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A Fundamental Study on the \([(\mu-\text{Cl})_3\text{Mg}_2(\text{THF})_6]^+\)
Dimer Electrolytes for Rechargeable Mg Batteries

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To address the global problems of escalating energy demand and increasing emissions of CO₂, it is critical to develop effective approaches to convert renewable but intermittently available energy sources (e.g. solar and wind energy) into reliable energy forms. Battery systems with low cost, high energy density and long cycling life have been suggested as viable technologies for storing sustainable energy.1 Recently rechargeable Magnesium (Mg) batteries have been advocated as promising battery systems alternative to lithium or sodium based batteries for grid scale energy storage, powering portable devices, and transportation applications.2,3 There are several technic advantages of Mg batteries over Li batteries and Na batteries. Mg is earth abundant and low cost (ca. 24 times cheaper than Li). As an anode material, Mg is safe to use without dendrite formation (vs Li, Li-ion or Na batteries) and high volumetric capacity (3832 Ah/L vs 2062 Ah/L for Li and 1165 Ah/L for Na) due to its two-electron redox chemistry. Mg based electrolytes are environmentally benign. Furthermore, Mg has a sufficient high reduction potential (-2.37 V vs SHE) amenable for assembling high energy density batteries with suitable cathode materials.

In 1990, Gregory et al. made a major breakthrough on electrochemical Mg deposition by using Al or B Lewis acid additives to enhance electrochemical performance of Grignard reagents and MgBu₂ electrolytes.4 Since then, there have been continuous efforts in developing Mg conductive electrolytes for rechargeable Mg batteries by Aurbach2 and others.5 We have put our efforts in developing advanced Mg electrolytes with improved oxidation stability and electrolyte compatibility by avoiding the use of nucleophilic Mg sources such as Grignard reagents or dialkyl magnesium.6 Poor oxidation stability and/or undesired nucleophilicity of the reported electrolytes prepared from nucleophilic Mg sources are attributed to reactive alkyl or aryl anions in these reagents.4b Based on a straightforward retrosynthesis analysis on the \([(\mu-\text{Cl})_3\text{Mg}_2(\text{THF})_6]^+\) dimer, a common recrystallized product of popular Mg-Cl complex electrolytes,4b,5c,7 we developed a facile approach (termed as \textit{mono-Cl abstraction, see Scheme 1}) using MgCl₂ and an Al Lewis acid to produce high performance Mg-dimer containing electrolytes.6

The MgCl₂-Al Lewis acid electrolytes are much cleaner compared to electrolytes made from nucleophilic Mg reagents in terms of composition because of the simply and clean Cl⁻ metathesis reaction without complication by nucleophilic alkyl or aryl species.5c Therefore, these electrolytes provide a unique opportunity to understand solution chemistry of the \([(\mu-\text{Cl})_3\text{Mg}_2(\text{THF})_6]^+\) dimer and further its electrochemical behaviors. Previously, we proposed a reaction mechanism for the formation of the Mg-dimer coordination complex from MgCl₂ and an Al Lewis acid in the mono-Cl approach (see Scheme 1). THF solvated MgCl₂, \textit{trans-MgCl₂(THF)₂}, undergoes THF dissociation and subsequent \textit{mono-Cl abstraction} by an Al Lewis acid (e.g. AlPh₃ in Scheme 1) to generate the reactive intermediate \([\text{MgCl}^+\text{Cl}]^-\) cation, \([\text{MgCl(THF)₃}]^+\). Coordination unsaturation of \([\text{MgCl(THF)₃}]^+\) can promote a dimerization reaction with MgCl₂(THF) to result in the Mg-dimer cation.4b,6 Both \([(\mu-\text{Cl})_3\text{Mg}_2(\text{THF})_6]^+\) and \([\text{MgCl(THF)₃}]^+\) have been proposed to be the active species responsible for electrochemical Mg cycling. However, there is no experimental evidence for the existence of THF solvated \([\text{MgCl}^+\text{Cl}]^-\) in the solution of dimer electrolytes.4b In this study, we report the first experimental evidence of a solution \([\text{MgCl}^+\text{Cl}]^-\) species, \([\text{MgCl(THF)₃}]^+\), in Mg dimer electrolytes using SPIN (Subambient Pressure Ionization with Nanoelectrospray) mass spectrometry.8
combination with additional results, the coordination chemistry of the Mg-dimer electrolytes is elucidated and electrochemical mechanisms involving Mg\(^{2+}\) and Cl\(^{-}\) transports for reversible Mg deposition and stripping of the Mg dimer electrolytes are proposed and discussed.

Figure 1. The M/Z isotopic distribution of the [MgCl(THF)]\(^{+}\) peak in the positive mode of SPIN MS of [(μ-Cl)\(_{3}\)Mg(THF)]\(_{2}\)AlPh\(_{3}\)Cl. Inset gives the calculated isotopic pattern for [MgCl(THF)]\(^{+}\).

In an effort to further understand the chemical nature of the Mg dimer electrolytes and elucidate the proposed reaction mechanism,\(^6\) we have attempted to detect the [MgCl]\(^{+}\) mono-cation in our electrolyte solution. We first tried conventional electrospray ionization (ESI) MS as the technique can resolve charged species through values of mass charge ratio (M/Z) and corresponding isotopic patterns. However, there was no meaningful species detected in the positive mode of ESI-MS with [(μ-Cl)\(_{3}\)MgCl(THF)]\(_{2}\)AlPh\(_{3}\)Cl at different concentrations (from 0.05 mM up to 5 mM). Similar results were obtained with [(μ-Cl)\(_{3}\)MgCl(THF)]\(_{2}\)AlEtCl\(_{3}\) and [(μ-Cl)\(_{3}\)MgCl(THF)]\(_{2}\)AlCl\(_{4}\), electrolytes.\(^6\) We rationalized Mg-THF coordination interaction is too labile to survive under typical ESI-MS interface configuration which operates with an inlet temperature of 200 °C. So we started to seek a MS technique with a softer ESI source. We found out Subambient Pressure Ionization with Nanoelectrospray Mass spectrometry (SPIN-MS)\(^8\) is a unique MS interface technique with high ionization efficiency and a softer ion source to enable the characterization of compounds with weak chemical bonds with high sensitivity. The SPIN source can be operated at a temperature below 80 °C and may provide a chance to detect positively charged THF solvated [MgCl]\(^{+}\) species. To our delight, a [MgCl]\(^{+}\) species with three THF coordinated, [MgCl(THF)]\(^{+}\), was consistently identified at 275.13 M/Z by SPIN-MS (Figure 1) for [(μ-Cl)\(_{3}\)Mg(THF)]\(_{2}\)AlPh\(_{3}\)Cl THF solution. The match of the experimental and calculated isotope patterns further confirms its identity (Figure 1). [MgCl(THF)]\(^{+}\) is either directly from the solution or generated from [MgCl(THF)]\(^{+}\) during the ionization process. In the later case, the result suggests Mg-THF bonds in [MgCl(THF)]\(^{+}\) is stronger than the ones in [MgCl(THF)]\(^{+}\). However, no signal was observed for the dimer cation. Preliminary DFT calculations suggest the [MgCl]\(^{+}\) monomer is more dominant over the dimer cation in the equilibrium, which explains a lower possibility to observe the dimer cation. A recent study also suggests the dimer is preferred in solid state.\(^9\) Another possibility is the lack of sufficient stability of the dimer cation under the testing conditions. Consistently, the [MgCl(THF)]\(^{+}\) cation was also clearly identified in the [(μ-Cl)\(_{3}\)MgCl(THF)]\(_{2}\)AlEtCl\(_{3}\) electrolyte (Figure S4). The SPIN-MS studies of the MgCl\(_{2}\) THF solution without adding an Al Lewis acid did no show the monocation signal.

To the best of our knowledge, the SPIN MS result is the first experimental evidence for the existence of THF solvated [MgCl]\(^{+}\) mono-cation in the solution of [(μ-Cl)\(_{3}\)MgCl(THF)]\(_{2}\) dimer electrolytes. The identification of the [MgCl(THF)]\(^{+}\) mono-cation establishes the reaction nature of mono-Cl abstraction of solvated MgCl\(_{2}\) by an Al Lewis acid as described above (see Scheme 1).

Raman spectroscopic studies of [(μ-Cl)\(_{3}\)MgCl(THF)]\(_{2}\)AlPh\(_{3}\)Cl in solution were first attempted but no resolved spectrum was obtained due to strong fluorescence. The Raman spectrum of [(μ-Cl)\(_{3}\)MgCl(THF)]\(_{2}\)AlPh\(_{3}\)Cl in solid state confirmed the presence of the solvated MgCl\(_{2}\) species, showing a diagnostic peak at 209 cm\(^{-1}\) (Figure 2). The weak peak at 252 cm\(^{-1}\) is assigned to [(μ-Cl)\(_{3}\)MgCl(THF)]\(_{2}\)^+ and two peaks at 618 and 653 cm\(^{-1}\) are attributed to phenyl ligand ring wagging of the AlPh\(_{3}\)Cl anion.\(^6\) The SPIN Mass, Raman spectroscopic results, and the established solid structure of [(μ-Cl)\(_{3}\)MgCl(THF)]\(_{2}\)AlPh\(_{3}\)Cl\(^5\) provide a full experimental picture on the equilibrium between [(μ-Cl)\(_{3}\)MgCl(THF)]\(_{2}\) and THF solvated [MgCl]\(^{+}\) and MgCl\(_{2}\).

CT\(^{−}\) is the primary ligand in supporting the coordination structures of solution Mg\(^{2+}\) species and defines their electronic structures, i.e. frontier HOMO and LUMO energy that are closely associated with redox potentials of electron transfer reactions. Studying CT ligand behaviors could lead to further insights into the [(μ-Cl)\(_{3}\)MgCl(THF)]\(^{+}\) dimer and its equilibrium in terms of CT\(^{−}\) and Mg\(^{2+}\) ion transports, and electrochemical Mg deposition and stripping. Aurbach \textit{et al.} reported external CT\(^{−}\) anion can affect the chemical equilibrium of [(μ-Cl)\(_{3}\)MgCl(THF)]\(^{+}\) and suggested [MgCl]\(^{2+}\) species is formed in the reaction of the APC electrolyte with Cl\(^{−}\). However, the presence of MgPh\(_{3}\) in the APC electrolyte\(^{6}\) can also react with external CT\(^{−}\) due to the Schlenk equilibrium and complicate data interpretation. As our electrolytes are less complex than Aurbach’s all phenyl complex (APC) electrolytes,\(^6\) we anticipate our electrolytes would provide new insights into the reactivities of the [(μ-Cl)\(_{3}\)MgCl(THF)]\(_{2}\) dimer electrolyte with CT\(^{−}\). So we have investigated the Cl\(^{−}\) effect on the [(μ-Cl)\(_{3}\)MgCl(THF)]\(_{2}\)AlPh\(_{3}\)Cl electrolyte using various physical methods.

\(^{25}\)Mg\(^{1}\)H NMR studies were carried out to monitor the effect of Cl\(^{−}\) on the \(^{25}\)Mg chemical shift of [(μ-Cl)\(_{3}\)MgCl(THF)]\(_{2}\). As shown in Figure 3, [(μ-Cl)\(_{3}\)MgCl(THF)]\(_{2}\)AlPh\(_{3}\)Cl exhibits a broad singlet at 5.3 ppm, consistent with an exchange process of the dimer with solvated [MgCl\(_{2}\)] and [MgCl]\(^{+}\) species. With addition of tetrabutylammonium chloride (TBACl) as the Cl\(^{−}\) source, the Mg resonance gradually shifts from 5.3 ppm to lower field up to 11.9
ppm at 0.1 M Cl (Figure 3 and Figure S1). In the same time, the peak width of the Mg resonances broadens from 207.5 Hz to 419 Hz (see Figure S1), indicating a slower exchange process with increasing Cl concentration. The changes of the chemical shift and its peak width are consistent with formation of new Mg species or a change of the chemical equilibrium of [(µ-Cl)3Mg2(THF)6]− upon addition of Cl−. During the addition of Cl−, there was no precipitate of MgCl2 formed. The solution after adding Cl− stayed in clear for days. No obvious spectroscopic change was observed in the 27Al (1H) NMR spectrum after addition of Cl− (Figure S2). To gain further understandings of the chemical equilibrium, the solution of 1:1 ratio of [(µ-Cl)3Mg2(THF)6]AlPh3Cl and TBACl was examined by SPIN MS. It was found out the signal of [MgCl(THF)3]− at 275.2 ppm became significantly more intensive than the sample of [(µ-Cl)3Mg2(THF)6]AlPh3Cl without adding Cl− (Figure S3). However, attempted SPIN-MS negative mode experiments did not identify new Mg species, either of [MgCl(THF)3]− or [MgCl4]2−.

![Figure 3. 25Mg NMR spectra of [(µ-Cl)3Mg2(THF)6]AlPh3Cl, 0.1 M in THF (collected at 25 °C) in the presence of tetrabutylammonium chloride (TBACl) at different concentrations as indicated.](image)

Figure 4 displays the cyclic voltammograms of [(µ-Cl)3Mg2(THF)6]AlPh3Cl at various concentrations of TBACl. With increasing Cl− concentration to 0.02 M, the current intensities of Mg deposition and stripping initially increased, consistent with enhanced solution conductivity (Figure S6). However, when Cl− concentration is above 0.03 M, the current start to decrease. To clearly demonstrate the experimental observation, a plot of Mg deposition and stripping charge versus Cl− concentration is given in Figure S6. In the meanwhile, solution conductivity (0.96 mS/cm at 0.02 M Cl−) started to decrease while there is not an apparent relationship between solution conductivity and Cl− concentration (Figure S6). The effect of Cl− on the electrochemical behaviors of [(µ-Cl)3Mg2(THF)6]AlPh3Cl is different from what was observed for Aurbach’s APC electrolyte.10 The difference is ascribed to the different solution constitutions of two electrolytes. During the addition of Cl−, coulombic efficiencies for Mg cycling are retained ca. 100% and overpotential stays at ca. 0.29 V. The decrease of Mg cycling current at higher Cl− concentration is believed to be ascribed to the change of the chemical equilibrium of the electrolyte solution with Cl− as indicated by the 25Mg NMR spectroscopic studies. There are two possibilities: (a) If the [(µ-Cl)3Mg2(THF)6]− dimer is the active specie for Mg deposition and stripping, then addition of Cl− can shift the equilibrium to the mononuclear Mg species and lower the concentration of [(µ-Cl)3Mg2(THF)6]−; (b) If the solvated [MgCl2]− is the active specie, the added Cl− can complex with [MgCl2]− to form solvated MgCl2 to lower the concentration of the mononic cation. If MgCl2 had formed, it would have precipitated out of the solution as it is barely soluble in THF. One explanation is the formed [MgCl2]− could complex with the dimer cation to form a soluble trimmer cation, [Mg2(µ2-Cl)2(µ2-Cl)(THF)4]11 (Scheme 2), consistent with the observed downshift of 25Mg resonance. It is less likely that [MgCl2]− can react with Cl− to form [MgCl4]2− and further [MgCl6]4− dianion10 as these Mg anions would lead to high-field shift of the averaged Mg resonance. We are conducting comprehensive DFT calculations to gain insights to these possible reactions.

![Figure 4. Cyclic voltammograms of [(µ-Cl)3Mg2(THF)6]AlPh3Cl, 0.1 M in the presence of various concentrations of TBACl as indicated. For clearance, cyclic voltammograms with TBACl of 0.01, 0.03, 0.05, 0.07, 0.09 M are given in Figure S5 in the supporting information; conditions: Pt working electrode, Mg reference electrode, glassy carbon counter electrode, 50 mV/s.](image)

In terms of the solution nature of the dimer electrolyte and its reactivity with external Cl− described above, comprehensive mechanisms are proposed for Mg2+ and Cl− transports and their involvements with Mg deposition and stripping for the dimer electrolyte in the operation of Mg batteries (scheme 2). The proposed mechanism is an updated version from Aurbach’s mechanism5 by providing new aspects on Mg electrolyte chemistry. For discussion purpose, [MgCl2(THF)3]− and cis-MgCl2(THF)2 will be used as primary equilibrumers with [(µ-Cl)3Mg2(THF)6]− although they can undergo THF solvation to form [MgCl2(THF)3]− and trans-MgCl2(THF)2 (Scheme 1). Either of [(µ-Cl)3Mg2(THF)6]− or [MgCl2(THF)3]− mono cation on the anodic electrode surface participates the first electron transfer reaction to form a neutral [MgCl2]0 absorbed on the electrode. It should be noted that our preliminary DFT calculations suggest the [MgCl2]− cation is predominant in solution. Then surface [MgCl2]0 species undergoes a second electron reduction to result in metallic Mg. It is believed the intermediate of [MgCl2]0 is the key to the dendrite free deposition of Mg using [(µ-Cl)3Mg2(THF)6]2− electrolytes and its formation is the rate determining step for Mg deposition, providing a kinetic control for overall electron transfer reactions. However, there is no experimental evidence reported for the surface bound species. Such principle can be applied to other anion supported Mg electrolytes.2b, 15 In the meanwhile, the released Cl− will be trapped by [MgCl2]− to form [MgCl4]2− species and subsequently [Mg2(µ2-Cl)2(µ2-Cl)(THF)4]− (Scheme 2a). Attentively, the solvated Mg2+ could directly react with [MgCl2]− to regenerate [MgCl2]− (Scheme 2b).
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Notes and references

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Conclusions

We have experimentally identified the long-sought solvated $[\text{MgCl}]^+$ species in the $[(\mu-\text{Cl})\text{Mg}_2(\text{THF})_6]^+$ electrolyte solution using the SPIN-MS technique, a unique tool to study weakly coordinating complexes. The identified $[\text{MgCl}]^+$ species confirmed the reaction nature of mono-Cl$^-$ abstraction of solvated MgCl$_2$ with an Al Lewis acid and its role in forming the well known $[(\mu-\text{Cl})\text{Mg}_2(\text{THF})_6]^+$ dimer. We suggest both $[(\mu-\text{Cl})\text{Mg}_2(\text{THF})_6]^+$ and $[\text{MgCl}(\text{THF})_3]^+$ as mono-cations can be the active species responsible for Mg deposition. Synergetic operation mechanisms of co-existing solvated MgCl$_2$ and $[\text{MgCl}]^+$ was proposed for electrochemical Mg cycling and Mg$^{2+}$ and Cl$^-$ transports in the operation of Mg batteries. The present fundamental understandings of the coordination chemistry of the Mg-Cl complex electrolytes will inspire synthetic design of next generation of Mg electrolytes for rechargeable Mg batteries.

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