Ultrafast fluxional exchange dynamics in electrolyte solvation sheath of lithium ion battery

Kyung-Koo Lee1, Kwanghee Park2,3, Hochan Lee2,3, Yohan Noh2, Dorota Kossowska2,3, Kyungwon Kwak2,3 & Minhaeng Cho2,3

Lithium cation is the charge carrier in lithium-ion battery. Electrolyte solution in lithium-ion battery is usually based on mixed solvents consisting of polar carbonates with different aliphatic chains. Despite various experimental evidences indicating that lithium ion forms a rigid and stable solvation sheath through electrostatic interactions with polar carbonates, both the lithium solvation structure and more importantly fluctuation dynamics and functional role of carbonate solvent molecules have not been fully elucidated yet with femtosecond vibrational spectroscopic methods. Here we investigate the ultrafast carbonate solvent exchange dynamics around lithium ions in electrolyte solutions with coherent two-dimensional infrared spectroscopy and find that the time constants of the formation and dissociation of lithium-ion \( \cdot \cdot \cdot \) carbonate complex in solvation sheaths are on a picosecond timescale. We anticipate that such ultrafast microscopic fluxional processes in lithium-solvent complexes could provide an important clue to understanding macroscopic mobility of lithium cation in lithium-ion battery on a molecular level.
Electrolyte solvent capable of dissolving lithium salt is an integral part of Li-ion battery (LIB) that has become an indispensable power source for current mobile ecosystem and electrical vehicles. Among various components in LIB, the properties and dynamics of carbonate-based solvents govern the mobility and stability of charge carrier, Li⁺ ion, in LIB. Because Li⁺ ion mobility is much lower than electrons in metal wires, the solvation dynamics of Li⁺ in electrolyte solutions is crucial in determining the LIB performance. Typical electrolyte solutions used in LIB are composed of non-aqueous carbonate solvents that have certain advantages. For instance, the oxidation potential can be matched to the output voltage of LIB and carbonate solvents are also building blocks of solid-electrolyte interphase.

To achieve high Li⁺ conductivity, mixed solvents have often been used not only because they increase the solubility of lithium salt but also because they serve as excellent media facilitating fast transport of Li⁺ ions. Cyclic carbonates, such as propylene carbonate (PC) and ethylene carbonate (EC), have high dielectric constants so that they enhance the dissociation of lithium salt by stabilizing ions. However, their high viscosity limits the fast Li⁺ transport. To enhance Li⁺ mobility as well as to improve low-temperature performance, linear alkyl carbonates like diethyl carbonate (DEC) and dimethyl carbonate (DMC) are added to the LIB electrolyte solutions as co-solvents. It has long been believed that such low viscosity co-solvents act as a medium for transporting the sheaths of Li⁺ ion-solvent complex. In the current commercial LIBs, lithium salt, 1.0 M LiPF₆, is dissolved in organic carbonate mixed solvents consisting of equal volumetric amounts of DEC, DMC, and EC. Li⁺ is believed to be preferentially solvated by EC, and DEC and/or DMC provides the medium for large Li⁺–EC complexes to flow between electrodes during charging and discharging processes. However, to our surprise, detailed microscopic solvation structures and ultrafast solvation dynamics around Li⁺ in such electrolyte solutions have not been fully understood and even they are still the key subjects under intense scholarly debate.

Therefore, elucidating the molecular solvation structure of organic carbonates around Li⁺ would, without a doubt, help one to better understand the working principles of LIB and to eventually design an improved LIB. Most of the previous studies primarily focused on the equilibrium structure of the first solvation shell of Li⁺, which has often been referred to as solvation sheath due to its rigidity. For instance, Raman spectroscopy of LIB electrolyte solutions revealed the existence of preferential solvation shell around Li⁺ in mixed solvent electrolyte solutions. This was also confirmed by Bogle et al. where ¹⁷O NMR experiments show that Li⁺ makes a direct electrostatic interaction with carbonyl oxygen atoms of surrounding carbonates. Even in the case of Li⁺ is preferentially solvated by EC or PC, the participation of co-solvent molecules, such as DEC and DMC, in the solvation sheath is possible not only because of their high-mole fractions but also because of extra entropic gain. Therefore, it is by now clear that such co-solvents could potentially play a dual-role of low-viscous medium and solvating molecule. Co-solvents like DEC and DMC would add an extra flexibility to solvation sheath, because they comparatively weakly interact with Li⁺ and have many conformational degrees of freedom. Nevertheless, no direct evidence of fast fluxional solvent exchange dynamics in Li⁺ solvation sheath has been reported due to their ultrafast nature and experimental difficulties in finding an appropriate spectroscopic probe.

In this regard, we believe that time-resolved nonlinear infrared spectroscopy is an ideal method capable of providing critical information on ultrafast chemical exchanges between different solvation structures that cannot be easily distinguished by NMR or other spectroscopic means due to their limited time-resolving power or to a lack of spectroscopic probe enabling one to track local configurational changes around Li⁺ ion. Here considering the carbonyl stretching vibration of DEC as an infrared probe, we carried out Fourier transform infrared (FTIR), femtosecond infrared pump-probe and two-dimensional (2D) infrared studies of chemical exchange processes between DEC molecules with and without Coulombic interaction with Li⁺. One of the most difficult problems in the present linear and nonlinear infrared spectroscopic investigations is that the infrared probe, DEC, itself is the solvent too. Thus, with normal infrared cells with several micrometre thickness at best, the linear absorption is saturated and the nonlinear infrared signal fields are self-attenuated, which renders a difficulty in detecting weak 2D infrared signal. To overcome this, we prepared a home-built thin sample cell coated with SiO₂, using radio frequency magnetron sputtering technique (see Fig. 1a,b and Supplementary Fig. 1). Here we show evidence on the ultrafast formation and dissociation dynamics of Li⁺-carbonate solvation complexes in the LIB electrolyte solutions by analysing the FTIR spectra and time-resolved infrared pump-probe and 2D infrared data of C=O stretch modes in lithium salt electrolyte solutions and carrying out quantum chemistry calculations. In particular, the time constant associated with the breaking of Li⁺–DEC electrostatic interaction is found to be 2.2 ps and that with the formation of the Li⁺–DEC bond is 17.5 ps at 1.0 M LiPF₆ concentration in DEC solvent. The timescales of making and breaking of Li⁺–DEC interaction bonds in PC/DMC mixed solvents are similar to those for DEC solutions, which indicates that the DEC molecules also participate in the formation of lithium cation solvation shell even in such mixed electrolyte solutions used in a real LIB. We anticipate that such ultrafast fluxional dynamics in solvation shell structures around lithium cations are of importance in understanding lithium-ion mobility in LIB solutions.

**Results**

**FTIR spectroscopy.** The infrared absorption spectra of the C=O stretch and O–C–O asymmetric stretch modes in LiPF₆ DEC solutions at different concentrations are shown in Fig. 1c,d, respectively. DEC has three strongly infrared-active modes in the fingerprint region, which are C=O stretching (1,747 cm⁻¹), O–C–O asymmetric stretching (1,258.9 cm⁻¹), and O–CH₂–CH₃ asymmetric stretching (1,022 cm⁻¹) vibrations. As LiPF₆ concentration in the DEC solution increases, two distinctively new peaks at 1,715.4 and 1,305.6 cm⁻¹ appear. The frequencies and spectral widths of the main peaks at 1,747 and 1,258.9 cm⁻¹ do not change much on increasing lithium salt concentration. Therefore, the two new peaks at 1,715.4 and 1,305.6 cm⁻¹ can be safely assigned to the C=O stretch and O–C–O asymmetric stretch modes of DEC molecules directly interacting with Li⁺.

However, it is noted that a DEC molecule has three different sites for electrostatic interaction with Li⁺. They are two ester ether oxygen atoms and one carbonyl oxygen atom. Therefore, at least two structures in Fig. 2 should be considered as possible Li–DEC complexes. Despite that a recent ¹⁵O NMR experiment provided evidence that the ester ether oxygen atoms do not strongly interact with Li-ion, it is not entirely clear whether this is the case even at high (1.0 M) lithium salt concentration. To pin point the Li–DEC complex structure among the two in Fig. 2, quantum chemistry calculations and vibrational analyses of
a few complexes were performed with DFT method at the level of B3LYP/6-311++G(3df,2pd). For an isolated DEC, the C–O stretch and O–C–O asymmetric stretch frequencies are found to be 1,714.4 cm$^{-1}$ and 1,235.5 cm$^{-1}$, respectively (detailed descriptions on the quantum chemistry calculation results are found in Supplementary Note 2). Then, the DFT calculations of two structures in Fig. 2 show that the C–O stretch frequency is red shifted by 112.2 cm$^{-1}$ and simultaneously the O–C–O asymmetric stretch frequency is blue shifted by 62.6 cm$^{-1}$ (Supplementary Table 1). In contrast, the complex formation of Li$^+$ with two ester ether oxygen atoms of DEC induces strong blue shift (112.4 cm$^{-1}$) of C–O frequency and red shift (96.8 cm$^{-1}$) of O–C–O asymmetric stretch frequency. These results can be understood by noting that the Li$^+$ ion interacting with carbonyl oxygen atom makes the double bond character (or bond length) of C–O group decrease (increase), which concomitantly makes the C–O bond lengths short (Supplementary Table 1). When Li$^+$ electrostatically interacts with ester ether oxygen atom, exactly the opposite charge redistribution occurs. Thus, our infrared results are consistent with the previous notion that Li$^+$ prefersentially interacts with carbonyl oxygen atom (Fig. 2a)$^{28,29}$. In fact, such frequency shifting behaviours of C=O and O–C–O modes in DEC on the formation of C=O···Li$^+$ complex are quite similar to the red and blue shifts of amide I and II modes of peptide bond, respectively, which are mainly C=O stretch and C–N stretch of the peptide bond, respectively, when the amide carbonyl oxygen atom forms a hydrogen bond with a water molecule. Despite that the quantum chemistry calculations for the two different complex structures in Fig. 2 are consistent with previous findings and our experimental FT-IR results discussed below, the C=O stretch frequency may depend also on the side chain (ethyl group) conformation. More specifically, the molecular dipole moment of the cis conformation can be larger than that of the trans conformation so that the cis conformer can make a bit stronger interaction with Li$^+$ and form a more stable Li–DEC complex.

The number of solvent molecules in the first solvation shell of a solute molecule is an important quantity providing information on the size and structure of Li-ion solvation sheath. Often, a series of lithium salt concentration-dependent spectroscopic experiments has been used to determine the solvation number around a Li$^+$ in various electrolyte solutions$^{28,29}$. However, Doucey et al.$^{13}$ showed that the formation of contact
ion pair (CIP) between two oppositely charged ions should also be taken into account in quantitatively interpreting the infrared spectra of LIB solutions. That means, the concentration of dissolved LiPF$_6$ is not necessarily identical to the concentration of DEC-solvated Li$^+$ ion because there could exist non-negligible amount of Li$^+$:PF$_6^-$ CIP. Thus, it is necessary to consider three different DEC species that are (i) DEC surrounded by other DECs, (ii) DEC interacting with free Li$^+$ and (iii) DEC interacting with Li$^+$ in the CIP state, Li$^+$:PF$_6^-$. In addition, the existence of two different types of anions, either PF$_6^-$ solvated by DECs or PF$_6^-$ in CIP imposes yet another complexity to the interpretation of linear and nonlinear infrared spectra. However, because the negatively charged PF$_6^-$ tends to be away from the carbonate part of DEC due to Coulomb repulsion and preferentially surrounded by the DEC ethyl groups$^{29}$, PF$_6^-$ does not strongly affect the DEC carbonyl stretch infrared spectrum.

To gain more insight into the vibrational properties of DEC...Li$^+$:PF$_6^-$ as compared to DEC...Li$^+$, we further analysed density functional theory calculation results of DEC interacting with Li$^+$:PF$_6^-$ (Supplementary Fig. 4). Even in the case of Li$^+$ in Li$^+$:PF$_6^-$, still the carbonyl oxygen atom of DEC is the binding site forming C=O...Li$^+$:PF$_6^-$ complex, though the C=O frequency red shift in that case is a bit smaller than that of C=O...Li$^+$. If the first solvation shell structure around free Li$^+$ is spectroscopically different from that around Li$^+$:PF$_6^-$, one might expect to see clear signature, that is, a new third infrared absorption band or shoulder in the lower-frequency region. A careful examination of the C=O band of LiPF$_6$ DEC solution, where the total C=O stretch infrared band was corrected by subtracting that of pure DEC liquid, shows no discernible spectroscopic feature of DEC-CIP (Supplementary Fig. 3). Thus, it is concluded that the carbonyl stretch infrared band consists of indistinguishable contributions from both C=O...Li$^+$ and C=O...Li$^+$:PF$_6^-$ complexes, which altogether will be referred to as Li--DEC complex.

**Polarization-controlled infrared pump-probe.** Figure 3a,b displays the dispersive isotropic pump-probe signals obtained from pure DEC liquid and 1.0 M LiPF$_6$ DEC solution, respectively. The time-resolved pump-probe spectra of LiPF$_6$ in DEC solution show complicated spectral features due to the destructive interference between the negative 1–2 transition peak (excited state absorption transition from |1> to |2>) of free DEC and the positive 0–1 peak (ground state |0> bleach and stimulated emission from |1> to |0>) of Li--DEC complex at around 1,720 cm$^{-1}$. Another notable feature is the prolonged positive signal (around 1,750 cm$^{-1}$ in Fig. 3a,b) that is associated with local heating effect due to the vibrational energy relaxation of infrared-pump-excited C=O stretch modes$^{31}$. The vibrational lifetime could be extracted from the fitting analysis of the 1–2 transition PP data that are relatively immune to local heating effect. Note that the decay constants of the excited state absorption signals at around 1,725 cm$^{-1}$ and 1,700 cm$^{-1}$, respectively, for pure DEC liquid and 1.0 M LiPF$_6$ DEC solution, are little dependent on the probe frequency (Supplementary Fig. 5). On the other hand, the exponential decay constants obtained from the fits of positive 0–1 signals at ∼1,750 cm$^{-1}$ show a strong probe-frequency dependence, which is mainly due to the local heating contribution to the infrared PP signal. Thus, the isotropic and anisotropic data at the 1–2 transition frequencies at ∼1,700 cm$^{-1}$ were taken into consideration to extract information on the vibrational and orientational relaxation times of Li--DEC complex.

The corresponding time constants of free DEC were obtained from the excited state absorption signal at 1,725 cm$^{-1}$ in the infrared PP data of pure DEC solvent (Fig. 3a).

Vibrational lifetime of Li--DEC complex (1.1 ± 0.1 ps) is found to be much shorter than that of free DEC (2.1 ± 0.1 ps). Detailed discussion on the origin of such increased vibrational relaxation rate of DEC upon its electrostatic interaction with Li$^+$ can be found in Supplementary Note 3. The anisotropic infrared pump-probe signals of free DEC and Li--DEC enabled us to determine the orientational relaxation rates. As expected, the Li--DEC complex with three or four DEC molecules with or without ion-pairing PF$_6^-$ rotates slowly as compared to free DEC and the second-order rotational correlation time constants are found to be 8.3 ± 0.6 ps and 1.5 ± 0.1 ps for Li--DEC complex and free DEC, respectively.

**2D infrared chemical exchange spectroscopy.** Figure 3c shows four representative time-resolved 2D infrared spectra of C=O stretch mode of DEC in 1.0 M LiPF$_6$ DEC solution at different waiting times. Each 2D infrared spectrum was normalized to the largest positive 0–1 peak magnitude. At a very short waiting time of $T_w = 0.3$ ps, there are only two peaks on the diagonal line (see the upper-left panel in Fig. 3c), which correspond to the two absorption peaks in the FTIR spectrum shown in Fig. 1c. The positive peaks (red colour) result from the ground state bleaching and stimulated emission contributions, whereas the negative peaks (blue colour) in the lower frequency region along the probe frequency ($\omega_o$) axis are the excited state absorption contributions. Both positive and negative peaks are elongated along the diagonal line at short waiting time up to $T_w = 1.0$ ps. However, after 2.0 ps, all peaks become round in shape due to the spectral diffusion process. Diagonally elongated peaks at short waiting times indicate structural heterogeneity, distribution of coordination number and ethyl side-chain conformational distribution (cis-cis, cis-trans, and trans-trans)$^{29}$ in DEC in solvation complexes around Li$^+$. We found that the C=O stretch frequency--frequency correlation decays on a few picosecond timescale, which could result from ultrafast internal rotations of the ethyl groups in DEC and hindered rotational motions of carbonates.

In principle, the inhomogeneous structural distribution can be studied by analysing the centre or nodal line slope and/or diagonal ellipticities of 2D infrared peaks$^{32–34}$. Unfortunately, due to the destructive interference between the positive 0–1 peak from Li--DEC and the negative 1–2 peak from free DEC, it is difficult to clearly identify the corresponding 2D infrared feature. Thus, we compared the infrared absorption line-widths of free DEC and Li--DEC complex with the help of pump-probe data. The FWHM (full-width at half-maximum) of the infrared spectrum of Li--DEC is found to be 22.2 cm$^{-1}$, which is slightly broader than that of free DEC (FWHM = 18.9 cm$^{-1}$). However, the vibrational lifetime of Li--DEC is just half of that of the free DEC, which means that the lifetime broadening of Li--DEC is larger than that of free DEC. These observations indicate that the inhomogeneous linewidth of Li--DEC may be narrower than free DEC, which is consistent with the notion that DEC molecules in Li--DEC complexes have stable conformers and the side-chain structure can be correlated with its solvation property$^{25}$.

In a real LIB electrolyte solution, detailed solvation dynamics around Li$^+$ could be affected by the presence of co-solvents in the first solvation shell. In a recent $^{17}$O NMR experiment, NMR peak of carbonyl oxygen atom in EC or DMC shows a motional narrowing behaviour, indicating the existence of fast equilibrium dynamics between free carbonate and Li-complexed carbonate molecules in nanosecond or even shorter timescales$^{20}$. 
Interestingly, it has long been believed that such fast solvent exchange dynamics can occur only in the second solvation shell because of the belief that the first solvation shell maintains its rigid structure during the Li-transport between electrodes\(^5,6\). However, if Li-solvent complexes undergo fast solvent exchange dynamics resulting in ultrafast fluctuation of coordination number and solvation structure, the conventional view on the robust and rigid solvation sheath formation must be reconsidered. Especially the coordination number fluctuation would modulate the diffusion rate of Li\(^+\) in LIB electrolyte solutions. To date, however, little has been understood about how fast the solvation dynamics and the structural fluctuation of solvation sheath.

In this regard, the 2D infrared spectroscopy is an ideal tool for studying such ultrafast structural changes inside each solvation shell, because it is capable of tracking frequency changes of infrared probe (C=O stretch mode of DEC in the present case) induced by conformational transitions between different solvation structures. Indeed, at longer waiting time (\(T_w>2\) ps), one can clearly observe cross peaks (Fig. 3c). The positive cross peak on the upper-left corner results from the breaking of Li\(^+\)···DEC, that is, transition from Li-DEC to free DEC, which reduces the number of DEC solvent molecules in the first solvation shell (the backward reaction in Fig. 4a). Another positive cross peak appears on the lower-right corner, even though its magnitude is comparatively weak because of its destructive interference with the negative diagonal peak from free DEC. The positive and negative cross peaks in the lower-right corner represent the formation of Li-DEC complex or the increase of coordination number around Li\(^+\) ion (the forward reaction in Fig. 4a).

The Li-DEC coordination breaking and forming dynamics are manifested in the \(T_w\)-dependent cross peak intensity changes on the 2D infrared spectra. As mentioned earlier, there are three different DEC species with DECs in electrolyte solutions that are free DEC, DEC···Li\(^+\)···DEC\(_{n-1}\) and DEC···Li\(^+\)···PF\(_6\)\(_n\)···DEC\(_{n-2}\). However, DEC in DEC···Li\(^+\)···DEC\(_{n-1}\) and that in DEC···Li\(^+\)···PF\(_6\)\(_n\)···DEC\(_{n-2}\) are neither spectrally nor kinetically

Figure 3 | Dispersed infrared pump-probe and absorptive 2D infrared spectra. Time-and-frequency-resolved infrared pump-probe data of pure DEC liquid (a) and 1.0 M LiPF\(_6\) DEC solution (b) are plotted. (c) Four 2D infrared spectra of 1.0 M LiPF\(_6\) DEC solution at different waiting times are shown, where each 2D infrared spectrum was normalized to the positive peak maximum value. The x and y axes are the infrared frequency of the pump (excitation) and probe (emission) fields, respectively. In these contour plot, red (blue) colour represent positive (negative) sign.
molecules with different transition frequencies can produce cross peaks of which intensities increase in time $T_\nu$ (ref. 36). Unlike the case of chemical exchange-induced cross peaks, the increase rate of the upper-left cross peak intensity should differ from that of the lower-right cross peak intensity, because they are produced by up- and down-hill transitions, respectively.

Unfortunately, the lower-right positive cross peak in Fig. 4b overlaps with the negative diagonal peak from the free DEC, which obscures the time-dependent features of the cross peaks. We thus compared the time-dependent increase of the upper-left positive cross peak and that of the lower-right negative peak, assuming that the chemical exchange dynamics of DECs (either free DEC or Li–DEC complex) in their excited states are similar to those in their ground states. To rule out the possibility that the 2D infrared cross peaks originate from vibrational excitation transfers between free DEC and Li–DEC complex, we further carried out a series of concentration-dependent pump-probe measurements at four different LiPF$_6$ concentrations in the range from 0.2 to 1.0 M (detailed analysis on pump-probe results are found in Supplementary Note 3). A decrease in LiPF$_6$ concentration would significantly reduce the chance for a free DEC to encounter Li-complexed DEC molecules. This should result in an increase of vibrational lifetime of free DEC, if the vibrational energy transfer between free DEC and complex DEC occurs within a few picosecond timescale. However, as shown in Supplementary Fig. 6, the vibrational lifetime of free DEC is independent of LiPF$_6$ concentration. This confirms that the cross peaks observed in our 2D infrared spectra originate from ultrafast chemical exchange processes in the solvation sheath of Li$^+$ in DEC electrolyte solution.

Figure 4 | Equilibrium structure and kinetic analysis of 2D infrared spectrum. (a) A schematic picture on the chemical equilibrium between two solvation structures. The left consists of Li-complex with three bound DECs and one free DEC, whereas the right represents the tetra-coordinated Li-DEC complex. (b) The time-dependent diagonal and cross peak volumes obtained from the experimentally measured 2D infrared spectra are plotted. The fitted (solid) curves are also shown here.

2D infrared chemical exchange spectroscopy of mixed solvent. Despite that the ultrafast chemical exchange dynamics between free DEC and Li–DEC complex was observed in pure DEC solutions, typical LIB’s use mixed solvents instead. To elucidate the separate roles and solvation properties of co-solvents with DMC and PC and to investigate the possibility of DMC playing dual roles both as a solvent medium for mobile Li–PC(DMC) complexes and as a ligand participating in the first solvation shell around Li$^+$, we carried out 2D infrared chemical exchange measurements for LiPF$_6$ solutions in the more realistic DMC and PC mixed solvents, which are (i) 1.0 M LiPF$_6$ in DMC:PC (1:1 in volume percent) solution, (ii) 1.5 M LiPF$_6$ in DMC:PC (1.5:1) solution and (iii) 2.0 M LiPF$_6$ in DMC:PC (1.5:1) solution. Here it should be noted that we considered DMC:PC mixed solvents instead of DEC:PC mixed solvents because of the following reasons. First, since the C = O stretching vibrational lifetime of DMC is comparatively longer than that of DEC, we could observe measurable diagonal and cross peaks of free DMC and Li–DMC complex in our 2D infrared spectra at long waiting times. Second, we here show that the chemical exchanges observed in Li–DEC (single carbonate solvent) solution do occur in different mixed solvent systems too.

Although our experimental observations clearly show the ultrafast chemical exchange dynamics in lithium-ion solvation sheath, we still need to consider another possibility that the increasing cross peaks could result from vibrational excitation transfers between free DEC and Li–DEC complex. Indeed, the vibration energy transfer between donor and acceptor...
modes of Li–DMC and free DMC molecules, respectively, and the two in the higher-frequency region at around 1,780 and 1,800 cm$^{-1}$ originate from the C=O stretch modes of Li–PC and free PC molecules (the FTIR spectra of three different mixed solvent systems (i)–(iii) can be found in Supplementary Fig. 9).

As waiting time increases, the cross peak amplitude at $\omega=(\omega_1=\omega_2=1,725$ cm$^{-1}$ and $\omega_3=\omega_4=1,760$ cm$^{-1}$ increases (see the 2D infrared spectra inside the red boxes in Fig. 5), which clearly indicates the chemical exchange or configurational transition from Li–DMC to free DMC. In contrast, we could not find any cross peak features between Li–PC and free PC in our 2D infrared spectra in Fig. 5 and Supplementary Figs 10–12 within our experimental timescale. This suggests that the electrostatic interaction between PC and Li$^+$ in LIB electrolyte solutions is stronger than that between DMC (or DEC) and Li$^+$, which is consistent with the notion that the PC molecule with large dipole moment compared to that of DEC molecule tends to make a stable and rigid solvation sheath around a lithium cation in LIB electrolyte solutions.

To approximately examine the timescale of chemical exchange processes between Li–DMC complex and free DMC in 1.0 M LiPF$_6$/DMC:PC (1:1 in volume percent) solution, we fitted the 2D infrared peaks with a 2D Gaussian function (Supplementary Figs 10–12), which allow us to determine the volumes of the two positive diagonal peaks at 1,725 and 1,760 cm$^{-1}$ and the corresponding positive cross peak. Note that the two diagonal peak intensities of Li–DMC and free DMC at 1,725 cm$^{-1}$ and 1,760 cm$^{-1}$, which are denoted as $I_{\text{Li–DMC|2}}$ and $I_{\text{DMC|2}}$, respectively, are determined by a few factors that include (i) the transition dipole moments, $|\mu_{\text{Li–DMC}}|^2$ and $|\mu_{\text{DMC}}|^2$, (ii) vibrational lifetimes, (iii) rotational relaxation times and (iv) survival probabilities of the two species within the waiting time. The cross peak intensity, denoted as $I_{\text{cross}}$, between Li–DMC and free DMC is determined by (i) the transition dipole moment, $|\mu_{\text{Li–DMC}}|^2|\mu_{\text{DMC}}|^2$, (ii) vibrational lifetimes, (iii) rotational relaxation times and (iv) survival probabilities of the two species, that is, Li–DMC complex and free DMC, as well as the conditional probability due to chemical exchange process from Li–DMC to free DMC. Thus, this intensity ratio is in fact quantitatively a good measure of the chemical exchange rate.

From the 2D Gaussian fit to the 2D infrared spectra (at 10 ps) for 1.0 M LiPF$_6$/DEC solution shown in Fig. 3c, we found that the intensity ratio, which is a dimensionless quantity, is 0.67. Now, for the 2D infrared spectra (10 ps) for 1.0 M LiPF$_6$/DMC:PC (1:1) solution in Fig. 5, our estimated intensity ratio is 0.76. In the cases of the 1.5 M LiPF$_6$/DMC:PC (1.5:1) and 2.0 M LiPF$_6$/DMC:PC (1.5:1) solutions, the intensity ratios are found to be 0.69 and 0.71, respectively (Supplementary Figs 10–12). Our observation that the ratio of cross peak intensity to the geometric mean value of the two diagonal peak intensities does not strongly depend on the solvent composition suggests that the chemical exchange timescale in LiPF$_6$/DEC solution is quantitatively similar to that in mixed solvents consisting of DMC and PC.

**Molecular dynamics simulations.** To examine the structures and dynamics of Li$^+$ and PF$_6^-$ solvation shells and ion aggregates, we carried out classical MD simulations for 1.0 M LiPF$_6$/DEC solution using the force field parameters for PF$_6^-$ in refs 37,38 with the general Amber force field parameters for DEC. A representative snapshot structure of the solution is shown in Fig. 6a, where Li$^+$ is represented by a yellow sphere and PF$_6^-$ by an octahedron. At this high lithium salt concentration, ions form not just CIPs but also large aggregates with a polydisperse distribution. Figure 6b depicts the configuration of ions only. Interestingly, certain large-size aggregates adopt ion network structures where lithium cations and hexafluorophosphate anions form spatially extended multi-branch chains and even three-dimensional network-like structures. We have recently found that cations and anions in high salt

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**Figure 5 | Four representative two dimensional IR spectra of 1.0 M LiPF$_6$/DMC:PC (1:1 v/v%) solution at different waiting times.** Each 2D infrared spectrum was normalized to the positive peak maximum value.

**Figure 6 | Snapshot structure of 1.0 M LiPF$_6$/DEC solution obtained from MD simulation trajectories.** (a) Li$^+$ and PF$_6^-$ are represented by yellow spheres and grey octagons, respectively. (b) The ions in this LIB electrolyte solution form three-dimensionally extended ion network structures.
aqueous solutions of KSCN, NaClO₄ and NaBF₄ show a strong propensity to form a large-scale spatially extended networks that are tightly intertwined with water H-bonding networks. The fact that Li⁺ and PF₆⁻ ions form three-dimensional ion networks in LIB electrolyte solutions could be a key to understand detailed mechanism of Li⁺ ion mobility, since Li⁺ can jump from one to the other neighbouring anions in the same or different ion networks in the presence of non-zero gradient of electric potential. Currently, we are carrying out a series of spectral graph analyses of the MD trajectories obtained for a variety of LIB electrolyte solutions and will present the results elsewhere since it is beyond the scope of this report.

**Discussion**

Ultrafast fluxional changes of Li⁺ solvation sheath structures in LIB electrolyte solutions are for the first time observed here with employing chemical exchange 2D infrared spectroscopic method. Combining FTIR spectroscopy, infrared pump-probe measurement and quantum chemistry calculation results, we completely characterized the vibrational properties of free DEC and Li–DEC complexes in both pure DEC solvent and mixed solvents consisting of DEC and PC. Subsequent 2D infrared data analyses revealed the ultrafast nature of chemical exchange dynamics in the immediate vicinity of Li⁺ in DEC solution. Here one of the most difficult challenges was to prepare very thin solution samples because the infrared probes are the C=O stretch modes of DEC and PC molecules that are also solvents. Using RF magnetron sputtering technique, we could control the thickness of coated SiO₂ layer on CaF₂ window for linear and nonlinear infrared spectroscopic studies. The picosecond Li–DEC complex making and breaking dynamics in both pure DEC solution and DMC:PC mixed solution suggest that the macroscopic Li transport during both charging and discharging processes might be interrelated to microscopic solvation fluctuation and discrete transitions between different solvation states of Li⁺ in LIB electrolyte solutions. We anticipate that the present ultrafast chemical exchange dynamics should also play an important role in solvation and de-solvation of Li⁺ at the solid-electrolyte interface on electrodes because of the following reason. If the rigidity of each Li⁺ solvation sheath is very high, any de-solvation of Li⁺ on the anode surface may not be efficient, which potentially restricts the function of LIB due to lowering the rate of lithium deposition on the anode. On the other hand, if the Li⁺ solvation sheath is not sufficiently stable due to a low dissolving power of solvent, the rate of Li⁺ release from the cathode could be slow. Therefore, it is naturally expected that ideal LIB solvent molecules should interact with Li⁺ with a properly balanced strength. Overall, the unexpected ultrafast fluxional solvation processes around Li⁺ in the LIB electrolyte solution is turned out to be prerequisite for optimally achieving the functional role of carbonate solvent in LIB. We anticipate that a further 2D infrared investigation on the Li-solvent fluctuation dynamics during both charging and discharging processes would provide critical information on the fluctuation-dissipation relationship between equilibrium macroscopic solvent fluctuation and non-equilibrium macroscopic charge transport phenomenon.

In the present work, the carbonyl stretch mode of DEC was used as the infrared probe for monitoring any local change and dynamics of Li-solvation sheath. Since the infrared chromophore, DEC, is also solvent, the transmittance of infrared beam is very low so that an infrared sample holder with extremely narrow path length was needed. Previously, to overcome a similar difficulty, Cowan et al. used thin sample cell consisting of 2 mm × 2 mm window of 800 nm-thick Si₃N₄ for 2D infrared study of pure water, where the effective thickness of water flow channel was controlled to be 500 nm (ref. 42). Here to make an infrared sample cell and holder for the commercially available FT-IR spectrometer and subsequent 2D infrared measurements, we used SiO₂ to coat on the 25 mm diameter CaF₂ window (Fig. 1a). Here SiO₂ was selected because it is commercially non-reactive and also it is strongly adhesive to CaF₂. RF (radio frequency) magnetron sputtering system (Supplementary Note 1) was used with three-inch diameter high-purity SiO₂ target. The thickness of sputtered SiO₂ layer on CaF₂ surface was verified by scanning electron microscopy and used the linear relationship between the sputtering time and the thickness of SiO₂ layer (Fig. 1b). In the infrared pump-probe and 2D infrared measurements, to minimize the re-absorption of the generated third-order infrared signal electric field, that is, self-attenuation problem, the absorbance of DEC carbonyl stretching mode was adjusted to be less than 0.4, which was achieved by using 800 nm path length. All the infrared spectra were measured on Jasco 6300 FV spectrometer with a circulating water bath for temperature control. This spectrometer is equipped with a DLATGS detector with Peltier element in the absorbance mode range from 450 to 7,800 cm⁻¹ with frequency resolution of 0.07 cm⁻¹. During the measurements, N₂ gas is purged to suppress any vapour noise and to minimize undesired water condensation on the CaF₂ window.

### Polarization selective infrared pump-probe and 2D infrared spectroscopy

Experimental details of pump-probe and 2D infrared spectroscopy employed in the present study have been described elsewhere. Briefly, pump pulses of duration of τ = 100 fs and 1.0 mJ were generated from a Ti:sapphire oscillator and retenerative amplifier operating at 1 KHz, which are then used to produce near infrared pulses at ~1.4 and ~1.9 μm. With AgGaS nonlinear optical crystal, the signal and idler pulses are mixed to generate mid-infrared pulses with ~8 μJ per pulse and ~100 fs duration centred at 2,070 cm⁻¹. For polarization-selective infrared pump-probe and 2D infrared measurements, grid polarizers were used to set the polarization directions of the pump and probe beams to be 45° and 0° before the sample. The probe beam was resolved after the sample to either +45° (parallel) or -45° (perpendicular) by a linear polarizer on a computer-controlled rotator, sent through a polarization analyzer fixed to 0° into the spectrometer, and detected by a 32-element array detector.

### MD simulation method

Molecular dynamic (MD) simulations were performed for LiPF₆-DEC solutions at 1.0 M LiPF₆ concentration, where 378 LiPF₆ molecules were dissolved in 1,000 DEC molecules. The force field parameters developed by Lopes and Padua and the general Amber force field parameters were used to describe PF₆⁻ ion and DEC molecule, respectively, where the corresponding restrained electrostatic potential charges were obtained by carrying out B3LYP/6-31G+G(3df,2pd) calculations. The particle mesh Ewald method was used for long-range electrostatic interaction and the cutoff distance for nonbonding interaction was set to be 10 Å. The solution system was first energy-minimized with the steepest descent method and the conjugate gradient method before running equilibrium ensemble simulation at 298 K. After initial equilibration simulation for 1 atm and 298 K was carried out for 2 ns to adjust the solution density. An additional 10 ns N, V and T ensemble simulation at 298 K was performed to make each LiPF₆-DEC solution system to reach its thermal equilibrium state. Finally, the production run was performed for 10 ns at constant N, V and T conditions, where the simulation time step was set to be 1 fs, and atomic coordinates were saved for every 1 ps.

### Data availability

The data that support the findings of this study are available from the authors on reasonable request, see author contributions for specific data sets.

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**Acknowledgements**

This work was supported by IBS-R023-D1. K.-K.L. thanks Dr Ho-Jung Sun (Kansan National University) for allowing us to use the sputtering system and for insightful discussion.

**Author contributions**

M.C., K.-K.L. and K.K. designed the research. K.-K.L., K.P., Y.N. and D.K. performed the research. M.C. and K.K. wrote the manuscript.

**Additional information**

Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

**Competing financial interests:** The authors declare no competing financial interests.

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**How to cite this article:** Lee, K.-K. et al. Ultrafast fluxional exchange dynamics in electrolyte solution sheath of lithium ion battery. *Nat. Commun.* 8, 14658 doi: 10.1038/ncomms14658 (2017).

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