ELECTROCHEMISTRY

Understanding the molecular mechanism of pulse current charging for stable lithium-metal batteries

Qi Li,1,* Shen Tan,1* Linlin Li,1,2 Yingying Lu,1,2† Yi He1†

High energy and safe electrochemical storage are critical components in multiple emerging fields of technologies. Rechargeable lithium-metal batteries are considered to be promising alternatives for current lithium-ion batteries, leading to as much as a 10-fold improvement in anode storage capacity (from 372 to 3860 mAh g\(^{-1}\)). One of the major challenges for commercializing lithium-metal batteries is the reliability and safety issue, which is often associated with uneven lithium electrodeposition (lithium dendrites) during the charging stage of the battery cycling process. We report that stable lithium-metal batteries can be achieved by simply charging cells with square-wave pulse current. We investigated the effects of charging period and frequency as well as the mechanisms that govern this process at the molecular level. Molecular simulations were performed to study the diffusion and the solvation structure of lithium cations (Li\(^{+}\)) in bulk electrolyte. The model predicts that loose association between cations and anions can enhance the transport of Li\(^{+}\) and eventually stabilize the lithium electrodeposition. We also performed galvanostatic measurements to evaluate the cycling behavior and cell lifetime under pulsed electric field and found that the cell lifetime can be more than doubled using certain pulse current waveforms. Both experimental and simulation results demonstrate that the effectiveness of pulse current charging on dendrite suppression can be optimized by choosing proper time- and frequency-dependent pulses. This work provides a molecular basis for understanding the mechanisms of pulse current charging to mitigating lithium dendrites and designing pulse current waveforms for stable lithium-metal batteries.

INTRODUCTION

Since the first commercial lithium-ion battery (LIB) appeared in the early 1990s, LIBs have been widely used in portable electronic devices and have recently been successfully applied in electric vehicles, unmanned aircraft, and large-scale deployment (1). However, the graphite anode of LIBs only has a theoretical specific capacity of 372 mAh g\(^{-1}\). This inherent limitation makes it unlikely to meet the growing demand for high-energy storage (2, 3). Li metal is thought to be one of the most promising alternative anodes because of its step-change improvement in capacity (3860 mAh g\(^{-1}\)) and its low redox potential (~3.04 V versus the standard hydrogen potential electrode) (2, 4). In addition, lithium-metal anode is an essential component for lithium-sulfur and lithium-air batteries, which are currently under intensive research for next-generation high-energy lithium batteries (5–7). Unfortunately, many challenges need to be overcome before Li metal batteries become a reliable technology, the most serious of which are the poor cycling stability and safety issues (8).

During a typical charge step of cell cycling in lithium-metal batteries, lithium cations gain electrons on the anode surface and become metallic lithium. The reductive electrodeposition is usually nonuniform and hard to control because of the complicated surface chemistry and morphology of the anode. This heterogeneous nature of the electrodeposition or so-called Li dendrite becomes more marked during the subsequent cycles because of the uneven charge distribution along the rough surface. The growth of lithium dendrites is known as the source of battery failure. On the other hand, the electrolyte and the lithium metal could react irreversibly and form a solid electrolyte interface (SEI) on the metal surface to passivate the highly active Li metal and reduce further consumption of active materials. The fragile SEI could be ruptured during cycling because of the huge volume expansion of the anode. The instability of an SEI is another reason for the propensity of lithium to deposit in dendritic structures (9, 10). Once the fresh Li encounters the electrolyte, chemical reactions happen again and form a new SEI, leading to irreversible consumption of Li and electrolyte. As a consequence, the coulombic efficiency of Li metal batteries decreases, and the capacity fades quickly. The sharp dendrites on the surface of Li metal anode could pierce the separator and cause short circuit, generating heat or even igniting the organic solvents (11). In addition, the huge volume change of Li metal anode leads to internal stress and interface fluctuation (12). These problems have hindered the practical applications of rechargeable Li-metal–based batteries over the past 40 years.

Many experimental studies have emerged to regulate the Li metal anodes and stabilize the electrochemical behavior of lithium-metal batteries (13–20). In addition, the formation of lithium dendrites has been monitored at various length scales. Researchers have reported and theoretically predicted the conditions of dendrite nucleation and propagation. The dendrite growth is understood to be driven by the diffusion and electromigration of both Li ion and the counter ion under electric driving force. During charging, lithium cations and their counter ions transport in opposite directions and cause ion concentration gradient. At high current densities, the situation for the anions is more marked because they cannot form electrodes. Eventually, anions pile up near the anode and deplete near the cathode. This phenomenon is called “space charge,” which determines the onset time of lithium dendrites (21–25).

To fundamentally understand the mechanisms of electrolyte transport properties and suppress or even eliminate dendrite growth from the nucleation step, we report on the relaxation of ion concentration gradient via pulse current charging instead of common constant current charging and the stabilization of lithium-metal batteries via simply charging cells with square-wave pulse current (Fig. 1A). Pulse current charging was first used for lead acid battery
to remove lead sulfate compound and extend battery lifetime (26). It was later used as an advanced charging technique for LIBs for the homogeneity of ion distribution between the two electrodes (27, 28). These reports showed that the pulse current charging could lower the impedance and temperature increment during the charging process and reduce side reactions compared to constant current charging. Only a few studies for the influence of pulse current charging in rechargeable lithium-metal batteries have been reported. To the best of our knowledge, the only experimental work for pulse current charging contains limited cycling information without appropriate simulation support (29). The underlying mechanism at the molecular level, which is essential for the rational design of pulse current charging, is also limited so far. Only two recent studies by Aryanfar et al. (30) and Mayers et al. (31) performed coarse-grained molecular simulations for lithium dendrite growth under pulse current charging. They both revealed a competition between the time scales for Li\(^+\) diffusion and reduction at the anode-SEI interface to understand the dendrite formation, and theoretically suggested that lithium dendrites could be inhibited by pulse charging with appropriate charging frequency.

To understand dendrite growth mechanism both theoretically and experimentally, we used rectangular cathodic pulses with various frequencies and rest time, and monitored their growth propensity by all-atom molecular dynamics (MD) simulations and voltage-time profiles. MD simulations were performed to examine the diffusivity of Li ions in an electrolyte under pulsed electric fields with different frequencies and strength. The solvation structures of Li ions were also considered to further understand the effectiveness of transport properties on dendrite growth. The model predicts that loose association between cations and anions can enhance the diffusion of Li\(^+\). We also evaluated the cycling behavior by galvanostatic cycling measurements and found that the cell lifetime can be more than doubled using certain pulse current waveforms. Both experimental and MD simulation results demonstrated that the effectiveness of pulse current charging can be optimized by choosing proper time- and frequency-dependent pulses. This work provides a molecular basis for understanding the mechanisms of pulse current charging for mitigating lithium dendrites and designing pulse current waveforms for stable performance in advanced lithium-metal batteries, such as high-energy lithium-sulfur and lithium-oxygen batteries.

**RESULTS**

**Diffusion of Li\(^+\) in pulsed electric fields**
Pulsed electric fields may affect both the diffusion and electrodeposition of Li\(^+\). Although it is still challenging to examine the electrodeposition of Li\(^+\) using classic all-atom MD at a large scale, it is, however, feasible to examine the diffusion of Li\(^+\) at the molecular level, which is the focus of this work. We abstract a local area in an electrolyte and analyze the microscopic motion of Li\(^+\) based on all-atom MD simulations. In all-atom models, the position and velocity of each individual atom are explicitly
represented in the simulations, which allow high-resolution analysis of simulation trajectories. The simulation system is constructed with 51 Li\(^{+}\), 51 TFSI\(^{-}\) [bis(trifluoromethanesulfonil)imide anion], and 600 PC (propylene carbonate) molecules, which gives a total of 8616 atoms and represents LiTFSI [lithium bis(trifluoromethanesulfonil)imide]-PC solution with a concentration of 1.0 M lithium salt. The system is in a cubic box with a dimension of 43.88 Å in length (x), width (y), and height (z) (Fig. 1B). The self-diffusion coefficient \( D \) of Li\(^{+}\) can be computed from the mean square displacement (MSD) of MD trajectories based on the Einstein relation

\[
D_{\text{MSD}} = \lim_{t \to \infty} \frac{\langle (\mathbf{r}(t) - \mathbf{r}(0))^2 \rangle}{6t}
\]

where \( \mathbf{r}(t) \) is the location of the center of mass of a specific ion at time \( t \); the brackets indicate the ensemble average. Figure 2 shows that the diffusion coefficients of Li\(^{+}\) in all cases are sensitive to applied electric fields. Pulsed electric fields may significantly enhance the diffusion of Li\(^{+}\), which is larger than that in a static electric field or in the control case, where no electric field is applied. The pulsed electric field with a \( T_{\text{on}}/T_{\text{off}} \) ratio of 1:3 \((T_{\text{on}}\) means the period of constant current charging or discharging, and \( T_{\text{off}} \) means the rest period followed by the constant current charging or discharging) is observed to offer the highest diffusion coefficient of Li\(^{+}\) among all the cases examined in this study. Moreover, in an electric field with a higher strength of \( 3 \times 10^8 \text{ V m}^{-1} \), a \( T_{\text{on}}/T_{\text{off}} \) ratio of 1:3 is also found to be the optimal condition for Li\(^{+}\) diffusion (Fig. S1A). This observation suggests that an innate nature of LiTFSI-PC solution may determine the optimal pulse current waveform for achieving the highest diffusivity of Li\(^{+}\), which is independent of the strength of the electric field within a certain range. For instance, in electric field with an even higher strength, such as \( 3 \times 10^8 \text{ V m}^{-1} \), the diffusion coefficient was observed to be dominated by the strength of electric field (Fig. S1B). This is consistent with a recent experimental study by Pei and co-workers (32), suggesting that high overpotentials are capable of effectively reducing the dendrite formation of lithium-metal batteries. Because the only intramolecular interactions of Li\(^{+}\) in the solutions are the ones with TFSI\(^{-}\) and PC, the solvation structures of Li\(^{+}\) and TFSI\(^{-}\) in PC solutions need to be examined for further insights.

It should be noted that the diffusion coefficients of Li\(^{+}\) obtained from this work are similar to those from other MD studies using the OPLS-AA (optimized potentials for liquid simulations for all-atom) force field (33, 34), which are typically about one order of magnitude lower than the experimental findings measured with the pulse gradient spin echo nuclear magnetic resonance (35). Despite the difference in value, the diffusion coefficients derived from simulations were understood to qualitatively agree with those revealed by experiments (33).

**Solvation structure of Li\(^{+}\) in PC solutions**

As shown in the simulation snapshot (Fig. 1, B and C), Li\(^{+}\) may associate with TFSI\(^{-}\) and PC, leading to certain solvation structures. The structures can be described with the radial distribution function (RDF), which shows how density of atom \( j \) varies as a function of distance from a reference atom \( i \) (in this case is Li\(^{+}\), and \( j \) could be either a PC molecule or TFSI\(^{-}\))

\[
g_{ij}(r) = \frac{\langle N_{ij}(r)V(r) \rangle}{\rho_j}
\]

where \( N_{ij}(r) \) is the ensemble averaged number of atoms \( j \) in a spherical shell of volume \( V(r) \) at a distance \( r \) from atom \( i \), and \( \rho_j \) is the bulk density of atom \( j \). The coordination numbers of the PC solvent molecules surrounding the solvation shell of the ions are calculated from the following integral

\[
n(R) = 4\pi\rho_j \int_0^R r^2 g_{ij}(r)dr
\]

where \( R \) is the minimum after the first peak in the RDF \( g_{ij}(r) \). The different coordination numbers may affect the dynamic properties of the Li\(^{+}\), such as their diffusivity (36).

Figure 3 (A and B) shows that the first peak in \( g(r) \) for Li\(^{+}\) appears at around 2.1 Å for both carbonyl oxygen atoms in PC and oxygen atoms in TFSI\(^{-}\). These results agree with previous reported studies of LiPF\(_6\) in PC (37) and LiTFSI-acetamide systems (36). It also indicates that Li\(^{+}\) has strong solvation structures with PC molecules. This association is more obvious between Li\(^{+}\) and TFSI\(^{-}\) ion pairs because the peak intensity in Fig. 3B is stronger than that in Fig. 3A.

As shown in Fig. 3 (C and D), the coordination number of oxygen atoms around Li\(^{+}\) is approximately six, among which four are from PC and one or two are from TFSI\(^{-}\). Similar results have been previously reported with Raman intensity experiments (38) and classical MD simulations (39–41) that, when a static electric field is applied, the coordination number of TFSI\(^{-}\) around Li\(^{+}\) gets smaller. In addition, Fig. 3 (C and D) demonstrates that both coordination numbers of cation-solvent \([\text{Li}\(^{+}\)-O(carbonyl in PC)]\) and cation-anion \([\text{Li}\(^{+}\)-O(TFSI\(^{-}\)])\) can be affected by applying pulsed electric fields. When a pulsed electric field with a \( T_{\text{on}}/T_{\text{off}} \) ratio of 1:3 is used, Li\(^{+}\) has a relatively larger coordination...
number of PC and a smaller coordination number of TFSI$^-$ compared with the control case where no electric field is applied, leading to a larger diffusivity of Li$^+$ (Fig. 2). It is demonstrated that the pulse current charging affects the structural properties of the electrolyte, which decreases the coordination number of TFSI$^-$ around Li$^+$ ions. That is, the number of TFSI$^-$ anions around Li$^+$ is reduced, and the repulsion effect between Li$^+$ and Li$^+$ is promoted. As a result, the Li$^+$ can transport easily, and the diffusion coefficient of Li$^+$ is increased. This enhanced diffusion would reduce the cluster behavior of Li$^+$ in the bulk electrolyte, promoting uniform distribution of ions and reducing ion concentration gradient.

Cycling behavior

Galvanostatic cycling measurements were used to mimic the real condition of battery cycling. Cells were cycled at 3 mA cm$^{-2}$ for 20 min in each half-cycle. Same symmetrical cells were cycled with pulse current at various $T_{on}$ durations and $T_{on}/T_{off}$ ratios ($T_{on}/T_{off} = 1:0.5, 1:1, 1:2, 1:3, 1:5$, and $1:10$; $T_{on} = 1, 5,$ and $10$ s). All pulse waveforms are galvanostatic square-wave pulses. The current density and total capacity passed through in each half-cycle were kept the same for cells with and without pulse current. To obtain a quantitative measure of the relative performance of the two cells, we arbitrarily defined the cell lifetime as the time before peak voltage amplitude increases to 1 V. The voltage becomes unstable, and postmortem scanning electron microscopy (SEM) analysis of the lithium electrode from cells interrupted at 1 V also indicates that the voltage change coincides with the appearance of lithium dendrites on the electrode surface. Figure 4A reports the voltage-time profiles for cells without pulse current at 3 mA cm$^{-2}$, and it shows a gradual increase in hysteresis (overpotential between Li plating and stripping) as the number of cycles increases. It is interesting to notice that a sudden increase in voltage appears at the 20th cycle, and the voltage profile becomes unstable in the following cycles. The sudden increase is known as an indication for cell instability that the internal impedance changes as the lithium dendrite shoots out from the substrate. Figure 4B shows the voltage profile for cells with pulse current ($T_{on}/T_{off} = 1:5; T_{on} = 1$ s) at the same current density. The cell can be cycled for at least 54 cycles without sudden increase or drop in voltage. The cutoff lifetime for cells with pulse current ($T_{on}/T_{off} = 1:5; T_{on} = 1$ s) at the same current density. The cell can be cycled for at least 54 cycles without sudden increase or drop in voltage. The cutoff lifetime for cells with pulse current is 48 cycles (192 hours), which is 2.4 times higher than that for cells with constant current cycling. Their detailed comparison in voltage at the 11th cycle can be found in Fig. 4 (C and D), which shows enlarged views of the rectangular areas corresponding to Fig. 4 (A and B, respectively). Cells under pulse current cycling exhibit a stable voltage response with small increase in hysteresis (about 0.8 V at the 11th cycle).

![Fig. 3. The solvation structures of Li$^+$ in PC solutions. RDFs of (A) cation-solvent [Li$^+$-O(carbonyl in PC)] and (B) cation-anion [Li$^+$-O(TFSI$^-$)] and the coordination numbers of (C) cation-solvent [Li$^+$-O(carbonyl in PC)] and (D) cation-anion [Li$^+$-O(TFSI$^-$)] under conditions of no electric field, static electric field, and pulsed electric fields with a strength of $3 \times 10^5$ V m$^{-1}$. The $T_{on}/T_{off}$ ratios of pulsed electric fields are 1:0.5, 1:1, 1:2, 1:3, 1:4, 1:5, and 1:10. For convenience, the static electric field is shown as the one with a $T_{on}/T_{off}$ ratio of 1:0. Pulse current charging can reduce the coordination number of TFSI$^-$ around Li$^+$ ions.](http://advances.sciencemag.org/)

Li et al., Sci. Adv. 2017;3:e1701246 21 July 2017 4 of 9

Downloaded from http://advances.sciencemag.org/ on June 27, 2021
compared to cells cycled without pulse current (about 1 V at the 11th cycle). Figure 4 (E and F) shows the details about the voltage response, current density, and $T_{on}/T_{off}$ settings in Fig. 4 (C and D). Generally, when cycled with pulse current, the cell shows lower voltage hysteresis and great improvement in cycling stability and cell lifetime.

To demonstrate the growth of lithium dendrites after galvanostatic cycling, we used SEM to analyze the surface and cross-section morphologies of Li-metal electrodes before cycling and after being cycled with constant current or pulse current. Symmetric Li cells were cycled at a fixed current density of 3 mA cm$^{-2}$ with a string/plating capacity of 1 mAh cm$^{-2}$ for each half-cycle. Then, the cells were disassembled in an Ar-protected glove box, and the Li-metal electrodes were washed by PC and dried rigorously before SEM test. As shown in fig. S2A, the surface of a pristine Li-metal electrode is uniform in general except for some small manufactural defects. After being cycled eight times, lithium foil harvested from cells without using pulse current shows rough, fibrous, and irregular jaggy-shaped structures, as displayed in fig. S2B. These nonuniform and sharp structures of lithium could eventually become Li dendrites, penetrate the separator, and cause cell short circuit. Figure S2C displays the surface morphology of the Li electrode obtained from pulse current cycling and shows uniform microstructure compared with that of the Li-metal anode with constant current cycling. We also tracked the thickness of the Li electrode from cross-section SEM analysis to study the electrode volume change upon cycling. The thickness of the pristine Li electrode is around 420 µm. After eight cycles, the electrode thickness increases by 30 and 84 µm for cells with and without pulse current cycling, respectively. The smaller increment in electrode thickness indicates that the electrode structure is more compact using pulse current cycling and the electrode volume expansion is significantly reduced. The results are consistent with the findings in galvanostatic cycling measurements and suggest that fewer lithium dendritic structures form in
cells with pulse current cycling. Cycling with pulsed current can increase the lithium coulombic efficiency and ultimately improve the cycling stability and cell lifetime.

To find the optimal conditions for dendrite suppression under pulse current cycling, we provide additional details about the effectiveness of pulse current cycling and statistically calculated the cell lifetime using galvanostatic measurements in Li/Li cells. The cell lifetime ($T_c$) is defined as the time when the amplitude of voltage exceeds 1 V. A high current density of 3 mA cm$^{-2}$ was implemented for practical rate capability. Figure 5 summarizes $T_c$ as a function of $T_{on}$ and frequency. The voltage of cells with constant current cycling exceeds 1 V at about the 20th cycle, whereas all cells with pulse current cycling show improvement in cell lifetime. It can also be seen in general that the cell lifetime increases with the increment of $T_{off}$ at fixed $T_{on}$ and increases when reducing the period of $T_{on}$ at a fixed ratio of $T_{on}/T_{off}$. The optimal enhancement in cell lifetime appears when $T_{on}/T_{off} = 1:5$, indicating that more rest time between the two periods of charging is not necessary for longer cell lifetime. The results also show that the cell lifetime depends on the pulse frequency and high-frequency pulse cycling is more efficient in promoting uniform Li electrodeposition and extending the cell lifetime.

**DISCUSSION**

Pulse current charging was demonstrated to be a promising method for stabilizing lithium-metal batteries compared with common constant current charging. We have used MD simulations to study the diffusion of Li$^+$ at the molecular level. The results suggest the existence of an optimal ratio, $T_{on}/T_{off}$, for achieving the highest diffusion coefficient of Li$^+$. We also analyzed the solvation of Li ions and revealed that a lower coordination number of TFSI$^-$ near Li$^+$ promotes the diffusivity of Li$^+$. In pulsed electric fields, Li$^+$ has an increased coordination number of PC and a reduced coordination number of TFSI$^-$ compared to the control case where no electric field is applied. The simulation results indicate that the transport of Li$^+$ in the bulk electrolytes and the underlying ion concentration gradient play a critical role in reducing dendrite formation and promoting stable cell cycling. A weaker association between Li cation and its counter ion is preferred. Electrochemical cycling performance was also evaluated for further evidence of the effectiveness of pulse current charging on cell stability. Consistent with the findings in MD simulations, all cells cycled with pulse current show enhanced cycling stability and lower hysteresis. Results from SEM analysis provide support of the cycling measurements and conclude that Li dendrites could be effectively hindered.

---

**Fig. 5. Cell lifetime at various frequencies with fixed $T_{on}$.** (A) Cell lifetime ($T_c$) of Li/Li symmetrical cells cycled with pulse current at various $T_{off}$ with fixed $T_{on} = 1$ s. (B) Cell lifetime ($T_c$) of Li/Li symmetrical cells cycled with pulse current at various $T_{off}$ with fixed $T_{on} = 5$ s. (C) Cell lifetime ($T_c$) of Li/Li symmetrical cells cycled with pulse current at various $T_{off}$ with fixed $T_{on} = 10$ s. The cell lifetime is calculated when the amplitude of the voltage exceeds 1 V. Fourteen cells under each cycling condition are tested, and their cell lifetimes are displayed in filled square with different colors. The red column shows the average lifetime from the 14 points. (D) Average cell lifetime as a function of $T_{on}/T_{off}$ ratios at fixed $T_{on}$ of 1, 5, or 10 s.
with pulse current and the volume expansion of the Li electrode can also be reduced. We found from both simulation and experimental results that the ratio of $T_{\text{on}}/T_{\text{off}}$ is critical for stabilizing the cycling behavior, and pulse charging with $T_{\text{on}}/T_{\text{off}} = 1.5$ or $T_{\text{on}}/T_{\text{off}} = 1.3$ is an optimal condition examined in this study. Considering the difference in time scales and other factors, such as ion migration and electrodeposition of Li ions on electrode surface, it is reasonable that some results are unable to have a one-to-one correspondence between simulations and experiments. To further enhance the benefits brought by the pulse current charging, future studies could include finding an anion with a lower coordination number around Li$^+$ to promote the Li diffusivity and eventually improve cycling stability and extend the lifetime of high-energy Li-metal batteries.

**MATERIALS AND METHODS**

**Simulation details**

The simulation system was constructed with 51 Li$^+$, 51 TFSI$^-$, and 600 PC molecules, which give a total of 8616 atoms and represent the LiTFSI solution with a concentration of 1.0 M lithium salt. The system is in a cubic box with a dimension of 43.88 Å in length ($x$), width ($y$), and height ($z$). The whole system, including Li$^+$, TFSI$^-$, and PC, were initially energy-minimized for 10,000 steps using the conjugate gradient algorithm to remove abnormally close contacts between molecules. Following the minimization, the system was heated from 50 to 300 K using a short MD run of 1 ns or 1.0 fs time steps and 50 K increments for every 100 ps until the system reached 300 K. After heating, an additional 1 ns or 1.0 fs MD steps were run to allow further isothermal equilibration of Li$^+$ and TFSI$^-$ in PC. The velocity Verlet method was used for the integration of the Newtonian equations in NPT (a simulation system that consists of a constant number of atoms at a constant pressure, in which the temperature of the system is controlled by using a thermostat) ensemble for equilibration. A pressure of 1 atm was used in all simulation cases. Cutoff of 1.2 and 1.0 nm was used for the Lennard-Jones and electrostatic terms, respectively. A neighbor searching was made up to this same distance from the central ion and was updated every five simulation steps. The electrostatic interactions beyond 1.0 nm were accounted for by the computationally efficient K-space method based on particle-particle particle-mesh (PPPM). It is important to use the PPPM method in the case of ionic systems, because electrostatic energy beyond the cutoff usually contributes a lot to the total electrostatic energy (42). The final configuration of the system has a density of ~1.329 g cm$^{-3}$, which agrees with the experimentally determined density of PC solution of LiTFSI (1.3205 g cm$^{-3}$, 1.11 M) (43).

For production runs, a series of 12-ns simulations at 300 K were carried out in the NVT (a simulation system that consists of a constant number of atoms at a constant volume, in which the temperature of the system is controlled by using a thermostat) ensemble using a Nosé-Hoover thermostat with a coupling constant of 0.1 ps. A pulsed electric field was set to be 0.01 ns. The on duration was set in a way so that the on/off value was 0.5, 1, 2, 3, 4, 5, and 10, respectively. Moreover, a simulation case with no applied electric fields was studied as a control. The simulation results were obtained by analyzing the last 4 ns trajectories of production runs.

The OPLS-AA force field (44, 45) was used. This force field has been widely used for biomolecules (46), hydrocarbon molecules (47, 48), and ionic liquids (49, 50). The basic functional form of potential energy in the OPLS-AA force field is defined as follows:

$$E = \sum_{i,j} K_{ij} (r_i - r_{ij})^2 + \sum_{i} K_{ii} (\theta_i - \theta_{ii})^2 + \sum_{i} \left[ \frac{1}{2} V_{1,ii}(1 + \cos(q_{i,j})) + \frac{1}{2} \frac{V_{2,ii}(1 - \cos(2q_{i,j}))}{1 - \cos(2q_{i,j})} \right] + \sum_{i,j} \frac{q_i q_j e^2}{r_{ij}} + 4\epsilon_{ij} \left[ \left( \frac{\sigma_{ij}}{r_{ij}} \right)^{12} - \left( \frac{\sigma_{ij}}{r_{ij}} \right)^6 \right]$$

The total energy $E$ of the system is evaluated as the sum of individual energies for the harmonic bond stretching and angle bending terms, a Fourier series for torsional energetics, and Coulomb and 12-6 Lennard-Jones terms for the nonbonded interactions. $K_{ij}$, $\theta_i$, $r_{ij}$, and $q_{i,j}$ represent the reference values. The Fourier coefficients $V$ and the partial atomic charges $q$ were fixed on each atom center. $i$ and $j$ represent all pairs of atoms ($i < j$), $r_{ij}$ is the distance between atoms $i$ and $j$, $\sigma_{ij}$ and $\epsilon_{ij}$ represent the Lennard-Jones radii and potential well depths, respectively, which are obtained from parameters for each type of atom by using geometric combination rules $\epsilon_{ij} = (\epsilon_i \epsilon_j)^{1/2}$ and $\sigma_{ij} = (\sigma_i \sigma_j)^{1/2}$ (48). The Lennard-Jones parameters and partial charges for Li$^+$ and PC were taken from the OPLS-AA parameterization (37, 49, 51). The bonded stretching, angle bending, dihedral torsion, and nonbonded parameters of TFSI$^-$ anions were obtained from Canongia Lopes and Pádua (49). The partial charge of atoms in TFSI$^-$ anions was calculated with the following protocol. First, geometry optimizations were carried out at the Hartree-Fock (HF) level by using a moderately sized polarized basis set at the 6-31G* level. The partial charges were computed by fitting the molecular electrostatic potential at the atomic centers. The effects of electron correlation and basis set extension of HF structures were considered by performing single-point calculations with second-order Moller-Plesset perturbation method MP2/cc-PVdz and the correlation consistent polarized valence cc-pVTZ(-f) basis set. This MP2/cc-pVTZ(-f) //HF/6-31G(d) method is a common practice for OPLS-AA parameterization (50, 52, 53). The calculated restrained electrostatic potential charges for TFSI$^-$ agree with other studies (49).

All MD simulations were performed with the LAMMPS (Large-scale Atomic/Molecular Massively Parallel Simulator) package on a 20-node Linux cluster (54). All quantum chemistry calculations were carried out with the Gaussian 09 package (55).

**Electrochemical measurements**

CR2032 coin cells were assembled in an argon-filled glove box. Bare Li metal was used as the reference and counter electrodes, and 1 M LiTFSI in PC was used as the electrolyte. Symmetrical lithium cells were galvanostatically cycled at 3 mA cm$^{-2}$ for 20 min in each half-cycle. The same symmetrical cells were cycled with pulse current charging and discharging at various $T_{\text{on}}$ and $T_{\text{off}}/T_{\text{off}}$ ratios ($T_{\text{on}}/T_{\text{off}} = 1.0, 1.5, 1.1, 1.2, 1.3, 1.5, 1.0$; $T_{\text{on}} = 1, 5$, and 10 s). $T_{\text{on}}$ means the period of constant current charging or discharging, and $T_{\text{off}}$ means the rest...
period followed by the constant current charging or discharging. The current density and total capacity in each half-cycle were kept the same for cells with and without using pulse current. Neware electrochemical testing system (CT4008) and Solartron Analytical Electrochemical Workstation were used for the electrochemical measurements.

**SEM analysis**

After galvanostatic cycling, the cells were carefully disassembled in a dry glove box filled with argon to obtain Li-metal electrodes. Then, Li-metal electrodes were gently washed with PC three times to remove residual lithium salts and fully dried under vacuum. To access the cross-section images of the Li-metal electrodes, the obtained lithium foils were cut with a sharp surgical knife blade and placed on a vertical sample holder. SEM analysis was conducted on a field-emission SEM (Hitachi SU8000 FE-SEM) at 5 kV.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/supplementary-materials a field-emission SEM (Hitachi SU8000 FE-SEM) at 5 kV.
Experimental and computational investigations. *Phys. Chem. Chem. Phys.* **18**, 9504–9513 (2016).

42. I. V. Vorobylova, V. V. Chaban, Atomistic force field for pyridinium-based ionic liquids: Reliable transport properties. *J. Phys. Chem. B* **118**, 10716–10724 (2014).

43. Z. Wang, W. Gao, X. Huang, Y. Mo, L. Chen, Spectroscopic studies on interactions and microstructures in propylene carbonate—LiTFSI electrolytes. *J. Raman Spectrosc.* **32**, 900–905 (2001).

44. W. L. Jorgensen, D. S. Maxwell, J. Tirado-Rives, Development and testing of the OPLS all-atom force field on conformational energetics and properties of organic liquids. *J. Am. Chem. Soc.* **118**, 11225–11236 (1996).

45. W. Damm, A. Frontera, J. Tirado-Rives, W. L. Jorgensen, OPLS all-atom force field for carbohydrates. *J. Comput. Chem.* **18**, 1955–1970 (1997).

46. Z. Xu, H. H. Luo, D. P. Tieleman, Modifying the OPLS-AA force field to improve hydration free energies for several amino acid side chains using new atomic charges and an off-plane charge model for aromatic residues. *J. Comput. Chem.* **28**, 689–697 (2007).

47. S. W. I. Siu, K. Pluhackova, R. Bockmann, Optimization of the OPLS-AA force field for long hydrocarbons. *J. Chem. Theory Comput.* **8**, 1459–1470 (2012).

48. W. L. Jorgensen, Optimized internuclear potential functions for liquid alcohols. *J. Phys. Chem.* **90**, 1276–1284 (1986).

49. J. N. Canongia Lopes, A. A. H. Pádua, Molecular force field for ionic liquids composed of triflate or bistriflylimide anions. *J. Phys. Chem. B* **108**, 16893–16898 (2004).

50. S. V. Sambasivarao, O. Acevedo, Development of OPLS-AA force field parameters for 68 unique ionic liquids. *J. Chem. Theory Comput.* **5**, 1038–1050 (2009).

51. K. P. Jensen, W. L. Jorgensen, Halide, ammonium, and alkali metal ion parameters for modeling aqueous solutions. *J. Chem. Theory Comput.* **2**, 1499–1509 (2006).

52. E. K. Watkins, W. L. Jorgensen, Perfluorooalkanes: Conformational analysis and liquid-state properties from ab initio and Monte Carlo calculations. *J. Phys. Chem. A* **105**, 4118–4125 (2001).

53. G. A. Kaminski, R. A. Friesner, J. Tirado-Rives, W. L. Jorgensen, Evaluation and reparametrization of the OPLS-AA force field for proteins via comparison with accurate quantum chemical calculations on peptides. *J. Phys. Chem. B* **105**, 6474–6487 (2001).

54. S. Plimpton, Fast parallel algorithms for short-range molecular dynamics. *J. Comput. Phys.* **117**, 1–19 (1995).

55. M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, B. Mennucci, G. A. Petersson, H. Nakatsuji, M. Caricato, X. Li, H. P. Hratchian, A. F. Izmaylov, J. Bloino, G. Zheng, J. L. Sonnenberg, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, J. A. Montgomery, Jr., J. E. Peralta, F. Ogliaro, M. Bearpark, J. J. Heyd, E. Brothers, K. N. Kudin, V. N. Staroverov, R. Kobayashi, J. Normand, K. Raghavachari, A. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, N. Rega, J. M. Millam, M. Klene, J. E. Knox, J. B. Cross, V. Bakken, C. Adamo, J. Jaramillo, R. Gomperts, R. E. Stratmann, O. Yazyev, A. J. Austin, R. Cammi, C. Pomelli, J. W. Ochterski, R. L. Martin, K. Morokuma, V. G. Zakrzewski, G. A. Voth, P. Salvador, J. J. Dannenberg, S. Dapprich, A. D. Daniels, O. Farkas, J. B. Foresman, J. V. Ortiz, J. Cioslowski, D. J. Fox, *Official Gaussian 09 Literature Citation* (Gaussian Inc., 2009).

Acknowledgments: We thank F. Chen (Department of Chemistry, Zhejiang University) for SEM analysis. Funding: This work was supported by the Natural Science Foundation of China (grants 21676242, 21476191, 91434110, and 21676245) and Ministry of Science and Technology of the People’s Republic of China (grant 2016YFA0202900). Author contributions: Y.L. and Y.H. designed the experiment, conceived the project, and wrote the manuscript. Q.L. performed the experimental work and analyzed the results with the help of Y.L. and S.T. S.T. conducted the simulation work and analyzed the results with the help of Y.H. and L.L. Competing interests: The authors declare that they have no competing interests. Data and materials availability: All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data related to this paper may be requested from the authors.

Submitted 19 April 2017
Accepted 13 June 2017
Published 21 July 2017
10.1126/sciadv.1701246

Citation: Q. Li, S. Tan, L. Li, Y. Lu, Y. He, Understanding the molecular mechanism of pulse current charging for stable lithium-metal batteries. *Sci. Adv.* **3**, e1701246 (2017).
Understanding the molecular mechanism of pulse current charging for stable lithium-metal batteries
Qi Li, Shen Tan, Linlin Li, Yingying Lu and Yi He

Sci Adv 3 (7), e1701246,
DOI: 10.1126/sciadv.1701246

ARTICLE TOOLS http://advances.sciencemag.org/content/3/7/e1701246
SUPPLEMENTARY MATERIALS http://advances.sciencemag.org/content/suppl/2017/07/17/3.7.e1701246.DC1
PERMISSIONS http://www.sciencemag.org/help/reprints-and-permissions

Use of this article is subject to the Terms of Service

Science Advances (ISSN 2375-2548) is published by the American Association for the Advancement of Science, 1200 New York Avenue NW, Washington, DC 20005. The title Science Advances is a registered trademark of AAAS.

Copyright © 2017 The Authors, some rights reserved; exclusive licensee American Association for the Advancement of Science. No claim to original U.S. Government Works. Distributed under a Creative Commons Attribution NonCommercial License 4.0 (CC BY-NC).