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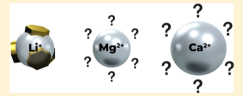
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# Cation Solvation and Physicochemical Properties of Ca Battery **Electrolytes**

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Supporting Information

ABSTRACT: Divalent-cation-based batteries are being considered as potential high energy density storage devices. The optimization of electrolytes for these technologies is, however, still largely lacking. Recent demonstration of the feasibility of Ca and Mg plating and stripping in the presence of a passivation layer or an artificial interphase has paved the way for more diverse electrolyte formulations. Here, we exhaustively evaluate several Ca-based electrolytes with different salts, solvents, and concentrations, via measuring physicochemical



properties and using vibrational spectroscopy. Some comparisons with Mg- and Li-based electrolytes are made to highlight the unique properties of the Ca<sup>2+</sup> cation. The Ca-salt solubility is found to be a major issue, calling for development of new highly dissociative salts. Nonetheless, reasonable salt solubility and dissociation are achieved using bis(trifluoromethanesulfonyl)imide (TFSI), BF<sub>4</sub>, and triflate anion based electrolytes and high-permittivity solvents, such as ethylene carbonate (EC), propylene carbonate (PC), γ-butyrolactone (gBL), and N,N-dimethylformamide (DMF). The local Ca<sup>2+</sup> coordination is concentrationdependent and rather complex, possibly involving bidentate coordination and participation of the nitrogen atom of DMF. The ionicity and the degree of ion-pair formation are both investigated and found to be strongly dependent on the nature of the cation, solvent donicity, and salt concentration. The large ion-ion interaction energies of the contact ion pairs, confirmed by density functional theory (DFT) calculations, are expected to play a major role in the interfacial processes, and thus, we here provide electrolyte design strategies to engineer the cation solvation and possibly improve the power performance of divalent battery systems.

### INTRODUCTION

Achieving sustainable and affordable energy sources and storage is one of the major challenges of our modern society-not the least to attack climate change and oil dependency. Considering only the transportation contribution, more than 1.2 billion vehicles are currently at use all around the world, and less than 0.2% are electric. While Li-based battery technologies undoubtedly will power more and more electric vehicles (EVs) in the next 5-10 years, the controversial debate on sustainability and production/extraction costs of lithium cannot be ignored. 1,2 The development of alternative battery chemistries based on abundant elements is thus urged for to accommodate a significant growth of the EV market. Calcium- and magnesium-based batteries are interesting candidates as they hold promise for safe use of metal anodes, thus potentially presenting high energy densities and low cost. Indeed, Ca and Mg electrodes have theoretical specific capacities of 1340 and 2000 mAh g<sup>-1</sup>, standard redox potentials of -2.76 and -2.38 V vs the standard hydrogen electrode (SHE), and are the fifth and eighth most abundant elements on the Earth's crust, respectively. 3,4

A myriad of Ca-based battery concepts have been developed since the first report of calcium as an electroactive element in

1964. Mostly, thermal batteries, using electrolytes with very high melting points, and the calcium equivalent to the commercial Li thionyl chloride (Li-SOCl<sub>2</sub>) batteries were developed.<sup>6</sup> However, in contrast to Mg, electrodeposition of calcium has for long been elusive and only primary cells were achieved. The lack of calcium plating upon cell reversal was even considered as a safety advantage for the Ca-SOCl2 cells as compared with its Li analogue.

Moving to room-temperature and liquid electrolytes, no Ca equivalents to Mg-based Grignard reagent electrolytes (allowing for Mg plating and stripping) are available. Seminal studies by Aurbach et al. using conventional aprotic organic electrolytes similar to those used for lithium-ion batteries (LIBs) allowed them to conclude that the electrochemical behavior of Ca metal anodes is surface-film controlled, as is the case for Mg. 9,10 The lack of divalent cation permeability of the formed passivation layer was suggested to be the origin of the impossibility to electrodeposit calcium or magnesium. While the electrolyte remains a major bottleneck for any divalent-

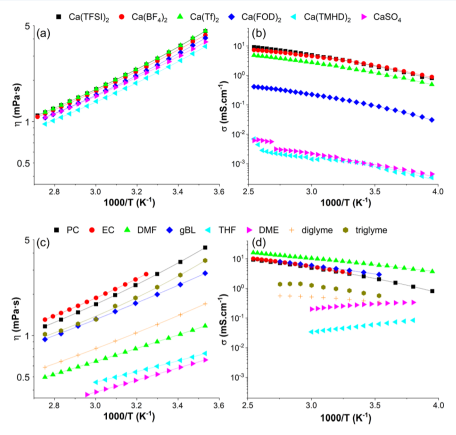
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**Figure 1.** Physical properties of calcium-based electrolytes. Arrhenius plots for (a) viscosity and (b) conductivity of 0.1 M calcium salt in PC. Arrhenius plots for (c) viscosity and (d) conductivity for 0.1 M Ca(TFSI)<sub>2</sub> in several solvents. Solid lines represent the Vogel—Tammann—Fulcher (VTF) fits.

cation-based batteries and most research studies so far have focused on electrolytes for which no passivation layers are formed (mostly based on ether solvents), solvents, recent demonstrations of Ca and Mg plating and stripping in the presence of a passivation layer or an artificial interphase the way for evaluation of new electrolyte formulations by allowing decoupling of interfacial and electrolyte requirements. The present study, thus, focuses on important parameters in the design of new electrolytes, such as cation mobility and desolvation energies, broadening the choices of electrolyte component candidate salts and solvents, potentially leading to formation of better-performing passivation layers.

As compared to Li<sup>+</sup> and Na<sup>+</sup> conducting electrolytes, divalent-cation-based electrolytes have a stronger tendency of ion-pair formation16 and larger cation desolvation energies.1 In addition, as the Li<sup>+</sup> and Na<sup>+</sup> mobilities are significantly lower than those of the anions and the desolvation energy plays a major role in the kinetics of Li+ insertion into graphite, 18-21 it can be expected that the power performance of any divalent-cation-based batteries will also depend greatly on the nature of the cation solvation shell. Here, we systematically investigate calcium-based electrolytes and report on the relationship between the cation solvation, coordination number and ion pairing, and physicochemical properties, such as ion conductivity and viscosity for different salts and concentrations and organic solvents. Some parallels are drawn with Li and Mg systems allowing for a better understanding of the strategies to optimize divalent cation mobility and desolvation energy.

## **■ EXPERIMENTAL SECTION**

**Materials.** The solvents used were propylene carbonate (PC, Aldrich, anhydrous, 99.7%), ethylene carbonate (EC, Aldrich, anhydrous, 99.0%), N,N-dimethylformamide (DMF, Alfa Aesar, 99.7%),  $\gamma$ -butyrolactone (gBL, Aldrich, anhydrous, 99.0%), tetrahydrofuran (THF, Aldrich, anhydrous, 99.9%), 1,2-dimethoxyethane (DME, Aldrich, anhydrous, 99.5%), diethyleneglycol dimethyl ether (diglyme, Aldrich, 99.0%), triethyleneglycol dimethyl ether (triglyme, Aldrich, 99.0%), and fluoroethylene carbonate (FEC, Aldrich, 99%).

The commercially available calcium salts used were calcium bis(trifluoromethanesulfonyl)imide ( $Ca(TFSI)_2$ , Solvionic, 99.5%), calcium tetrafluoroborate ( $Ca(BF_4)_2$ , Alfa Aesar, hydrated), calcium trifluoromethanesulfonate ( $Ca(Tf)_2$ , Sigma-Aldrich, 99.9%), calcium sulfate ( $CaSO_4$ , Sigma-Aldrich, 99.99%), calcium bis(6,6,7,7,8,8,8-heptafluoro-2,2-dimethyl-3,5-octanedionate) ( $Ca(FOD)_2$ , Sigma-Aldrich, >99.9%), and calcium bis(2,2,6,6-tetramethyl-3,5-heptanedionate) ( $Ca(TMHD)_2$ , Sigma-Aldrich). LiTFSI (Solvionic, 99.9%) and Mg(TFSI)<sub>2</sub> (Solvionic, 99.5%) were also used to create reference electrolyte systems. All salts and solvents were kept and used under a dry atmosphere (<1 ppm  $O_2$  and  $O_2$  and  $O_2$  in an argon-filled Jacomex glovebox at 25 °C.

**Electrolytes.** First, two families of Ca-based electrolytes were made by dissolving (i) six different Ca-salts in PC at a concentration of 0.1 M and (ii) 0.1 M Ca(TFSI)<sub>2</sub> in eight different solvents. Subsequently, a third family was evaluated for the best solvent and 0.1–1.5 M Ca(TFSI)<sub>2</sub>, and this was also further compared with LiTFSI- and Mg(TFSI)<sub>2</sub>-based electrolytes. All electrolytes were prepared under a dry

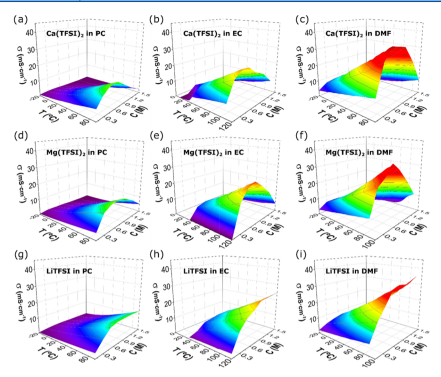


Figure 2. Ion conductivity plots for (a-c)  $Ca(TFSI)_2$ , (d-f)  $Mg(TFSI)_2$ , and (g-i) LiTFSI in (a, d, g) PC, (b, e, h) EC, and (c, f, i) DMF, at different concentrations and temperatures.

atmosphere in the glovebox and remained there until analysis. Molecular sieves (Alfa Aesar, 3 Å) were used as drying agents for each electrolyte. Prior to any characterization, the water contents were measured using an 899 Karl Fisher Coulometer (Metrohm), resulting in <30 ppm for all electrolytes.

**Experimental Methods.** Ion conductivity measurements were performed using a FRA-based Multiplexed Conductivity Meter MCM 10 (BioLogic Science Instruments) from -20 to +120 °C (10 °C steps with 20 min equilibration time).

Viscosity and density measurements were carried out from +10 to +90 °C using a rolling-ball viscometer (Lovis 2000 M/ME, Anton Parr). The cell temperature was regulated within  $\pm 0.02$  °C. Ultrapure water was used to calibrate the viscometer and densitometer. The uncertainties of the density and viscosity measurements were less than  $5 \times 10^{-5}$  g cm<sup>-3</sup> and 0.5%, respectively.

Fourier transform infrared (FT-IR) spectroscopy was carried out using a Jasco FT/IR-4700 Spectrometer with the universal ATR Pro One accessory (equipped with a diamond crystal) for the range 500–4000 cm<sup>-1</sup> at a resolution of 2 cm<sup>-1</sup>.

Raman spectra were recorded using a Bruker MultiRAM FT-Raman spectrometer with a spectral resolution of 2 cm<sup>-1</sup> using a Nd:YAG laser (1064 nm, 500 mW) as the excitation source. Data were typically averaged over 500 scans and recorded at 40 °C for the carbonate-based electrolytes and 25 °C for the DMF-based electrolytes. All bands in the resulting spectra were fitted as Voigt functions, typically with a unity Gaussian/Lorentzian mix by width, using Origin Software.

**Computational Details.** The Raman activities of the complexes  $[Ca(DMF)_6]^{2+}$ ,  $[Ca(PC)_6]^{2+}$ , and  $[Ca(FEC)_6]^{2+}$  were obtained by first creating the initial structures by employing the minima hopping global optimization algorithm (MH) as implemented in the Atomistic Simulation Environment (ASE) coupled with TURBOMOLE. 22–24 This method carries out short ab initio molecular dynamics (AIMD) and

local optimizations to explore the energy landscape where the temperature is used to escape any local minima. The AIMD runs were performed in the microcanonical ensemble with a time step of 1 fs. The BP86 functional was employed together with a SVP basis set as implemented in TURBOMOLE. 25,26 The MH algorithm started at 300 K and an initial energy threshold of 48 kJ mol<sup>-1</sup>. The subsequent calculations of gasphase geometries and Raman activities for the most stable structures used the M06-2X functional and the 6-31G\*\* basis set as implemented in the Jaguar package. 27-31 All obtained structures appear to be minima, with no imaginary vibrational frequencies.

## ■ RESULTS AND DISCUSSION

Viscosity and Ion Conductivity. To separate out the relative influence of the solvent and the nature and concentration of the salt, two different families of electrolytes were made. First, the six different Ca-salts were dissolved in PC aiming at a concentration of 0.1 M to target salt solubility i.e., can we at all make electrolytes from these salts? Notably, 0.1 M solutions in PC were easily prepared for the majority of salts under study, but CaSO<sub>4</sub> and Ca(TMHD)<sub>2</sub> presented poor solubilities, and therefore, their saturated electrolytes were used. While the viscosities of all electrolytes are relatively similar, their ionic conductivities differ greatly for the different salts under study (Figure 1a,b). The Ca(TMHD)2- and CaSO<sub>4</sub>-based electrolytes show the lowest ion conductivities, mainly due to the poor salt solubility, while Ca(TFSI)<sub>2</sub>, Ca(BF<sub>4</sub>)<sub>2</sub>, and Ca(Tf)<sub>2</sub> all perform better with their ion conductivities being 3 orders of magnitude higher (Figure 1b). This is consistent with the ease of ion-ion dissociation, facilitating solubility and creation of charge carriers. These results highlight one of the major issues in the development of new divalent-cation-based electrolytes: the salt solubility and the need for new highly dissociating salts.

Turning to the role of the electrolyte solvent, 0.1 M  $Ca(TFSI)_2$  electrolytes were studied (Figure 1c,d). It appears that the ion conductivity and the ability to dissolve Ca-salt both follow the solvent donor number (Table S1); highest for DMF and lowest for PC. This trend agrees well with the computed solubility of calcium salts in a wide variety of solvents.<sup>32</sup>

Tentatively, salt solubility, and not viscosity, seems to be the determining factor for ion conductivity as we find, for example, one of the highest ion conductivities for the EC-based electrolyte (cf. Figure 1d) while being the most viscous electrolyte (Figure 1c). The THF- and glyme-based electrolytes show decreased ion conductivities as a function of temperature, an unusual behavior attributed to the presence and augmentation of strong ion pairing, as previously reported for Mg-glyme systems.<sup>33</sup>

Overall, the DMF-based electrolytes preliminarily seem to offer the best performance, and together with more standard PC- and EC-based electrolytes, maximum ionic conductivities are obtained for salt concentrations of 0.42-0.57 M (Figure 2), after which a steep decrease is recorded, especially at lower temperatures. The maxima correspond to the lowest activation energies ( $E_a$ ) in the VTF fits (Figure S6 and Table S3). Notably, these maxima are found at much lower salt concentrations than those for standard Li<sup>+</sup> and Na<sup>+</sup> conducting electrolytes, with maxima typically at 1.0-1.2 M. Therefore, as far as macroscopic properties (viscosity and ionic conductivity) are concerned, moderate salt concentrations (around 0.5 M), highly dissociating anions (such as TFSI or BF<sub>4</sub><sup>-</sup>), and solvents with high donor numbers (DMF, EC, or PC in that order) are required for Ca-battery electrolytes.

lonicity. While viscosity, in general, limits ion motion in electrolytes, intermolecular and electrostatic interactions are also important as they affect both the solvation, i.e., nature and radius of the diffusing (cationic) species, and ion pairing, i.e., concentration of charge carriers. All of these interactions are taken into account in the ionicity of the electrolyte within the framework of the empirical Walden's rule.<sup>34</sup> While originally formulated for infinitely diluted aqueous electrolytes with solvated ions of the same radius, it can be used to compare nonideal, nonaqueous electrolytes. In a Walden plot, the logarithm of the molar conductivity,  $\Lambda$ , is plotted against the fluidity, i.e., the inverse of the viscosity,  $1/\eta$ , and most often an ideal (infinitely diluted) aqueous KCl solution is chosen as a reference system. 35,36 Deviations below the ideal KCl line indicate that other actions than viscosity hinder the ionic motion, e.g., ion pairing.

The Walden plots for the Ca, Mg, and Li electrolytes show all data to fall below the ideal KCl line, and hence they are subject to ion pairing. The anion influence is clear as the ionicity (Figure 3a) and the ion conductivity (Figure 1b) covary:  $Ca(TFSI)_2 > Ca(BF_4)_2 > Ca(Tf)_2 > Ca(FOD)_2 > CaSO_4 > Ca(TMHD)_2$ , in clear agreement with how delocalized the negative charge is for the three former anions and the poor salt solubility and ion—ion dissociation for the three latter salts.

For the solvents, the ionicity of the electrolytes increases with increasing dielectric constant (Figure 3b and Table S1): EC > PC > gBL > DMF > triglyme > diglyme > DME > THF. For THF and the glymes (incl. DME), the unusual behavior, illustrated by the negative slope in Figure 3b, can be attributed to their low dielectric constants and the formation of ionic aggregates.<sup>37</sup> In contrast, the EC, PC, gBL, and DMF

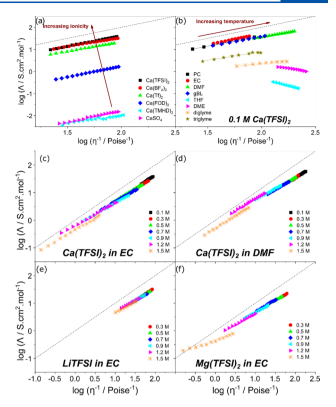


Figure 3. Walden plots of (a) different Ca-salts (0.1 M) in PC, (b) 0.1 M  $Ca(TFSI)_2$  in different solvents and 0.1 to 1.5 M (c)  $Ca(TFSI)_2$  in EC, (d)  $Ca(TFSI)_2$  in DMF, (e) LiTFSI in EC, and (f) Mg(TFSI)<sub>2</sub> in EC. The dashed line represents an aqueous KCl solution as a reference for an ideal, fully dissociated electrolyte.

electrolytes have data in close vicinity to the ideal KCl standard line, implying a high ion—ion dissociation degree. The Walden plot slopes are 1.01, 0.95, 0.90, and 0.87 for EC, PC, gBL, and DMF, respectively, which shows that while the DMF electrolytes have the highest ion conductivities and lowest viscosities, they do seem slightly less dissociated than the gBL-, EC-, and even PC-based electrolytes.

Increasing the salt concentration results in very slight shifts toward the ideal KCl line and decreased slopes for both EC and DMF electrolytes (Figure 3c-f). This is clearly observed for Ca(TFSI)<sub>2</sub> and Mg(TFSI)<sub>2</sub> but is much subtler for the LiTFSI electrolytes. It can be understood in terms of ionicity as increasing salt concentration for divalent cation electrolytes promotes extensive ion—ion interactions.

In contrast to the macroscopic view on electrolyte performance presented in the previous section, ionicity is a more local criterion, and an electrolyte with a high dielectric constant shows a better performance: EC > PC > gBL > DMF. Therefore, there is a clear discrepancy between the micro- and macroscopic points of view. To understand this discrepancy better, the molecular interactions and speciation were further investigated by vibrational spectroscopy.

**Cation–Anion and Cation–Solvent Interactions.** As the cation–solvent interactions compete with the cation–anion interactions, we turn to vibrational spectroscopy to assess the local coordination in more detail. Starting with infrared spectroscopy, the very intense IR absorption band of the carbonyl C=O stretching mode should be an interesting probe for EC, PC, and DMF as solvents. However, for cyclic carbonates, multiple mechanisms prevent any straightforward analysis. The fundamental stretching frequency of C=O ( $\nu_2$ )

at 1793 cm<sup>-1</sup> couples strongly with the first overtone of the skeletal breathing of the ring  $(2\nu_7)$  and produces an intense band at 1767 cm<sup>-1</sup>; a Fermi resonance phenomenon (Figure 4).<sup>38</sup> Upon coordination with a cation, these bands can, thus,

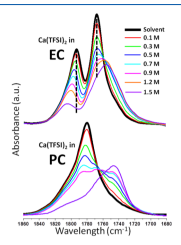


Figure 4. Carbonyl stretching region IR spectra of Ca(TFSI)<sub>2</sub> in ECand PC-based electrolytes as a function of salt concentrations.

shift due to different mechanisms: losing the Fermi coupling condition would return the mode to its "unperturbed" frequency (gas-phase IR<sup>39</sup> ca. 1870 cm<sup>-1</sup>, upward shift), decreased electron density of the C=O bond (downward shift), and/or reduced intensity, and possibly a shift, of the  $2\nu_7$  overtone band due to the lack of Fermi resonance (upward shift). This complex combination of mechanisms prevents any unambiguous use of the carbonyl region.

Instead, the out-of-phase wagging of the CH<sub>2</sub> groups ( $\nu_{15}$ , 1390 cm<sup>-1</sup>) and the fundamental skeletal breathing of the ring ( $\nu_{7}$ , 892 cm<sup>-1</sup>) are used, and they both shift to higher wavenumbers as the Ca-salt concentration increases (Figure S7a). At 1.5 M, these "free" solvent bands are almost gone, suggesting that almost all ECs are coordinated by the Ca<sup>2+</sup> cations. As compared to Li-salt-based electrolytes, where free EC is found even at 3 M,<sup>35</sup> the Ca<sup>2+</sup> cations are indeed expected to have larger first solvation shells.

The carbonyl region of the IR spectra of PC-based electrolytes is laden with the same problems as those for EC, but with the additional disadvantage that the lower symmetry of the PC produces, in general, broader bands.<sup>36</sup> For the PC electrolytes, we indeed discern that all of the same modes are affected by Ca<sup>2+</sup> coordination, but the shifts are not sufficiently large to allow us to correctly determine the contributions from free and coordinated PC, respectively.

In the IR spectra of DMF electrolytes (Figure 5), the C=O stretching mode gets clearly shifted toward lower wavenumbers upon coordination by Li<sup>+</sup>, Mg<sup>2+</sup>, or Ca<sup>2+</sup>. However, the  $\delta$  O=C-N vibration at 658 cm<sup>-1</sup> is a better option to analyze the cation–DMF interaction, with a significantly larger shift obtained for the Mg<sup>2+</sup>-containing electrolytes, attributed to the more polarizing character of the magnesium cation, resulting in a tighter first solvation shell. Significant differences in terms of solvation shell dynamics and desolvation energies are, thus, expected between the Mg-, Ca-, and Li-based electrolytes.

Ohashi et al. recently reported that formamide coordinates Ca<sup>2+</sup> cations only through the oxygen atom, <sup>40</sup> but the nitrogen of the amide group might also take part in the cation

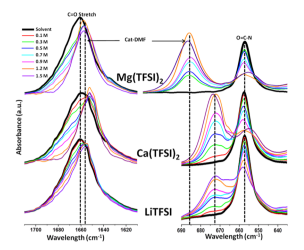


Figure 5. IR spectra of DMF-based electrolytes as a function of  $Mg(TFSI)_{2}$ ,  $Ca(TFSI)_{2}$ , and LiTFSI salt concentrations.

coordination. Al, As compared to formamide, the nitrogen atom in DMF has a higher electronic density due to the electron donor character of the methyl groups—further supported by computational studies showing that its electron lone-pair occupies a  $\pi$  nonbonding molecular orbital, which also involves the methyl groups. The distinct 20 cm<sup>-1</sup> shift of the symmetric bending of the  $-CH_3$  groups upon coordination by  $Ca^{2+}$  or  $Mg^{2+}$  points to a direct participation of the nitrogen atom in the cation solvation (Figure S7b).

For the cation solvation number determination, we turn to Raman spectroscopy and a few relatively intense bands for each solvent, which are minimally affected by other vibration modes but highly sensitive to cation coordination (Figure 6): ring breathing mode for EC (at 894 cm<sup>-1</sup>),  $\gamma$ C=O for PC (sum of two bands at 706 and 712 cm<sup>-1</sup>), and  $\delta$ O=C-N for DMF (at 658 cm<sup>-1</sup>). At this point, we also include fluoroethylene carbonate (FEC) as a model solvent with a very low donor number, and the band analyzed is the ring breathing mode (at 905 cm<sup>-1</sup>).

For all regions, the free solvent band shifts toward higher wavenumbers upon cation coordination, and by deconvolution of the spectra (Figure S8), the solvation numbers (SNs) of the cations were calculated using eq 1

$$SN = \left(\frac{A_{\text{bound}}}{A_{\text{bound}} + \theta A_{\text{free}}}\right) \left(\frac{C_{\text{s}}}{C_{\text{Ca}^{2+}}}\right)$$
(1)

where  $C_{\rm S}$  and  $C_{\rm Ca}^{^{2+}}$  represent the molar concentrations of the solvent and Ca-salt, respectively. The parameter,  $\theta$ , is the ratio of the Raman activity of free and coordinated solvent molecules as obtained by density functional theory (DFT) calculations ( $\theta = RA_{\rm coord}/RA_{\rm free}$ ).

Similarly, the number of TFSI anions coordinating each cation, as a measure of the contact ion-pair concentration, was calculated using eq 2

TFSI coordinated by 
$$M^{2+} = \left(\frac{2A_{\text{coord.}}}{A_{\text{coord.}} + A_{\text{free}}}\right)$$
 (2)

This is derived from eq 1 and considers the 2:1 TFSI/Ca<sup>2+</sup> relation and, furthermore, assumes that the TFSI all breathing mode (free at 740 cm<sup>-1</sup>) is equally Raman-active when "coordinated" ( $\theta = 1$ , confirmed for Li-based electrolytes<sup>44</sup>).

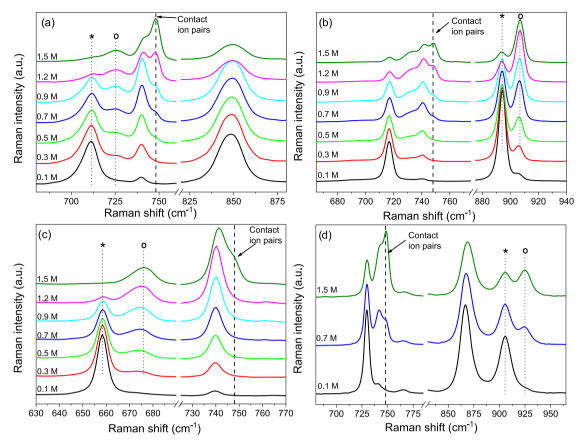


Figure 6. Raman spectra of  $Ca(TFSI)_2$  in (a) PC, (b) EC, (c) DMF, and (d) FEC, at different salt concentrations. \* denotes free solvent, while of denotes  $Ca^{2+}$ -coordinated solvent. The band assigned to cation—anion contact ion pairs appears in all cases at 748 cm<sup>-1</sup> and that of free TFSI at 740 cm<sup>-1</sup>.

The number of solvent molecules in the first solvation shell  $(SN_{solvent})$  decreases upon increasing the salt concentration (Figure 7a), which can be explained by the displacement of solvent molecules by anions, thus forming contact ion pairs. The concentration threshold, when this process is triggered, is

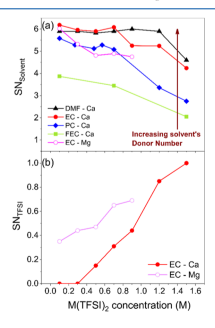


Figure 7. SN of (a) solvent and (b) TFSI as a function of  $M(TFSI)_2$  salt concentration (M = Ca and Mg).

highly dependent on the electrolyte solvent (Figure 7a), where DMF is most resilient to be displaced, followed by EC, PC, and finally FEC—which correlates with their donor numbers (Table S1).

The formation of contact ion pairs can also be clearly observed in the Raman spectra of the electrolytes by the appearance of a band at 748 cm<sup>-1</sup> attributed to Ca<sup>2+</sup>-TFSI complexes (Figure 6). Indeed, DMF is proven to be a very good solvent as no contact ion pairs were detected below 1.2 M salt concentration, while for carbonate solvents (EC and PC) they appeared already at 0.5 M.

The formation of contact ion pairs also heavily depends on the salt employed; as TFSI is flexible and its charge is highly delocalized, it is less prone to contact ion-pair formation and is, thus, preferred over other anions (such as  $BF_4^-$ ), which are involved in the solvation shell at lower concentrations (Figure S9). For Mg electrolytes, however, the polarizing character of  $Mg^{2+}$  promotes the formation of contact ion pairs even for 0.1 M  $Mg(TFSI)_2$  in EC (Figure 7b).

While the formation of contact ion pairs is moderated by the solvent's donor numbers, we propose that its permittivity (dielectric constant) plays a major role in the formation of solvent-separated ion pairs. In the Raman spectra of DMF electrolytes (a low-polarity solvent), the band of free TFSI (740 cm<sup>-1</sup>) grows more asymmetric in the range from 0.3–1.2 M. This asymmetry evidences a certain cation—anion interaction even if there is no direct contact. The presence of such solvent-separated ion pairs could, at least partially, explain why DMF-based electrolytes do not provide the

highest ionicities (Figure 4b). On the other hand, the high permittivities of EC and PC effectively shield the cation charge, decreasing the tendency to form solvent-separated ion pairs, and hence, result in higher ionicities. Yet, as the donor numbers of EC and PC are lower, they are more easily displaced by the TFSI anions as the salt concentration increases, forming contact ion pairs and possibly higher aggregates.

We also observe that the total SN (solvent + anion) decreases as a function of salt concentration, and hence, each anion entering displaces more than one solvent molecule. If the coordination sites of the cation are to be constant, bidentate coordination can be expected by either TFSI or the solvent. For cyclic carbonates, bidentate coordination to Li<sup>+</sup> and Na<sup>+</sup> has been proposed for highly concentrated electrolytes, <sup>45</sup> and as the band for the coordinated solvent is actually composed of two contributions shifted by ca. 5 cm<sup>-1</sup> (Figure S10), bidentate coordination could also be the case for Ca<sup>2+</sup>.

The evolution of the amount of free solvent molecules with salt concentration also appears to be solvent-dependent. For the PC- and DMF-based electrolytes almost no free solvent band can be detected (Figure 6a,c) for salt concentrations above 1.2 M, while they can still be detected for higher salt concentrations for both EC and FEC, most likely due to the bidentate coordination of the solvent in the first case and to the high degree of ion-pair formation in the latter. For Mg(TFSI)<sub>2</sub> in EC, a larger free solvent feature was obtained as compared to the Ca system, in agreement with the higher degree of ion-pair formation. In all cases, the threshold concentration needed to reach the highly concentrated electrolyte domain is much lower for systems with divalent cations as compared to monovalent cation based electrolytes (>3 M), in agreement with the higher coordination number of Ca and Mg (typically > 6) than that of Li (typically 4). 12,16

Understanding the solvation structure of battery electrolytes is a crucial step in the quest for electrolyte optimization. Indeed, the formation of cation-anion pairs effectively diminishes the amount of charge carriers, and a cation in solution needs to get rid of its solvation shell before plating or insertion can take place at the anode and cathode, respectively. The ease with which the cation desolvates is related to its binding energy with other species. Binding energies were calculated for the M-X complexes (Table S6), and much larger energies were obtained for the ion pairs; M-TFSI with 1162 and 1442 kJ  $\text{mol}^{-1}$  for M = Ca and Mg, respectively, as compared to the cation-solvent interactions; M-EC with 380 and 544 kJ  $\text{mol}^{-1}$  for M = Ca or Mg, respectively. Thus, as far as interfacial processes are concerned, Ca-based systems present prospects for better power performance than Mg. For both systems, a solvation shell composed mainly of solvent molecules is preferred (absence of ion pairs), which can be achieved by using high donor number solvents and moderate salt concentrations.

### CONCLUSIONS

Ca-based electrolytes were studied, and the influence of salt, concentration, and the solvent was investigated in terms of ionic conductivity, viscosity, ionicity, and solvation structure and in some cases compared with Li and Mg solutions. One of the main issues for further development of divalent-cation-based electrolytes, in general and also true for Ca batteries, is the salt solubility—resulting in only a few Ca-salts possible and only when combined with high-permittivity solvents. The

degree of contact ion-pair formation and mono- vs bidentate coordination were both found to evolve with salt concentration and solvent donor number. The amount of free solvent molecules evolves accordingly, and divalent cation based electrolytes enter the domain of highly concentrated electrolytes at concentrations as low as 1.2 M. We found that a high donor number solvent, such as DMF, allows for both low viscosities, high ion conductivities, as well as few contact ion pairs (although solvent-separated ion pairs can be envisaged, resulting in low ionicity). The lower probability for contact ion pairs, together with the lower interaction energies within their solvation shell, is fundamentally promising for better kinetics of metal plating and cation insertion.

## ASSOCIATED CONTENT

# **S** Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpcc.9b07308.

Physicochemical properties of salts, solvents, and calcium electrolytes; Vogel—Tammann—Fulcher (VTF) parameters of viscosities and conductivities; infrared and Raman spectra of calcium electrolytes; deconvolution of Raman spectra; interaction/binding energies; Raman parameters for fitting (PDF)

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### **Notes**

The authors declare no competing financial interest.

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