>-</sup> anions largely coordinate *via* their nitrogen atom to magnesium, meaning that Li<sup>+</sup> has to bind to sulfur (ESI† Fig. S3 and S4). This results

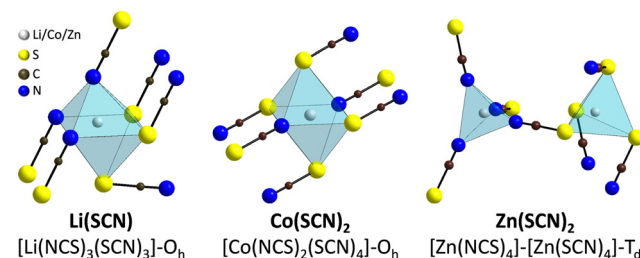


Fig. 4 Comparison of the metal cation coordination polyhedra in Li(SCN),<sup>31</sup> Co(SCN)<sub>2</sub>,<sup>55</sup> and Zn(SCN)<sub>2</sub>.<sup>56</sup> The shown structure of Zn(SCN)<sub>2</sub> is the  $\beta$ -modification (for more details, cf. ESI†).



in a more facile formation of  $V'_{Li}$ , increasing the concentration of mobile defects and lifting the ionic conductivity by more than four orders of magnitude. In contrast, both  $Zn(SCN)_2$  and  $Co(SCN)_2$  form stable M-S bonds in their anhydrous form and are not hygroscopic or prone to form coordination compounds with oxygen containing ligands.<sup>55–57</sup> Thus it is expected that both  $Zn^{\bullet}_{Li}$  and  $Co^{\bullet}_{Li}$  have a far lower tendency to form associates with  $V'_{Li}$  than  $Mg^{\bullet}_{Li}$ . In  $Zn^{2+}$ -doped  $Li(SCN)$  this negligible association is clearly reflected in the conductivity data (Fig. 3), while in  $Co^{2+}$ -doped  $Li(SCN)$  it is overshadowed by an electronic contribution.

Assuming ideally dilute behavior, the mass action law for reaction (2) can be written as:

$$K_S = [V'_{Li}][V^{\bullet}_{SCN}] = N_{Li}N_{SCN} \cdot \exp\left(\frac{\Delta_S S^\circ}{k_B}\right) \exp\left(-\frac{\Delta_S H^\circ}{k_B T}\right) \quad (5)$$

in which  $K_S$  is the Schottky equilibrium constant,  $[V'_{Li}]$  and  $[V^{\bullet}_{SCN}]$  are the respective defect concentrations,  $N_{Li}$  and  $N_{SCN}$  (both equal to  $4/V_{UC}$ ) are the number of available defect sites ( $V_{UC} = 240.56 \text{ \AA}^3$  unit cell volume),<sup>31</sup>  $\Delta_S S^\circ$  and  $\Delta_S H^\circ$  are the standard entropy and enthalpy of defect formation, and  $k_B$  and  $T$  are Boltzmann's constant and temperature. Defect association can be accounted for by reaction (4), and the mass action law reads:

$$K_A = \frac{[(D_{Li}V_{Li})]}{[D^{\bullet}_{Li}][V'_{Li}]} = \frac{N_A}{N_D N_{Li}} \cdot \exp\left(\frac{\Delta_A S^\circ}{k_B}\right) \exp\left(-\frac{\Delta_A H^\circ}{k_B T}\right) \quad (6)$$

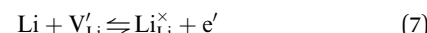
$K_A$  is the association equilibrium constant,  $[(D_{Li}V_{Li})]$ ,  $[D^{\bullet}_{Li}]$  and  $[V'_{Li}]$  are associate, dopant and vacancy concentration, and  $\Delta_A S^\circ$  and  $\Delta_A H^\circ$  are the standard entropy and enthalpy of association. Since the doping occurs on the  $Li^+$  sites,  $N_D$  is equivalent to  $N_{Li}$ , and  $N_A$  equals  $2N_{Li}$ . The total dopant concentration is  $[D_{total}] = [(D_{Li}V_{Li})] + [D^{\bullet}_{Li}]$ , and the Brouwer approximations in Table 1 can be derived.<sup>10,58</sup>

It can be reasonably assumed that the concentrations of electronic carriers are negligible compared to ionic defects (for  $Co^{2+}$ -doping some additional considerations are discussed below). If the enthalpies and entropies are known, the changes in defect concentration with temperature can be semi-quantitatively drawn as done in Fig. 5a.

The results of the doping experiments were used to construct the Brouwer diagrams displayed in Fig. 5b. The defect

chemical analysis of  $Li(SCN)$  shows that changes in ion transport induced by doping are specific to the coordination chemistry of the dopant;  $Mg^{\bullet}_{Li}$  forms associates,  $Zn^{\bullet}_{Li}$  does not and  $Co^{\bullet}_{Li}$  can even induce predominant electronic conduction. This phenomenon of dopant specific transport behavior is most likely connected with the specific coordination chemistry of hard-cation – soft-anion ion conductors.

The electronic conductivity for  $Co^{2+}$ -doped  $Li(SCN)$  cannot unambiguously be interpreted. As long as  $[V'_{Li}]$  is fixed *via* the electroneutrality condition by the donor dopant concentration, and also the elemental lithium activity remains constant, the reaction:



with

$$K_{Li} = \frac{[e']}{a_{Li} [V'_{Li}]} \quad (8)$$

( $K_{Li}$  being the lithium incorporation equilibrium constant and  $a_{Li}$  the activity of elemental lithium) keeps the electronic defect concentration fixed. To obtain an increased electronic conductivity with  $Co^{2+}$ -doping, either further defect chemical complications (e.g. trapping reactions) or strongly modified electronic carrier mobilities would be required. Another possibility is related to the fact that the lithium activity is not strictly fixed in the measurement setup. If the presence of redox-active  $Co^{2+}$  in the synthesis leads to a minor concentration of electronic holes  $h^{\bullet}$  (from traces of oxygen, remaining in the sample also under measurement conditions) with a high mobility, this might create an electronic conductivity that becomes perceptible relative to the low ionic conductivity at low temperature. This would also correspond to a decreased elemental lithium activity.

### 2.3. Determination of defect mobility and defect concentrations

The mobility  $u_{V'_{Li}}$  of  $V'_{Li}$  can be calculated from  $\sigma_{V'_{Li}}$  and  $[V'_{Li}]$  in the extrinsic regime:

$$u_{V'_{Li}} T = \frac{\sigma_{V'_{Li}} T}{z_{V'_{Li}} e \cdot [V'_{Li}]} \quad (9)$$

Table 1 Brouwer approximations of defect concentrations for all employed dopants in  $Li(SCN)$

	Mg	Zn	Co
Regime I intrinsic	$[V'_{Li}] = \sqrt{K_S} = [V^{\bullet}_{SCN}]$ $[Mg^{\bullet}_{Li}] = [D_{total}] > [(Mg_{Li}V_{Li})]$ $[(Mg_{Li}V_{Li})] = K_A \sqrt{K_S} [D_{total}]$	$[V'_{Li}] = \sqrt{K_S} = [V^{\bullet}_{SCN}]$ $[Zn^{\bullet}_{Li}] = [D_{total}]$	$[V'_{Li}] = \sqrt{K_S} = [V^{\bullet}_{SCN}]$ $[Co^{\bullet}_{Li}] = [D_{total}]$
Regime II extrinsic	$[V'_{Li}] = [D_{total}] = [Mg^{\bullet}_{Li}]$ $[V^{\bullet}_{SCN}] = K_S / [D_{total}]$ $[(Mg_{Li}V_{Li})] = K_A [D_{total}]^2$	$[V'_{Li}] = [D_{total}] = [Zn^{\bullet}_{Li}]$ $[V^{\bullet}_{SCN}] = K_S / [D_{total}]$	$[V'_{Li}] = [D_{total}] = [Co^{\bullet}_{Li}]$ $[V^{\bullet}_{SCN}] = K_S / [D_{total}]$
Regime III association	$[V'_{Li}] = \sqrt{[D_{total}] / K_A} = [Mg^{\bullet}_{Li}]$ $[V^{\bullet}_{SCN}] = K_S \sqrt{K_A / [D_{total}]}$ $[(Mg_{Li}V_{Li})] = [D_{total}]$		

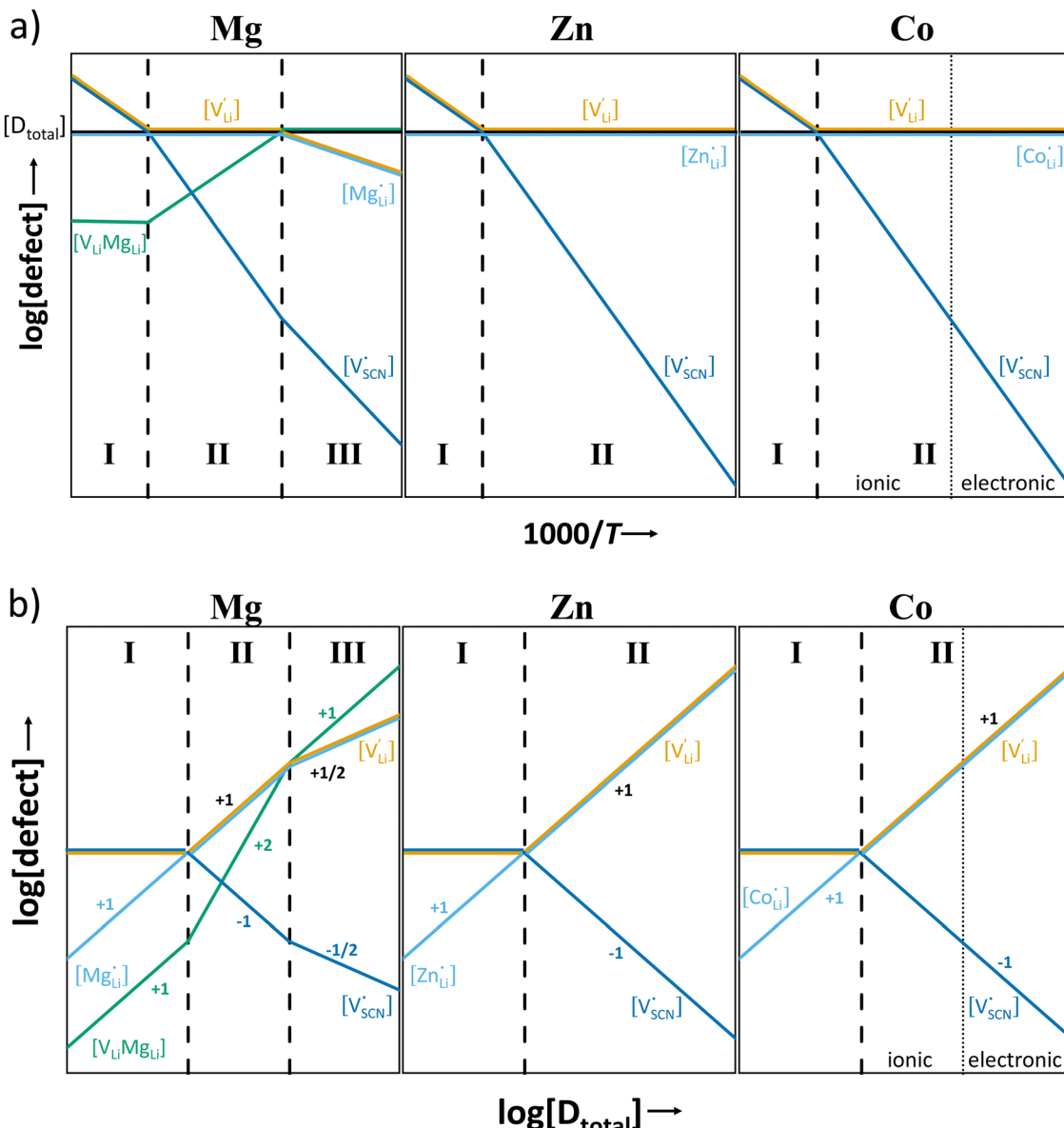


Fig. 5 Defect concentrations of ionic defects in Li(SCN)-doped systems; (a) as a function of temperature, and (b) as a function of dopant concentration (Brouwer diagrams). The dotted line denotes the transition, where for  $\text{Co}^{2+}$ -doping the  $\text{V}'_{\text{Li}}$  ionic conductivity surpasses the one of electronic charge carriers.

where  $z_{\text{V}'_{\text{Li}}}$  and  $e$  are the charge of  $\text{V}'_{\text{Li}}$  (equal to 1) and of an electron.

According to:

$$u_{\text{V}'_{\text{Li}}} T = \frac{r_{\text{Li}}^2 e}{N k_B} \nu_0 \cdot \exp\left(\frac{\Delta_m S_{\text{V}'_{\text{Li}}}}{k_B}\right) \exp\left(-\frac{\Delta_m H_{\text{V}'_{\text{Li}}}}{k_B T}\right) \quad (10)$$

the linear fits shown in Fig. 6a were averaged to yield:

$$u_{\text{V}'_{\text{Li}}} T = 10^{(5.1 \pm 0.4)} \exp\left(-\frac{(0.89 \pm 0.08) \text{ eV}}{k_B T}\right) \text{ cm}^2 \text{ K} \text{ V}^{-1} \text{ s}^{-1} \quad (11)$$

$r_{\text{Li}}$  is the distance to a neighboring available site of  $\text{V}'_{\text{Li}}$  (3.16 Å),  $N$  is the number of neighboring sites (equal to 2), and  $\nu_0$  is the jump attempt frequency ( $\sim 10^{13}$  Hz).<sup>10,58</sup> Knowing the mobility, the Schottky mass action constant  $K_S$  is calculated

from the linear fit of regime I in undoped Li(SCN) (Fig. 3a) by inserting  $[\text{V}'_{\text{Li}}] = \sqrt{K_S}$  and eqn (11) into eqn (9) with the result:

$$K_S = 10^{(45 \pm 2)} \exp\left(-\frac{(0.6 \pm 0.3) \text{ eV}}{k_B T}\right) \text{ cm}^{-6} \quad (12)$$

The association equilibrium constant  $K_A$  is calculated by linearly fitting the conductivity data in regime III of  $\text{Mg}^{2+}$ -doped Li(SCN) (Fig. 3b) and inserting that expression together with  $[\text{V}'_{\text{Li}}] = \sqrt{[\text{D}_{\text{total}}]/K_A}$  into eqn (9), which yields:

$$K_A = 10^{(-25 \pm 3)} \exp\left(-\frac{(-0.3 \pm 0.2) \text{ eV}}{k_B T}\right) \text{ cm}^3 \quad (13)$$



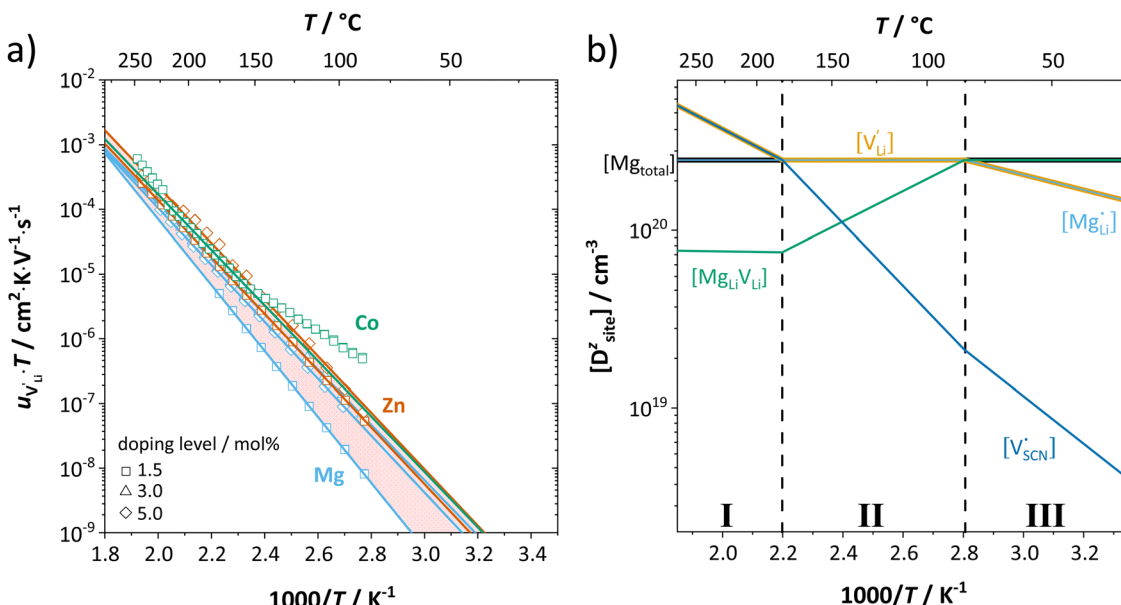


Fig. 6 (a) Calculated mobilities of lithium vacancies for all dopants. Solid lines correspond to linear fits. In case of Co<sup>2+</sup>-doping, only data above 153 °C were used for the fitting. (b) Calculated defect concentrations in 1.5 mol% Mg<sup>2+</sup>-doped Li(SCN).

The corresponding entropies are obtained from eqn (5), (6), (10) and the enthalpies in eqn (11)–(13). The results are summarized in Table 2.

#### 2.4. DFT calculations

To complement the experimental results, DFT calculations were performed for the possible defect formation enthalpies in Li(SCN) using the PBE exchange–correlation functional<sup>59</sup> which is broadly applied for calculations of ionic solids. For a fully relaxed cell, lattice parameters of 12.988 Å, 3.682 Å, and 5.311 Å were obtained. The cell volume is noticeably larger compared to the experimental value (+5.5%). In particular the lattice parameter in *a* direction (between the “layers” of Li(SCN) double chains, *cf.* Fig. S2, ESI<sup>†</sup>) is overestimated by 6.9%. This suggests that in this direction van der Waals forces make an important contribution, which tend to be underestimated in standard functionals.

The formation energies of defect pairs in undoped Li(SCN) were calculated, and found to be lower for Schottky pairs (V<sub>Li</sub>' + V<sub>SCN</sub>', 0.34 eV) than Frenkel pairs (V<sub>Li</sub>' + Li<sub>*i*</sub>', 1.1 eV).

Table 2 Experimental thermodynamic and kinetic data of Li(SCN) to calculate defect concentrations (calculated defect concentrations of 1.5 mol% Mg<sup>2+</sup>-doped Li(SCN) can be found in Fig. 6b)

Schottky disorder	$\Delta_s H^\circ$ (eV)	$0.6 \pm 0.3$
V <sub>Li</sub> ' migration	$\Delta_s S^\circ (k_B)$	$5 \pm 2^a$
	$\Delta_m H_{V_{Li}'} (eV)$	$0.89 \pm 0.08$
	$\Delta_m S_{V_{Li}'} (k_B)$	$7.7 \pm 0.9$
(Mg <sub>Li</sub> V <sub>Li</sub> ) association	$\Delta_A H^\circ$ (eV)	$-0.3 \pm 0.2$
	$\Delta_A S^\circ (k_B)$	$-8 \pm 6$

<sup>a</sup> Values from different samples could deviate even beyond the estimated error.

This sequence matches with the results from doping experiments in section 2.3. The numerical value is rather low compared to the experimental result in Table 2 ( $0.6 \pm 0.3$  eV), although within the estimated experimental error. Migration barriers were calculated for V<sub>Li</sub>' and Li<sub>*i*</sub>' (technical details are specified in the ESI<sup>†</sup>). For V<sub>Li</sub>' two possibilities were considered (Fig. S2c and d, ESI<sup>†</sup>): (i) straight along *b* direction, (ii) “zigzag” path *via* the shortest distances between regular Li sites. The “zigzag” path yields the lower barrier of 0.08 eV compared to the direct path (0.26 eV). This relative magnitude appears reasonable, as the “zigzag” path has the shorter individual jump distance. However, the absolute values are significantly lower than the experimental values (Table 2). This is most probably related to the overestimation of the cell volume in the present calculations. The barrier for the Li interstitial in *b* direction amounts to 0.88 eV. Thus, Li interstitials are not only less favorable with respect to defect formation, but also regarding defect migration.

#### 2.5. Comparison to other lithium ion conductors

It is interesting to compare Li(SCN) with other, chemically similar Li<sup>+</sup> cation conductors (Fig. 7). Naively, one might expect the conductivity of Li(SCN) between Li<sub>3</sub>N and Li<sub>2</sub>S, which is evidently not the case. In addition, while the chemical properties and behavior (including structure and reactivity) of Li(SCN) are similar to those of the lithium halides, the situation is more complex concerning ion transport.

Table 3 compares data of relevant Li<sup>+</sup> systems to Li(SCN) data of the present work. The literature entropy values or association energy data (as far as available) are very similar to the present Li(SCN) data, giving confidence in their magnitudes. Despite Li(SCN) and all LiX compounds being Schottky defective,

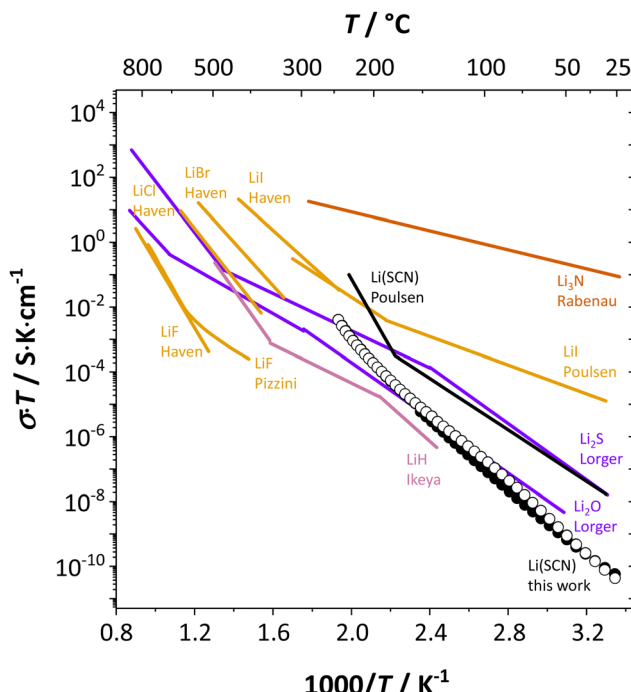


Fig. 7 Comparison of Li(SCN) conductivity (black circles) with literature data (black line)<sup>30</sup> and different Li<sup>+</sup> cation conducting materials; LiH (magenta),<sup>7</sup> Li<sub>3</sub>N (red),<sup>60</sup> chalcogenides (purple),<sup>10,11</sup> and halides (orange).<sup>13,14,23</sup>

Li(SCN) deviates from some trends seen for LiX. In the series H<sup>-</sup>/F<sup>-</sup>, Cl<sup>-</sup>, Br<sup>-</sup> and I<sup>-</sup>, defect formation and migration enthalpies decrease, but defect formation is always energetically more costly than migration. Li(SCN) breaks this relation, with a relatively low formation enthalpy and a high migration enthalpy. As will be shown in more detail in Part II,<sup>40</sup> this deviation is closely related to the specific ion transport mechanism in Li(SCN). The important difference between Li(SCN) and the lithium halides is the bidentate, anisotropic (SCN)<sup>-</sup> anion, since Li<sup>+</sup> can coordinate both to  $-\overline{S} - C \equiv N|$  and  $-\langle N=C=S \rangle$ , with Li-N bonds being more favorable than Li-S bonds. While defect formation is comparatively facile (weak Li-S bonds), ion migration is inhibited by the rigid anion lattice (strong Li-N bonds) and a slow relaxation process (anisotropic shape) after an initial ion jump. This emphasizes the importance of specific chemical interactions, which affect ionic mobilities at least as much as (simplistic) geometrical/size and electrostatic arguments.

### 3. Conclusion

This investigation of anhydrous Li(SCN) shows its specific ion transport properties and defect chemistry. The material forms Schottky pairs, of which lithium vacancies are the mobile defects throughout the investigated temperature range. The solubility limit for Mg<sup>2+</sup>, Zn<sup>2+</sup> and Co<sup>2+</sup> donor doping is estimated to be around 3–5 mol%. Higher Mg<sup>2+</sup>-concentrations led to the formation of a previously unknown phase Mg<sub>1.02</sub>Li<sub>3.96</sub>(SCN)<sub>6</sub>, whose crystal structure was solved from XRPD data. The derived defect model for anhydrous Li(SCN) shows that the formation of defect associates depends on the employed dopant; more specifically on the (dis-)favor to form M-S bonds with the (SCN)<sup>-</sup> anion. The comparison of the defect formation and migration enthalpy of Li(SCN) with similar materials emphasizes the impact of the (SCN)<sup>-</sup> anion on ion transport. Although the highly polarizable anion facilitates defect formation, its very asymmetric interaction of sulfur and nitrogen with Li<sup>+</sup> cations hinders the migration of mobile defects, which renders the material a poor conductor. More details about this impact as well as the defect chemistry close to the melting point will be given in separate publications (Part II<sup>40</sup> and III<sup>41</sup>).

### 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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Table 3 Comparison of thermodynamic and kinetic data from Li(SCN) with LiX (X = H<sup>-</sup>, F<sup>-</sup>, Cl<sup>-</sup>, Br<sup>-</sup>, I<sup>-</sup>)

Compound	$\Delta_S H^\circ$ (eV)	$\Delta_S S^\circ$ (k <sub>B</sub> )	$\Delta_m H_{V'_{Li}}$ (eV)	$\Delta_m S_{V'_{Li}}$ (k <sub>B</sub> )	$\Delta_A H^\circ$ (eV)	$\Delta_A S^\circ$ (k <sub>B</sub> )	Ref.
LiH expt.	2.3 ± 0.3		0.54 ± 0.02		-0.50 ± 0.05 (Mg <sub>Li</sub> V <sub>Li</sub> )		7
LiF expt.	2.6 ± 0.2		0.67 ± 0.02				12, 14, 61, 62
DFT	2.2–2.9		0.6				63
LiCl expt.	2.12		0.41				12, 14
LiBr expt.	1.80		0.39				12, 14
LiI expt.	1.2 ± 0.1	4.5	0.41 ± 0.03	4.9			12, 14, 20
Li(SCN) expt.	0.6 ± 0.3	5 ± 2	0.89 ± 0.08	7.7 ± 0.9	-0.3 ± 0.2 (Mg <sub>Li</sub> V <sub>Li</sub> )	-8 ± 6 (Mg <sub>Li</sub> V <sub>Li</sub> )	This work



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