

Insight into SEI Growth in Li-Ion Batteries using Molecular Dynamics and Accelerated Chemical Reactions[†]

Lorena Alzate-Vargas,^{*,‡} Samuel M. Blau,[¶] Evan Walter Clark Spotte-Smith,^{§,||}
Srikanth Allu,[‡] Kristin A. Persson,^{§,⊥} and Jean-Luc Fattebert[‡]

[‡]*Computational Sciences and Engineering Division, Oak Ridge National Laboratory, Oak Ridge, TN, USA*

[¶]*Energy Technologies Area, Lawrence Berkeley National Laboratory, Berkeley, CA, USA*

[§]*Department of Materials Science and Engineering, University of California, Berkeley, CA, USA*

^{||}*Materials Science Division, Lawrence Berkeley National Laboratory, Berkeley, CA, USA*

[⊥]*Molecular Foundry, Lawrence Berkeley National Laboratory, Berkeley, CA, USA*

E-mail: alzatevargll@ornl.gov

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Abstract

The performance of lithium-ion batteries (LIB) using organic electrolytes strongly depends on the formation of a stable solid electrolyte interphase (SEI) film. Elucidating the dynamic evolution and spatial composition of the SEI can be very useful to study the stability of SEI components and help optimize the formation cycles of LIB. We propose a classical molecular dynamics simulation protocol for predicting the first stages of SEI formation using a reaction method involving the decomposition of EC and LiPF₆ molecules in the electrolyte. We accelerate the formation of SEI components near the anode surface by increasing the probability of reactions, implemented through a geometry matching scheme, followed by a force-field reconfiguration. We observe the formation of gases (C₂H₄), inorganic (Li₂CO₃ and LiF) and organic (LEDC) components. This protocol shows promise to be able to evaluate the effects of varying electrolyte compositions and additives on SEI layer structure and composition.

1 Introduction

Lithium-ion batteries (LIBs) are key for transforming the energy infrastructure and transportation towards the storage of renewable electricity and enabling long range electrical vehicles. LIBs have been widely used in an extensive range of applications from small portable devices to large-scale energy storage systems.^{1,2} The solid electrolyte interphase (SEI) is one of the most critical components due to its influence on the battery performance i.e., capacity degradation, safety, calendar life, and cycle life.^{3{5}} At the same time, it is the least understood components of lithium-ion batteries. The SEI usually consists of reduction products formed through reactions between electrode and electrolytes due to the electron leakage from the anode.⁶ The SEI layer can be viewed as a multi-layered structure — an inorganic inner layer near the electrode that allows lithium ion transport and an organic outer layer permeable to electrolyte solvent molecules.⁷ Numerous experimental techniques have been developed to characterize the film composition and contribute to an understanding of the

SEI formation.^{8,9} Nevertheless, fully characterizing the SEI remains a challenge. The development of computational models that can contribute to its understanding have become more popular during the last years.^{10,11} These techniques can unveil the details of SEI formation at a molecular level, providing insights that cannot be obtained from experiment alone.

Atomistic-level studies using first principles calculations have been able to predict the reductive reaction mechanism of the solvent ethylene carbonate (EC) and reaction pathways to multiple SEI components, such as LEDC.^{12{14} Similarly, first principles molecular dynamics (FPMD) studies have focused on the reductive decomposition mechanisms of electrolyte solvent molecules on different electrodes.^{15{20} Although FPMD can predict chemistry — it includes the atom’s electronic structure, and can properly describe atomic bonds and forces, allowing bond breaking and bond formation — it is restricted to small systems and time resolution remains limited on the 10s of picoseconds in practice. That drastically reduces the number of chemical reactions one can observe and would provide limited information about the SEI growth mechanism.

Classical molecular dynamics (MD) have become more popular to investigate the behavior of LIBs, since they allow simulations with bigger systems on nanosecond timescales. In general, such MD simulations do not allow reactive events, limiting their utility for studies of the SEI. Still, they have enabled simulations that capture the transport and structural properties of the electrolyte and SEI^{21{25} and lead to an accurate description of the properties of the main SEI components.^{26,27} Some MD simulations have modeled electrode-electrolyte structural changes under electric field, ignoring any chemical reactions or buildup of the SEI.^{28{30} On the other hand, MD simulations accounting for reactivity with a reactive force field (ReaxFF) can calculate bond breaking/bond forming events. They have been used to investigate the pathways and activation energies of the reduction reactions of EC solvent molecules in LIB.^{31,32} Some simulations have elucidated certain aspects of the SEI buildup on Li-metal and silicon based anodes, capturing the electrolyte decomposition but on a very short time scale (hundreds of picoseconds), more limited than non-reactive force-field

simulations can achieve.^{33,35} While ReaxFF can predict reactions on the fly, based on the dynamic evolution of the system, it requires the parametrization of all the atomic interactions in the system, limiting its applicability. Recently, Takenaka and co-workers³⁶ employed a hybrid MC/MD method to study the SEI formation mechanism on a lithium and sodium-ion batteries demonstrating the impact of larger and longer space-time scale simulations to understand complex chemical reaction processes. It is noted that their simulations are limited to a small set of pre-defined chemical reactions, that occur based on a interatomic distance, whereas our simulations allow the specification of additional geometry constrains.

Given the importance of the SEI in lithium batteries, improved molecular level understanding of SEI structure and Li ion transport can help design novel solvents and/or additives that would form SEIs with enhanced transport properties and increased voltage stability windows. Our goal is to develop a methodology that helps us understand SEI formation at the atomistic level, using classical MD simulations, by growing the SEI one molecule at a time, simulating the most relevant chemical reactions occurring within the electrolyte near the anode. MD simulations give us in particular a spatial component that continuum methodologies cannot determine, useful for designing SEI layers with improved properties.¹¹ In reality, chemical reactions are based on atomistic configurations, not only concentrations. Achieving this goal is however quite challenging. First of all, the time scale needed to simulate all these chemical reactions is typically way beyond the time scales reached even by classical MD. To properly model the atomic vibrations in an MD calculation, a step on the femtosecond time scale is required. On the other hand, chemical reactions happen on a pico/nanosecond time scale, and the SEI growth occurs on a much longer time. To overcome this time-scale challenge, we have developed a strategy to accelerate reactive classical molecular dynamics simulations and simulate hundreds or even thousands of chemical reactions within the electrolyte. Our approach resembles the one proposed by Takenaka et al.,³⁶ but with a richer set of chemical reactions as well as a more sophisticated algorithm to capture reactions based on the recent work of Gissinger et al.^{37,38} Our work also differs in the way

we handle the flow of lithium ions in the electrolyte and through the anode.

In this article, we present an atomistic MD simulation methodology, useful to investigate the early stages of SEI formation, described in section 2. We used this protocol to predict the SEI growth for an EC-based electrolyte. Our results are in concordance with the existing literature, and can be found in section 3, followed by concluding remarks in section 4.

2 Methods

2.1 Model systems

As illustrated in Figure 1, simulations were conducted using a system representing a half-cell (anode side) of a LIB; the rectangular simulation cell is comprised of a liquid electrolyte, 1 M LiPF_6 in EC, and a nonreactive amorphous silicon electrode.

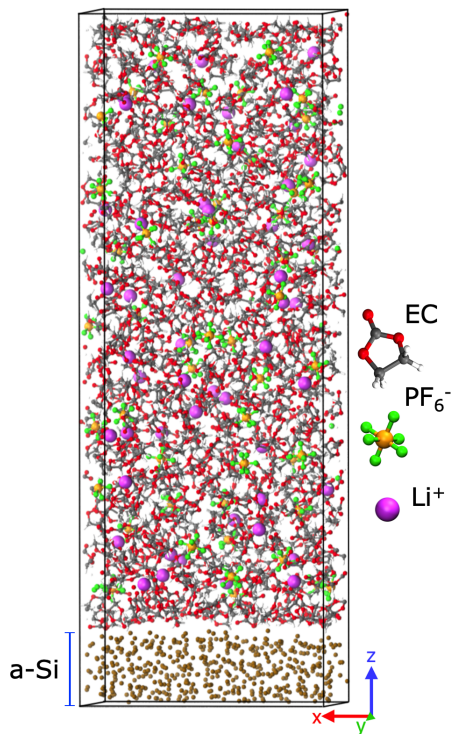


Figure 1: Simulation cell snapshot comprising 700 EC and 52 $\text{Li}^+ - \text{PF}_6^-$ pairs and an amorphous silicon anode (ochre).

An amorphous silicon (a-Si) slab of size of $32.58 \text{ \AA} \times 32.58 \text{ \AA} \times 8 \text{ \AA}$, was created first

using the melt-quench method and further relaxed for 2 ns at 298 K. An initial electrolyte system consisting of 350 EC molecules and 26 salt molecules randomly packed in a periodic cubic box of size $32 \text{ \AA} \times 32 \text{ \AA} \times 32 \text{ \AA}$ was generated using packmol.³⁹ The simulation domain was then replicated along the z direction, doubling the number of molecules in the system. The energy of the latter configuration was minimized by a conjugated-gradient minimization scheme, followed by an equilibration, with periodic boundary conditions (PBC) applied in x and y directions, for 2 ns in the isothermal-isobaric (NPT) ensemble at a pressure of 1 atm and a temperature of 298 K, with a time constant of 1 ps. An annealing process was conducted, and the electrolyte was heated from 298 K to 400 K for 1 ns, and maintained at 400 K for 1 ns, and subsequently annealed from 400 to 298 K in 1 ns. Subsequently, the simulation box was deformed to a final size of 32.58 \AA in both x and y directions, in order to match the size of the silicon electrode, followed by a relaxation in the z direction for 1 ns in the isothermal-isobaric (NPT) ensemble at a pressure of 1 atm and a temperature of 298 K. The final electrolyte density is 1.32 g/cm^3 , in good agreement with the experimental value.⁴ Finally, the electrode/electrolyte system was relaxed for 1 ns in the canonical ensemble (NVT) under a Nose-Hoover thermostat at 298 K and periodic boundary conditions were used in all directions.

2.2 MD reaction method

To investigate the reaction processes that lead to the SEI formation we have employed the REACTER algorithm,^{37,38} implemented in the Large-scale Atomic/Molecular Massively Parallel Simulator (LAMMPS) code⁴⁰ as *fix bond/react*. As pointed out before, one strong limitation with classical MD simulations is that non-reactive force fields do not allow chemical reactions to happen and reactive potential development is quite difficult. To overcome this restriction, one can identify when a chemical reaction is about to happen, and explicitly restart the MD simulation with different atomic potential parameters and potential function forms, that describe the reaction products.

In the present study, we have included a set of reactions involving ethylene carbonate reduction and PF_6^- reduction process based on both previous experiments and theoretical studies.^{13,41,45} The set of reactions shown in Table 1 leads to the formation of the various key SEI components detected experimentally, such as Li_2EDC , Li_2CO_3 , and LiF .^{46,50}

In order to carry out a reaction, the REACTER protocol consists in the following processes: (i) Search for potential reaction by checking that the distance between reaction site atoms is below the specified cutoff distance, (ii) For reaction sites that satisfy that condition, a matching algorithm compares the configuration of the reaction site and its surroundings at that time step with a pre-reaction template, (iii) The reaction is carried out if matching is successful. Every reaction requires the specification of pre-reaction (reactants) and post-reaction (products) templates. They contain not only the geometry and atomic charges of the reactants, but also the force field information, such that when a chemical reaction occurs, atomic potential parameters in the reactant molecules are replaced by those of the product.

Additional reaction constraints such as interatomic distances or angles can be specified in the templates and must be satisfied for a pre-reaction topology to be converted into a post-reaction topology. It is important to mention that this differs from the approach of Takenaka et al.³⁶ where the geometry of the reactants is ignored and only interatomic distances are used.

Two extra arguments are specified for each reaction: (i) frequency, *i.e.* the number of timesteps between checks for potential reactive sites, and (ii) the probability of an eligible reaction occurring. The probability can be tuned for different reaction pathways to proceed from the same reaction sites. In principle we could use actual reaction rates for the probabilities, but that would require accurate and reliable data which is hard to obtain. Also, these rates would need to be re-scaled to accelerate the products formation. In our simulation setup, the choice of these parameters was subsequent to the need of accelerating the reactions rates while conserving a reasonable overall behavior of the system, *i.e.* we observe the formation of the expected SEI components. We use a baseline frequency of 50 timesteps

Table 1: Set of electrolyte decomposition reaction used in our simulation. Also shown are the constraint types used for the REACTER protocol as well as the frequencies at which reaction templates are checked (in number of steps) and the probabilities associated with each reaction.

	Reaction	Reaction Constraints	Freq	Prob
EC	$\text{Li}^+ + \text{EC} \xrightleftharpoons{*} \text{Li}^+(\text{EC})$	1 distance (Oc-Li ⁺)	400	0.4 0.6
	$\text{Li}^+(\text{EC}) + \text{e}^{\zeta} \longrightarrow \text{o-LiEC}$	1 distance (Oe-Li ⁺) 1 angle (Oc-Cc-Oe)	200	0.4
	$2(\text{o-LiEC}) \longrightarrow \text{Li}_2\text{EDC} + \text{C}_2\text{H}_4 \uparrow$	1 distance (Oe ₁ -C ₂) ^a	50	1
	$2(\text{o-LiEC}) \longrightarrow \text{Li}_2\text{BDC}$	1 distance (C ₁ -C ₂) ^a	50	1
	$\text{o-LiEC} + \text{e}^{\zeta} \longrightarrow \text{o-LiEC}^{\zeta}$	1 distance (Cc-Li ⁺)	400	0.5
	$\text{o-LiEC}^{\zeta} \longrightarrow \text{LiCO}_3^{\zeta} + \text{C}_2\text{H}_4 \uparrow$	1 distance (O-Li ⁺)	2000	1
	$\text{LiCO}_3^{\zeta} + \text{Li}^+ \longrightarrow \text{Li}_2\text{CO}_3$	1 distance (O-Li ⁺)	50	1
	$\text{LiCO}_3^{\zeta} + \text{Li}^+(\text{EC}) \longrightarrow \text{Li}_2\text{EDC}$	1 distance (O ₁ -C ₂) ^a	50	1
PF ₆ ^ζ	$\text{PF}_6^{\zeta} + \text{Li}^+ + \text{e}^{\zeta} \longrightarrow \text{PF}_5^{\zeta} + \text{LiF} \downarrow$	1 distance (F-Li ⁺) ^b	200	1
	$\text{PF}_5^{\zeta} + \text{Li}^+ + \text{e}^{\zeta} \longrightarrow \text{PF}_4^{\zeta} + \text{LiF} \downarrow$	1 distance (F-Li ⁺) ^b	50	1
	$\text{PF}_4^{\zeta} + \text{Li}^+ \longrightarrow \text{PF}_3 + \text{LiF} \downarrow$	1 distance (F-Li ⁺) ^b	50	1

Oc: carbonyl oxygen, Cc: carbonyl carbon, Oe: ether oxygen

Probability for reversible reaction ($\text{Li}^+(\text{EC}) \longrightarrow \text{Li}^+ + \text{EC}$)

^a Involves one atom from molecule 1 and one from molecule 2

^b Any F atom in PF_x⁻

— given the high correlation between consecutive MD timesteps — and a probability of 1. Variations to these numbers will be explained in section 2.2.1. For each reaction listed in Table 1, we specify the type of constraints to be met, the template-matching frequency for the REACTER algorithm and probability to initiate a reaction. More detailed information about the reaction templates can be found in the supplementary material. It should be noted that the reactions implemented in our simulations can occur in any order.

For simplicity, the silicon anode is kept fixed during the reaction scheme, however it is crucial to our simulations since it acts as a barrier and a guide to as where reactions can

occur. Reduction reactions are constrained to happen only in a finite *redox region* near the silicon electrode. We initially set that region to be of 5 Å, then let it evolve with time, growing each 5 ns, and reaching thickness of 15 Å after 90 nanoseconds. Since electrons are not explicitly included in MD simulations, reduction reactions are based simply on the geometry of the reactant molecule and charge transfer is represented by changing the partial atomic charges to the product ones. Since it is easy to satisfy the geometry constraint in this case, reductive processes could occur spontaneously. To slow down this process, reactions involving a “free” electron are given a lower probability to occur and/or a longer period between between templates checking.

In order to neutralize the whole system after the electrolyte solvents or lithium salts are reduced, the same number of Li^+ cations as that of virtually injected electrons are placed at the electrode-electrolyte interface (non-reactive side) on the next time step after a reaction occurs. This does not affect significantly the pressure of the system. However, every time atoms are inserted, a 20 fs NVT run with no reactions is performed to equilibrate the system before returning to the reaction scheme.

All the reactions implemented happen on a much faster time-scale than they would in reality since our reaction rates are on the order of 1 if a reaction template is satisfied. With this acceleration, we can quickly end up with a deficiency in lithium ions needed to feed the reactions that lead to SEI, due to the slow diffusion of lithium ions in the electrolyte (see supplementary information). To avoid stalling the reactions, we also accelerate the lithium diffusion in the electrolyte to “match” the acceleration in reaction rates. This is done with an external artificial force in the z direction with magnitude of 0.02 eV/Å acting only on the lithium cations in the electrolyte. We choose this force large enough to increase lithium ion diffusion by an order of magnitude, but not too large not to affect the properties of the electrolyte. The radial distribution functions of the Li ions were calculated to verify that the structural properties of the electrolyte were not changed significantly by the external force on Li^+ ions and are available in the supplementary material.

If a lithium ion does not react in the SEI, it can pass through the silicon, under the influence of an external force in the Z direction and reappear on the electrolyte. This force — we use a value of $4.3 \text{ eV}/\text{\AA}$ — will ensure that lithium ions do not get trapped in the silicon cages, and quickly return to the electrolyte.⁵¹ It is only active within 1.2 \AA from the anode and drives lithium ions into the anode when they get close to it. In the vicinity of the anode, it can be interpreted as a classical substitute to the chemical force that would lithiate the silicon.

2.2.1 Reductive dissociation of EC and their implementations.

In this section we describe in more details the reactions involving the reductive process of ethylene carbonate solvents. These reactions were introduced based on the reported experimental data and theoretical studies. The first-principles data used for each EC solvent reaction described here and their reactants and products was obtained from the Lithium-Ion Battery Electrolyte (LIBE) dataset described in Ref. 52. Note that to be able to use reactants and products in the REACTER protocol in LAMMPS, some artificial covalent bonds in the templates had to be created in the force-field parameterization. Our reaction scheme centers on the reduction mechanism of ethylene carbonate molecules and the formation of three expected SEI components: an inorganic product Li_2CO_3 and two organic lithium bicarbonate compounds, $(\text{ROCO}_2\text{Li})_2$ ($\text{R} = \text{CH}_2, \text{CH}_2\text{CH}_2$). Studies using DFT calculations^{12,13,44,45} have predicted reaction pathways and confirmed the formation of the latter products. Several specific reaction pathways to LEOC obtained using a reaction network approach have been included.⁵³

To onset the reduction process, the first reaction captures a Li^+ coordinated solvent configuration $\text{Li}^+(\text{EC})$ by identifying when a Li^+ ion coordinates with a carbonyl oxygen (Oc) of a EC solvent molecule within a distance less than 1.85 \AA . The post-reacted topology includes a created Li^+-Oc bond to make it a single “molecule” to be used in the next reaction. We make this artificial bond soft enough not to affect the dynamics. We consider

this reaction reversible, such that when the Li^+ ion coordinating moves further from the solvent molecule, the bond can break and all atom types revert to the initial ones.

Often this $\text{Li}^+(\text{EC})$ complex is short-lived. Since EC preferentially coordinates to Li^+ cations, this reaction is often found in our simulations so a check for a new reaction site is performed only every 400 MD timesteps and the probability was set to 0.4. These parameters allowed us to reduce the potential reactive sites and yet accelerate the reaction rates. In order to favor the reversibility, while allowing some $\text{Li}^+(\text{EC})$ complex to survive longer and potentially react further, a probability of 0.6 is set for the $\text{Li}^+(\text{EC}) \longrightarrow \text{Li}^+ + \text{EC}$ reaction, while performing a template matching check every 400 timesteps.

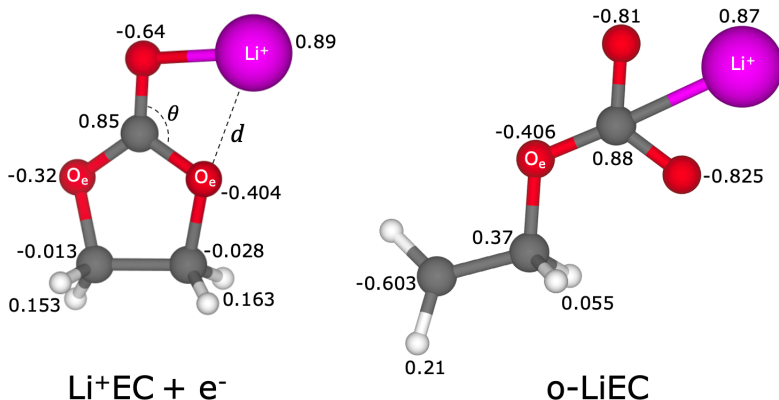


Figure 2: Pre-reacted (left) and post-reacted (right) templates for the one-electron reduction of $\text{Li}^+(\text{EC})$. The configuration of a Li^+ coordinated with an EC molecule is used as pre-reacted topology (left) and the post-reacted topology is the ring-opened $o\text{-LiEC}$. The reaction distance cutoff d does not exceed 2.1 Å and the angle θ must measure between 118° and 120° for the reaction to occur. The partial atomic charges of each atom before and after the reaction are shown.

$\text{Li}^+(\text{EC})$ molecules can undergo the one-electron reduction (see Fig. 2 for reaction templates). This redox process can only occur if the reaction site atoms are contained within the artificial redox region and the distance between the Li^+ cation and an ether oxygen (Oe) in the EC does not exceed the reaction distance cutoff. An angle constraint between a carbonyl oxygen, carbonyl carbon and ether oxygen must be satisfied as well. This will ensure that the $\text{Li}^+(\text{EC})$ geometry is ready for the C–Oe bond-breaking to form a ring-opened com-

plex o-LiEC. We found that checking for a potential reactant configuration only every 200 timesteps and lower the probability to 0.4 was necessary to leave some $\text{Li}^+(\text{EC})$ available to react with LiCO_3^\ddagger to generate Li_2EDC (last reaction in Table 1).

After the formation of o-LiEC molecules, three different pathways can follow: (i) A second reduction of the complex to form the negative charged molecule o-LiEC^{⋅-}, (ii) the direct combination of two o-LiEC to form dilithium butylene dicarbonate $(\text{CH}_2\text{CH}_2\text{OCO}_2\text{Li})_2$, or (iii) the formation of dilithium ethylene dicarbonate $(\text{CH}_2\text{OCO}_2\text{Li})_2$, the most common found product experimentally^{46,54} and usually considered as a dominant SEI component, via combination of two o-LiEC. For the second reduction a lower probability and larger frequency than other reactions has been assigned. In such a manner that this reaction does not consume all the products of the first reduction reaction as soon as they are formed, since it is simply based on the geometry and location of the reactant molecule in the simulation box. Without these modifications, the formation of LEDC and LBDC by combination of two ring-opened EC radicals could indeed never occur due to a lack of o-LiEC. The combination of two ring-opened EC radicals has been proposed as a potential reaction mechanism to form these molecules.^{13,50}

As proposed in multiple studies,¹³ we split the two-electron reduction path of $\text{Li}^+(\text{EC})$



into two reactions: a second reduction on the o-LiEC molecule and a time delayed bond-breaking — implemented though a less frequent check for reactants matching — by which a ethylene gas and an unpaired nucleophilic carbonate anion (LiCO_3^\ddagger) are generated.

The reaction $\text{o-LiEC}^\ddagger \longrightarrow \text{LiCO}_3^\ddagger + \text{C}_2\text{H}_4$ effectively should occur spontaneously once the o-LiEC^{⋅-} are formed. In our simulations, the o-LiEC^{⋅-} is formed near the anode (as required for a reduction) and rapidly undergoes the bond breaking to create two products, LiCO_3^\ddagger and a gas molecule. As the o-LiEC^{⋅-} molecules do not have time to diffuse in the SEI

before the bond breaking, we implemented a time delay by checking a possible reaction every 2000 timesteps, giving time to some of these molecules to move away from the anode. This reduces gas trapping between the anode and the SEI, since we do not expect gas buildup at the anode, a side effect of the acceleration.

Once the carbonate anion (LiCO_3^{\ominus}) is formed, two possible further reaction pathways are implemented: it may react with another Li^+ (EC) to form lithium ethylene dicarbonate (LEDC) or be paired with Li^+ to precipitate as insoluble inorganic lithium carbonate Li_2CO_3 .

2.2.2 Salt decomposition.

Several theoretical and experimental studies have discussed the decomposition of PF_6^{\ominus} anion, dissociated from the LiPF_6 salt, in EC-based electrolytes, suggesting possible reduction pathways, some of which involve water traces present in the electrolyte.^{16,18,41,55,56}

Our reaction scheme includes the PF_6^{\ominus} decomposition reaction mechanism proposed by Aurbach et al.⁴¹ to form a gaseous PF_3 molecule and three LiF complexes



Note that reaction has also been implemented in the hybrid MC/MD reaction method done by Takenaka et. al.³⁶ However, due to the complexity of the reaction which requires a very unlikely configuration with an anion surrounded by three Li^+ ions simultaneously at the right place and two electrons, along with the limitations of the REACTER algorithm, we have chosen to split the reaction into three consecutive reaction steps as shown in Table 1.

The reaction mechanism follows two reduction reactions and three Li^+ cations binding to form three LiF : (i) A Li^+ binds with a fluorine atom in the PF_6^{\ominus} anion creating a LiF and a virtually injected electron forms a PF_5^{\ominus} molecule. Since this reaction requires an electron it must occur near the anode surface in the defined redox region. Also, since it competes for Li^+ with the first EC reduction mechanism, we use the same period of 200 steps to check a

reaction. On the other hand, we kept a probability of 1 since PF_6^ζ anions are fewer than EC molecules. (ii) A reduction on the PF_5^ζ molecule and a second Li^+ cation binds with any F atom to form $\text{PF}_4^\zeta + \text{LiF}$, and (iii) a fluorine atom from the PF_4^ζ molecule bonds with a third Li^+ to reach the final products.

2.3 Computational details

The bonded interactions (bonds, angles, dihedrals, and impropers) were modeled as harmonic functions and the non-bonded included van der Waals interactions and Coulombic forces. Silicon anode interactions were modeled with the Stillinger-Weber (SW) potential.⁵⁷ Non-bonded interactions between amorphous silicon and electrolyte-SEI molecules were modeled by a simple Lennard-Jones potential. The bonded and non-bonded parameters for the electrolyte and SEI molecules were obtained from the non-polarizable force field OPLS-AA,⁵⁸ the PF_6^ζ anion from Lopes et al.,⁵⁹ and the parameters for the Li^+ and F^ζ ions were taken from Jensen et al.⁶⁰

The partial atomic charges for the electrolyte molecules EC, PF_6^ζ , and Li^+ were obtained by a full domain DFT calculation with our initial system and the RESP method⁶¹ at the PBE/DVZP level of theory using the CP2K⁶² package, and averaging charges for each atomic species. The geometries and partial charges for all EC reaction species were obtained from the LIBE dataset.⁵² These were calculated using the !B97X-V/def2-TZVPPD/SMD DFT level of theory, but were verified to give consistent charges compared to RESP in CP2K. Geometries and partial charges for the molecules in the PF_6^ζ decomposition reactions were calculated in CP2K at the PBE/DVZP level of theory.

The particle-particle particle-mesh (PPPM) method was applied to compute long-range Coulombic interactions. A cutoff distance of 12 Å was used for electrostatic interactions. A 12—6 Lennard-Jones potential was used for the van der Waals interactions. Production runs were carried out in the canonical (NVT) ensemble at 298 K using the Nose-Hoover

thermostat with a MD timestep of 1 fs.

3 Results & Discussion

We examined the formation processes of the SEI film on the anode surface via MD reaction simulations using the parameters described in the methodology. Figure 3 shows snapshots of the SEI reaction products, displaying how the electrolyte solution develops into the SEI film and gases during an MD reaction simulation up to 100 ns. Throughout the MD simulation o-LiEC complexes (yellow for EC and blue for Li^+ atoms) are constantly produced via reduction of EC solvent molecules on the anode surface as described in section 2.2.1. If single electron reduced species stay near the silicon surface long enough, they undergo a second reduction reaction forming o-LiEC[⋅] molecules, by dissociation the later ones generate ethylene gases (black) and LiCO_3^{\ominus} complexes (pink). At the same time, LiEC complexes can form Li_2EDC and Li_2BDC (cyan and green) via their radical polymerization. In our reaction scheme LiCO_3^{\ominus} molecules can either form Li_2EDC (cyan) or the inorganic compound Li_2CO_3 (red). Additionally, it is also shown that some PF_6^{\ominus} anions are reduced to form PF_3 gases (orange) and LiF complexes (F^{\ominus} colored in light purple and Li^+ cations in blue).

During the early stages, most of the produced open-ring LiEC complexes experience a second electron reduction as shown in Figure 4, where the production of o-LiEC and further evolution and dissociation into LiCO_3^{\ominus} is constant. Throughout the first nanoseconds, most of the LiCO_3^{\ominus} complexes react to form Li_2EDC followed by the formation of lithium carbonate Li_2CO_3 . We found that about 88% of the Li_2EDC molecules were produced from the two-electron transfer reduction mechanism. This is in agreement with previous studies,^{12,63} in which two-electron transfer reduction reactions are faster than the one-electron EC reduction. LiF and Li_2EDC are the predominant species of the initially formed SEI, according to our simulations. As the simulation continues, between 25 and 50 ns, the production of Li_2EDC slows down, whereas the formation of Li_2CO_3 increases continuously up to 60 ns.

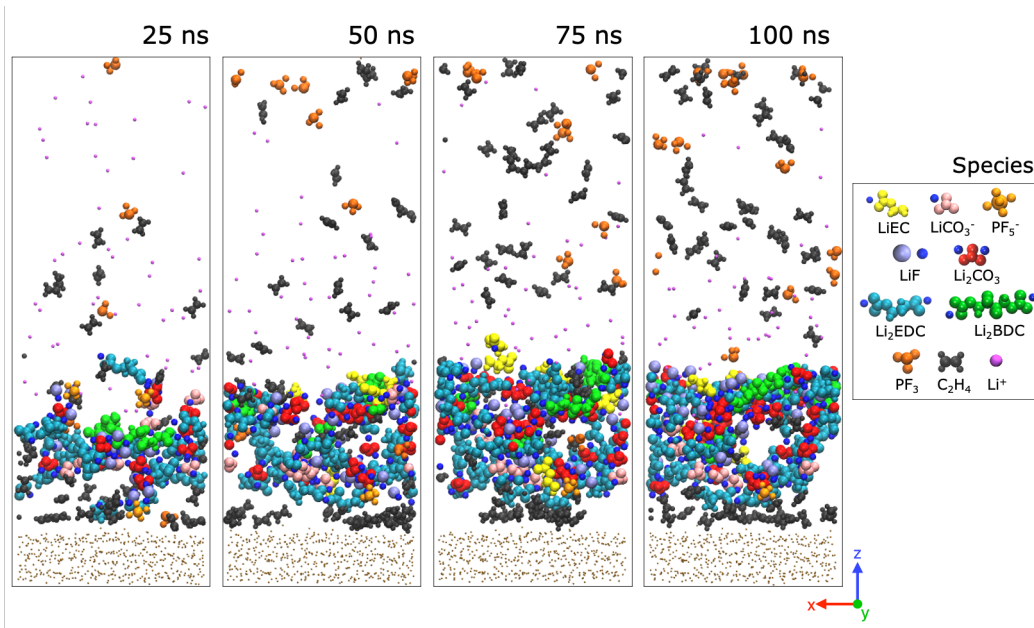


Figure 3: SEI film formation process in the EC-based electrolyte. Snapshots of the reaction products during 100 ns, shown every 25 ns. For visualization purposes, bulk EC and PF_6^- anions are not shown. Li^+ that have not been involved in any reaction are colored in magenta.

This is consistent with the idea that Li_2EDC is likely to form at a high EC concentration, while Li_2CO_3 tends to form at a low EC concentration.¹³ During the last 50 ns, the production of EC decomposition products slows down significantly, the formation of LiCO_3^- stops after 80 ns, since most of the o-LiEC complexes have moved far away from the redox region impeding their further reduction. In the outer regions of the SEI, LiEC produces lithium ethylene dicarbonate and an ethylene gas (C_2H_4) via a dimerization process.

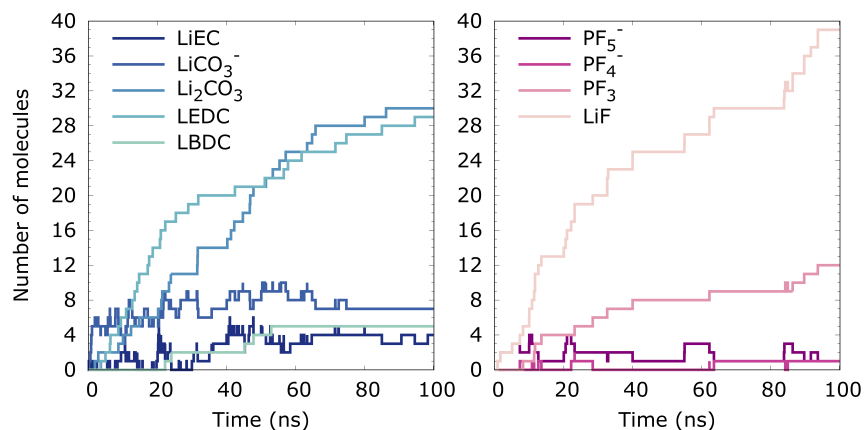


Figure 4: Evolution of reaction product counts during the 100 ns simulation.

As it can be observed from Fig. 4, LiF formation increases again rapidly after 80 ns. This is partially due to reaction of PF_5^{\ddagger} and PF_4^{\ddagger} molecules (both in light orange in Fig. 3) that have remained unreacted for long periods of time.

The SEI film at 100 ns is rather homogeneous in composition and does not display a clear double-layer structure. While the relatively short time scales of our simulations prevent us from observing an inorganic core and organic outer layer, we nonetheless observe structural trends that are consistent with the two-layer Peled model.⁶⁴ Figure 5 shows the density of the reaction products as a function of time. It is interesting to notice that at early stages the compounds LiF and Li_2EDC are more abundant near the anode surface, consistent with the interpretation of experimental results.^{48,65,66} It is also interesting to notice that at longer time scales, these complexes were distributed rather widely and diffusely over in the whole SEI film. The Li_2EDC molecules formed via dimerization (mostly after 75 ns of simulation) are mainly located on the outer SEI, in concurrence with the idea that organic complexes lay farther away from the anode. This is observed as well with Li_2BDC molecules, which start forming as LiEC complexes diffuse in the SEI, away from the anode to finally distribute on the outer SEI. Li_2CO_3 molecules tend to concentrate at the inner part of the SEI initially, but later diffuse, mostly aggregating in the middle.

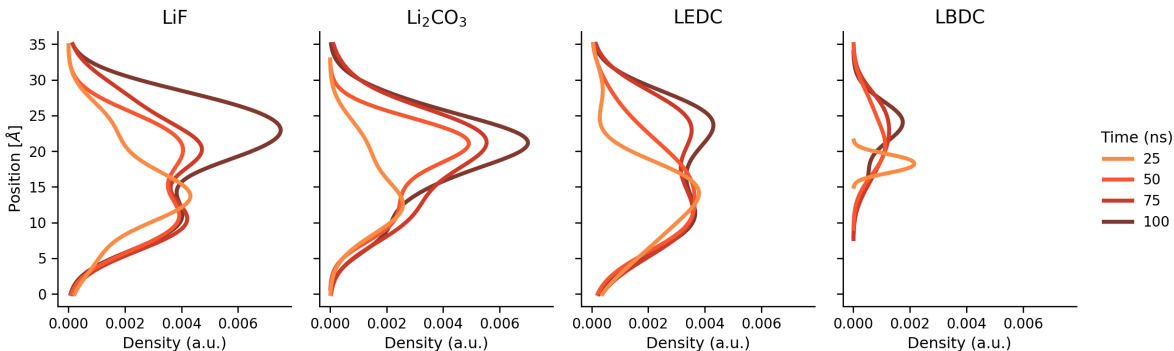


Figure 5: Density distributions of the product components (SEI film) in the present atomistic reaction simulations. The origin of the z coordinate is taken at the position of the silicon atoms on the anode surface in contact with the electrolyte.

A final note about the presence of C_2H_4 in our simulations. We know that in experiments,

gas is produced and released during the SEI growth. We could mimic its release by taking C_2H_4 out of our simulation cells, however we never reached the point where gas molecules agglomerate into bubbles. In addition, we do not want to assume that these molecules do not get trapped into the SEI or at least temporarily affect its growth. Finally, removing the gas molecules could change the density of the system. Based on this, we decided to keep the gas molecules in our simulations.

4 Conclusions

Time-scale is a major challenge when simulating SEI formation at the atomistic level. With recent advances in computer hardware, larger all-atom simulations can be carried out, but reducing time-to-solution for each time-step is more challenging with the end of Moore's law and the sequential nature of time. The approach described in this paper shows a way of accelerating the sequence of reactions by essentially associating artificially high reaction rates to each reaction, and making these reactions happen as soon as an appropriate reactants geometry is reached. We have established a set of parameters that leads to simulation results consistent with the current conjectures. We are aware that the results can be affected by these choices however this gives us a baseline for further studies. Indeed, the proposed computational approach can also be applied to investigate relative effects such as the impact of additives in the electrolyte on the initial SEI formation, giving helpful insights on how it could impact its evolution and growth. One limitation is obviously to have a good knowledge of all the reactions involved — either through first-principles simulations or experiments — and there is certainly room for improvement in this direction. Another limitation of our simulations is that, even with 100 ns of MD and an artificially high rate of reactions, we are far away from a fully formed SEI micro-structure. However, capturing the behavior at this initial stage is important to set the conditions for more extended modeling, particularly on the mesoscale, where longer time scales can be reached.

Supporting Information Available

Visualization of reactants and products and cutoffs or constraints used in the reaction templates, atomic coordinates and partial atomic charges for all species involved in PF_6^- decomposition reactions. Continuum diffusion analysis at atomistic scale and RDF plot for Li^+ under external force.

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Graphical TOC Entry

