

Cite this: *J. Mater. Chem. A*, 2022, 10, 11426

Li nucleation on the graphite anode under potential control in Li-ion batteries†

Arihant Bhandari,^{ab} Chao Peng,^{bc} Jacek Dziedzic,^{abd} John R. Owen,^{ab} Denis Kramer^{bce} and Chris-Kriton Skylaris^{*ab}

Application of Li-ion batteries in electric vehicles requires improved safety, increased lifetime and high charging rates. One of the most commonly used intercalation anode material for Li-ion batteries, graphite, is vulnerable to Li nucleation, a side reaction which competes with the intercalation process and leads to loss of reversible capacity of the battery, ageing and short-circuits. In this study, we deploy a combined grand canonical large-scale electronic density-functional theory (DFT) and Poisson–Boltzmann electrolyte theory to study the nucleation and growth of Li clusters on the graphite anode in the presence of its surrounding electrolyte environment at different applied voltages with respect to the Li metal reference electrode. We find the voltage below which the nucleation energy becomes negative (corresponding to Li nucleation becoming energetically favourable), the ‘potential of zero nucleation energy’ (U_{PZN}). We observe a distinct minimum in the plots of U_{PZN} as a function of the size of nucleated clusters. When the applied voltage on the graphite electrode is below the minimum value of U_{PZN} , the nucleated clusters start growing unbounded on graphite electrode. This potential for cluster growth (U_{PCG}) is found to be -0.12 V on the periodic basal plane of un lithiated graphite and -0.08 V on lithiated graphite. The corresponding potential for the zigzag edge termination is -0.06 V on un lithiated graphite and -0.04 V on lithiated graphite. Thus, the nucleation and cluster growth is favored on the zigzag edge termination of the graphite electrode as compared to the periodic basal plane and on the lithiated graphite as compared to the un lithiated graphite. We find that the surrounding environment plays a significant role and that nucleation is more likely to occur in electrolyte environment than that predicted from calculations in vacuum. We observe that the potentials obtained with grand canonical ensemble DFT method in electrolyte are close to experimentally available data. The study has profound implications for the nucleation, growth and control of metal dendrites in a battery cell.

Received 25th March 2022
Accepted 8th May 2022

DOI: 10.1039/d2ta02420a

rsc.li/materials-a

1 Introduction

Li-ion batteries have revolutionized the portable electronics industry and are being intensively pursued for application in the electrification of the transport sector.^{1–3} Key requirements of Li-ion batteries for successful implementation in electric vehicles are fast charging, long lifetime and high safety. Graphite-based carbonaceous materials are one of the most widely used

intercalation anode materials for Li-ion batteries since their inception.⁴ Thermodynamic and kinetic aspects of intercalation have been computationally studied for periodic graphite,⁵ edge-terminated graphite,⁶ and disordered carbon.⁷ The surface of graphite is susceptible to Li nucleation, a competing side-reaction which leads to a loss of reversible capacity. Moreover, Li nucleation followed by growth of dendrites can lead to a short-circuiting of the battery and is, hence, a critical safety challenge with Li-ion batteries.⁸ Li nucleation and dendrite growth also occurs on Li metal anodes.^{9,10}

In order to improve the performance of Li-ion batteries and gain more insight into this phenomenon, the process of Li nucleation has been extensively studied both experimentally and computationally in the recent years.^{11–13} Li nucleation is thermodynamically possible when the graphite electrode is at a lower potential (voltage) with respect to the Li/Li⁺ reference electrode.¹⁴ The voltage required for Li intercalation during a full charge of the graphite electrode corresponding to the phase transition from stage 2 to stage 1 graphite is around 0.065 – 0.085 V.¹⁵ When the overpotential for the Li intercalation

^aSchool of Chemistry, University of Southampton, Highfield, Southampton SO17 1BJ, UK

^bThe Faraday Institution, Quad One, Becquerel Avenue, Harwell Campus, Didcot, OX11 0RA, UK

^cEngineering Sciences, University of Southampton, Southampton SO17 1BJ, UK

^dFaculty of Applied Physics and Mathematics, Gdansk University of Technology, Gdansk 80-233, Poland

^eHelmut-Schmidt-University, University of the Armed Forces, 22043 Hamburg, Germany. E-mail: C.Skylaris@soton.ac.uk

† Electronic supplementary information (ESI) available: Files for all calculations are attached for reference and reproduction of results. Fig. S1: Enlarged plot of Fig. 2 for small Li clusters ($1 \leq n \leq 12$). See <https://doi.org/10.1039/d2ta02420a>



is large enough, the voltage of the graphite electrode will drop below 0.0 V with respect to Li/Li^+ , and Li nucleation becomes thermodynamically possible.¹⁶ Large overpotential for Li intercalation can easily occur at high current density during fast charging or at low temperature due to the Arrhenius dependence of the charge transfer resistance.¹² Detailed *in situ* experiments were performed by Gao *et al.* demonstrating the competition between Li intercalation and Li nucleation on graphite.¹³ Their observations were mostly limited to the kinetics of the two processes, with the observation of the intercalation of Li into the graphite until the graphite is fully lithiated, followed by nucleation of Li clusters at the graphite edges leading to dendrite formation and growth.

With the development of advanced atomistic methods, it is possible to gain more insight into the thermodynamics of the Li nucleation process.¹⁷ Fan *et al.* studied the adsorption of small Li clusters on graphene using canonical density functional theory calculations in vacuum (cDFTv) and found higher binding energy of Li clusters on graphite surface than on the (001) surface of Li metal, suggesting a possible cause for the initiation of Li nucleation.¹⁸ Liu *et al.* further studied the formation of Li clusters on graphene using cDFTv and found that the nucleation barrier is largely dependent on Li concentration ranging from over 3.2 eV at dilute concentrations to below 0.3 eV at high Li concentrations.¹⁹ Cui *et al.* also employed cDFTv to study the nucleation of small Li clusters with 2–6 Li atoms on graphene and found that the nucleation energy is positive and increases with the size of nucleated clusters, predominantly due to the lattice incompatibilities between the hexagonally-arranged carbon and the bcc structure of metallic Li.²⁰ They further found a respective decrease in the nucleation energy on amorphous C, N-doped C, carboxylated C, hydroxylated C and epoxied C, showing favorable nucleation. Chen *et al.* also studied Li binding on heteroatom-doped carbon using cDFTv and found that O-doped and O/B-co-doped carbons exhibit high lithiophilicity.²¹ We note here that all these aforementioned studies of Li nucleation using cDFTv are on the basal plane of single layer graphene, without considering the effect of a finite edge termination of the basal plane. Several studies on Li intercalation have shown a large effect of the presence of a finite edge termination of the graphite basal plane.^{6,22} Therefore, it is also important to study Li nucleation at finitely terminated graphite edges.

In our previous work we used cDFTv to study the nucleation of larger clusters with 1–65 Li atoms on multi-layered graphite for the first time, while also considering the effect of the presence of a finite edge termination of graphite basal plane.²³ We found that the nucleation energy barrier near the zigzag edge termination of the graphite is reduced by more than 1 eV relative to that on the periodic basal plane. We also found that chemical doping with nitrogen can increase the nucleation energy barrier and suppress Li nucleation. Most importantly, we observed a significant effect of the voltage on the nucleation process, with a decrease in the nucleation energy and energy barrier on lowering the potential. While these conclusions have significant impact on the understanding of Li nucleation, there are inherent limitations in state-of-the-art cDFTv methods.

cDFTv calculations are performed in ultra high vacuum conditions under a charge control regime, where the charge on the entire structural model is fixed to zero. This regime is schematically shown in Fig. 1a. Here the calculations are performed for an isolated neutral Li slab and an isolated neutral $\text{Li}_n|\text{G}$ system, where an n -atom Li cluster has been nucleated on graphite (G). The two systems are completely independent and do not interact electrically *via* a ‘wire’ that would allow transfer of electrons under the constraint on their electrochemical potentials. To overcome these limitations and to simulate electrochemical systems under experimental conditions, we have developed a model for grand canonical DFT calculations.²⁴ Here, electrons can flow *via* an electrical connection between the working electrode and the reference electrode under potential control. The electrochemical potential of electrons at the working electrode ($\tilde{\mu}_e^-$) can be set at a certain voltage (U) with respect to that at a reference electrode: $\tilde{\mu}_e^-(U) = \tilde{\mu}_e^{\text{ref}} (U = 0) - e \cdot U$. Our model parameters have been calibrated according to the reduction potential of the Li metal electrode.²⁴ As the number of electrons varies in response to the electrochemical potential, the electrode can have a net charge. The grand canonical calculations can be performed in vacuum (gcDFTv), as shown in Fig. 1b or in an implicit solvent-electrolyte environment (gcDFTe), shown in Fig. 1(c). In vacuum, the net charge on the electrode is neutralized by a uniform opposite background charge (‘jellium’). In a solvent environment, the charged electrode system is neutralized by a Poisson–Boltzmann electrolyte. The presence of a net charge on the electrode system leads to a shift in the electrolyte concentrations near the electrode surface, such that there is more of the oppositely charged ‘counter-ions’ than the ‘co-ions’ leading to the formation of double layers. The concentrations of the electrolyte ions reach their asymptotic bulk values far away from the surface. This experimentally observed phenomenon is implemented within the neutralization by electrolyte concentration shift (NECS) method.²⁵ This method overcomes the limitations of the prevalent ‘jellium’ based approaches which introduce artificial opposite uniform background charge to neutralize the electrode charge.²⁵ Our method works *via* the solution of a modified Poisson–Boltzmann equation using a highly-parallel, multigrid library DL_MG .^{26,27} The entire electrolyte model has been implemented in the order- N electronic total energy program (ONETEP), which allows large-scale DFT computations with the computational cost scaling linearly with the number of atoms (N).²⁸ The model has been previously tested to accurately predict activity coefficients of LiPF_6 in ethylene carbonate (EC),²⁹ and the differential capacitance of graphite electrodes.²⁴

In this paper we extend our previous work with cDFTv,²³ to apply our newly developed grand canonical model to study Li nucleation on graphite electrodes in realistic electrochemical conditions, in order to answer the following questions: (1) What voltage marks the onset of Li nucleation? (2) What is the critical voltage below which the nucleated Li clusters start growing spontaneously? (3) How does the presence of environment affect the nucleation energetics? (4) Where on graphite do the Li clusters prefer to nucleate? (5) What is the effect of the degree of



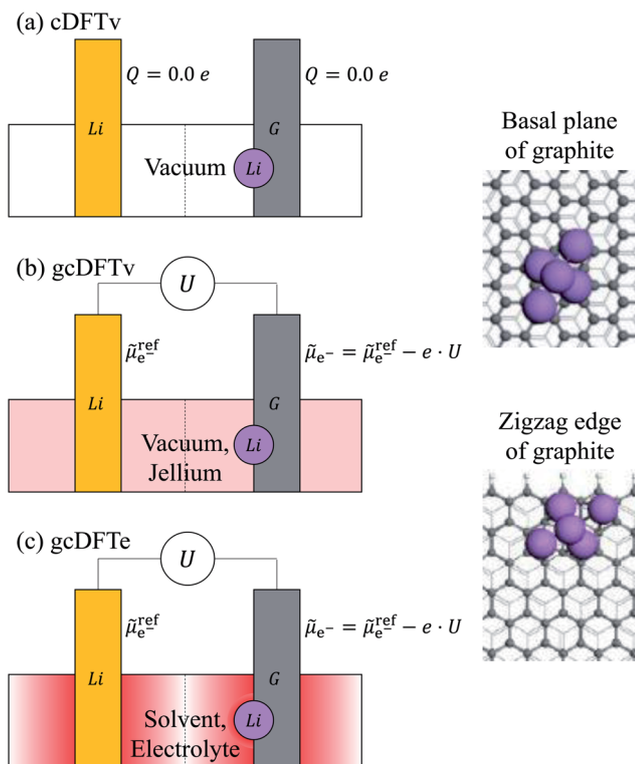
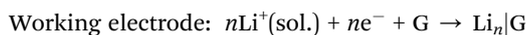


Fig. 1 Schematic of a Li nucleation process in different regimes: (a) zero charge regime using canonical DFT calculations in vacuum (cDFTv). The total charge on each of the electrodes is constrained to be zero, while there is no constraint on the electrochemical potentials of electrons at the two electrodes. (b) Potential control regime using grand canonical DFT calculations in vacuum (gcDFTv). The net charge on the electrode is neutralized by a uniform opposite background charge ('jellium'). (c) Potential control regime using grand canonical DFT calculations in electrolyte (gcDFTe). In a solvent environment, the charged electrode system is neutralized by a Poisson–Boltzmann electrolyte. In grand canonical methods (b and c), the applied potential (U) controls the electrochemical potential of electrons ($\tilde{\mu}_{e^-}$) at the graphite electrode (G) with respect to that at the Li reference electrode ($\tilde{\mu}_e^{\text{ref}}$).

lithiation of graphite on the nucleation process? We now present our theory (Sec. 2), computational details (Sec. 3), results and discussion (Sec. 4), followed by conclusions (Sec. 5).

2 Theory

The Li nucleation reaction on graphite can be written as:



The Li^+ ions move through the solution, while the electrons (e^-) travel through the outer electrical circuit. G represents the graphite and $\text{Li}_n|\text{G}$ represents an n -atom Li cluster nucleated on a graphite working electrode. The nucleation energy is the change in free energy due the reaction (ΔQ). For an electrochemical cell, the ionic path is usually

thermodynamically reversible but the electronic path may be interrupted by a potentiometer, in which case the potential difference (U) on the potentiometer gives an experimental value for the nucleation energy ($\Delta Q = -n \cdot e \cdot U$, where e is the elementary charge). Conversely, in a potential controlled experiment, the voltage U is applied which results in the nucleation of a Li cluster assuming thermodynamic equilibrium. Similarly, in the present study we could expect the nucleation energy ($\Delta Q(U)$) to indicate whether a cluster of a given size n will form at a voltage (U). The nucleation energy ($\Delta Q(U)$) is calculated as:

$$\Delta Q(U) = Q_{\text{Li}_n|\text{G}}(U) - Q_{\text{G}}(U) - n \cdot \tilde{\mu}_{\text{Li}}^{\text{ref}}(U=0), \quad (1)$$

where Q is the grand free energy calculated in a grand canonical ensemble of electrons and electrolyte ions.²⁴ U is the voltage of the graphite electrode with respect to the Li reference (ref) electrode which is at $U = 0$ V. $\tilde{\mu}_{\text{Li}}^{\text{ref}}$ is the electrochemical potential of Li at the reference electrode, which is calculated from the change in the grand free energy ($Q_{\text{Li}}^{\text{ref}}$) of the Li reference electrode slab with respect to the number of Li atoms in the slab ($n_{\text{Li}}^{\text{ref}}$):

$$\tilde{\mu}_{\text{Li}}^{\text{ref}}(U=0) = \frac{\Delta Q_{\text{Li}}^{\text{ref}}(U=0)}{\Delta n_{\text{Li}}^{\text{ref}}} \quad (2)$$

3 Computational details

We perform simulations for Li nucleation using the ONETEP linear-scaling density functional theory (DFT) program,^{28,30} where the computational cost scales linearly with the number of atoms as opposed to the cubic scaling of conventional DFT. The simulations under potential control are performed using the grand canonical model for electrochemistry simulations,²⁴ utilizing the neutralization by electrolyte concentration shift (NECS) method for a modified Poisson–Boltzmann equation (P-BE).^{25,29} The P-BE is solved using a bespoke, highly parallel, multigrid solver DL_MG .^{26,27}

The kinetic energy cutoff for the basis set, which consists of psinc functions and is equivalent to plane waves, is 1000.0 eV.³¹ The radius of the localized orbitals called as the non-orthogonal generalized Wannier functions (NGWFs) is set to 8.0 a_0 for all calculations.³² The convergence threshold for the electronic energy is 10^{-6} eV and that for the NGWF rms gradient is 5×10^{-7} a.u. The convergence threshold on the number of electrons in the grand canonical ensemble is 10^{-6} electrons per atom. For van der Waals interactions, a D2 dispersion correction has been used.³³ In the grand canonical calculations the electrochemical potential of electrons in the Li reference electrode is set to -1.39 eV (*cf.* Table 1 of ref. 24) corresponding to the experimental value, while we set the applied voltage of the graphite electrode at $U = -0.1, 0.0, 0.1$ V with respect to the Li metal reference electrode. The structural geometries for Li clusters with 1–65 atoms and graphite electrodes have been obtained from AIMD simulations in our previous work using cDFTv as shown in Fig. 1, S3 and S5 of ref. 23. The chemical formula of



Table 1 Chemical formula and simulation cell size of various structural models

Graphite model (G)	Chemical formula (Li _n G, 1 ≤ n ≤ 65)	X (Å)	Y (Å)	Z (Å)
Unlithiated basal plane	Li _n C ₅₁₂	34.17	19.73	100.03
Lithiated basal plane	Li _n Li ₁₀₀ C ₉₀₀	37.38	21.58	100.00
Unlithiated zigzag edge	Li _n C ₆₄₈ H ₃₆	53.41	22.19	67.73
Lithiated zigzag edge	Li _n Li ₁₀₂ C ₉₇₂ H ₅₄	57.38	22.42	67.77

structural models and size of simulation cell are summarized in Table 1.

The size of the graphite model is much larger than the number of atoms (n) in the Li cluster, so that the interaction between periodic images is negligible. The equilibrium distance of Li clusters from the graphite electrode is further optimized in electrolyte environment for each applied voltage by minimizing the grand free energy of the system as a function of the distance of the Li cluster from the graphite surface.

The surrounding environment is described by an implicit 1.0 M LiPF₆ electrolyte in ethylene carbonate solvent (EC). The dielectric permittivity of the EC solvent is described using the soft-sphere model,³⁴ and varies smoothly from 1.0 near the electrode to that of the bulk solvent which is set to 90.7 as per the experimental value at 298.15 K.³⁵ For an even closer approximation of the environment in Li-ion batteries, one can consider using the dielectric constant of a mixture of solvents.³⁶ The soft-sphere radius for the transition of the dielectric permittivity is calibrated from the reduction potential of standard reference electrodes (*cf.* Tables 2 and 3 of ref. 24). Correspondingly, soft-sphere radii of 3.048 a_0 for Li, 3.907 a_0 for C and 2.644 a_0 for H are used. The solute–solvent cavitation, dispersion and repulsion interactions are taken as proportional to the solvent accessible surface area, with the proportionality constant being the scaled surface tension of the solvent.³⁷ The surface tension of EC is taken to be 0.0506 N m⁻¹ as per the experimental value.³⁸ The accessibility function of the implicit electrolyte is also described *via* a soft-sphere model,²⁹ varying from 0.0 near the electrode to 1.0 in the bulk electrolyte. The radius of transition spheres is set by considering the total size of the solute surrounded by a solvation shell. The solute size is described by isoradii of the electronic density of isolated atoms at an isovalue of 0.001 e/a_0^3 . A solvation shell radius of 3.0 a_0 is used. The values of these parameters have been previously calibrated to yield realistic activity coefficients of LiPF₆ in EC.²⁹

We calculate the nucleation energy for Li clusters with 1–65 atoms on the graphite electrode at different voltages ($U = -0.1, 0.0, 0.1$ V) with respect to the Li reference electrode. From the trend of the variation of the nucleation energy with voltage, we find the voltage which would give a zero nucleation energy. This potential of zero nucleation energy (U_{PZN}) is then plotted as a function of the size of the cluster. The minimum in the potential of zero nucleation energy is the critical voltage for cluster growth, below which the nucleated clusters start growing unbounded on graphite. Regarding the location of nucleation on graphite, we consider Li nucleation on the

periodic basal plane (far from the edge), as well as close to the zigzag edge termination of the basal plane. We further consider the two end cases where graphite is fully unlithiated and fully lithiated.

4 Results and discussion

4.1 Li reference chemical potential ($\tilde{\mu}_{\text{Li}}^{\text{ref}}$)

For the Li reference electrode, we consider (100) slabs of Li metal with 11, 13, 15 layers at an applied voltage of $U = 0.0$ V and compute the grand free energy using the grand canonical DFT method in electrolyte (gcDFTe).²⁴ The electrolyte is 1 M LiPF₆ in ethylene carbonate (EC) solvent. From the change in grand free energy with the number of Li atoms, we compute the chemical potential of Li at the reference electrode ($\tilde{\mu}_{\text{Li}}^{\text{ref}}$) using eqn (2). We also consider a grand canonical calculation in vacuum (gcDFTv), where the charge of the electrode is neutralized by an opposite uniform background charge ('jellium'). The computed values of ($\tilde{\mu}_{\text{Li}}^{\text{ref}}$) are shown in Table 2. We see a large difference in the computed values of ($\tilde{\mu}_{\text{Li}}^{\text{ref}}$) with the choice of environment. The values in vacuum are higher by approximately half an electron volt (eV) than those in EC/1 M LiPF₆, which is more representative of the actual environment in Li-ion batteries. The computed chemical potential of Li at the reference electrode ($\tilde{\mu}_{\text{Li}}^{\text{ref}}$) feeds into the calculation of the nucleation energy of Li clusters at the graphite electrode *via* eqn (1).

4.2 Nucleation energy ($\Delta\Omega$)

The nucleation energy ($\Delta\Omega$) calculated from eqn (1) informs whether the formation of a cluster of a particular size (n) is thermodynamically feasible. A negative nucleation energy suggests a thermodynamically feasible nucleation, while a positive nucleation energy indicates that nucleation is not feasible. In Fig. 2, we show the nucleation energy in the grand canonical ensemble ($\Delta\Omega$) computed for Li clusters of 1–65 atoms on the graphite electrode. The simulations are performed under potential control at different voltages ($U = -0.1, 0.0, 0.1$

Table 2 The computed chemical potential of Li metal at the reference electrode ($\tilde{\mu}_{\text{Li}}^{\text{ref}}$) using eqn (2)

Environment	Vacuum, jellium	EC, 1 M LiPF ₆
$(\tilde{\mu}_{\text{Li}}^{\text{ref}})(U = 0)$ (eV)	-200.6018	-201.0472



V) with respect to the Li reference electrode using the grand canonical ensemble DFT method.²⁴ We have considered both lithiated and unlithiated graphite electrodes, while considering nucleation on the periodic basal plane, as well as near the finite zigzag edge termination of graphite. The nucleation energy is calculated using the grand canonical DFT method in an electrolyte solution (gcDFTe), as well as in vacuum (gcDFTv). In vacuum, the surface charge on the electrode is neutralized by an opposite uniform background charge ('jellium'). In implicit solvent, the charged electrode is neutralized by the excess of oppositely charged electrolyte near the surface *via* the NECS method.²⁵ The bulk concentration of LiPF₆ electrolyte is 1 M.

As a general observation for large clusters, the nucleation energy for Li clusters on the graphite electrode at voltages $U \geq 0.0$ V with respect to the Li reference electrode is positive and monotonically increasing. This suggests that nucleation is generally unlikely to occur at non-negative voltages ($U \geq 0.0$ V). The nucleation energy on a graphite electrode at a voltage, $U = -0.1$ V with respect to the Li reference electrode becomes negative at least beyond a certain cluster size for all the cases considered, suggesting that Li nucleation is possible on a graphite electrode at a negative voltage with respect to the Li reference electrode. To find the voltage below which nucleation energy becomes negative, we compute the potential of zero nucleation energy, which is described next.

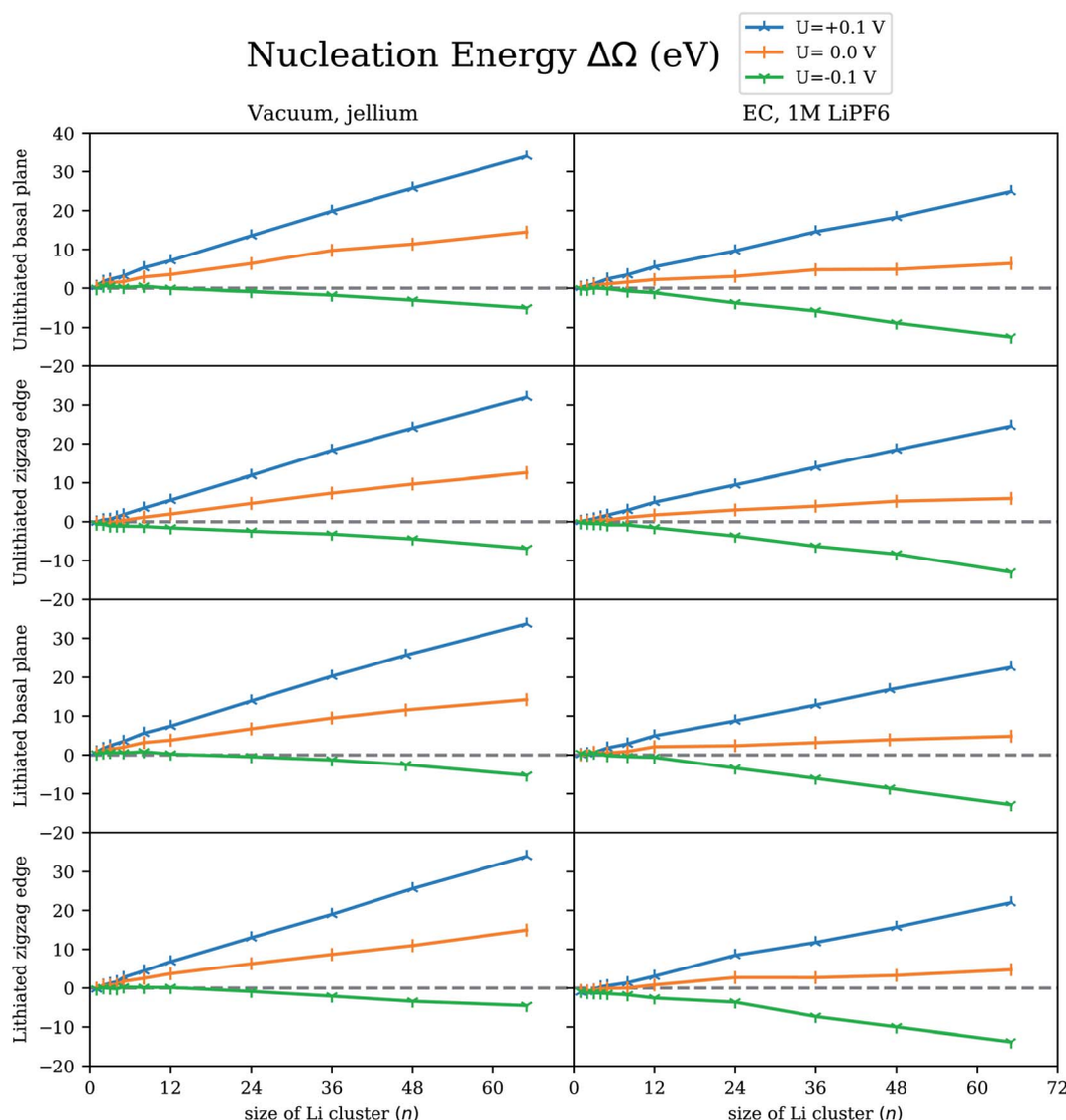


Fig. 2 The computed nucleation energy ($\Delta\Omega$) for Li clusters with 1–65 atoms on the graphite electrode at voltages $U = -0.1, 0.0, 0.1$ V with respect to the Li reference electrode using the grand canonical ensemble DFT method.²⁴ We have considered both lithiated and unlithiated graphite electrodes, while considering nucleation on the periodic basal plane as well as near the finite zigzag edge termination of graphite. The nucleation energy is calculated in vacuum as well as in ethylene carbonate (EC) solvent environments. Two types of neutralization schemes are considered: an opposite uniform background charge ('jellium') and neutralization by electrolyte concentration shift (NECS).²⁵ The electrolyte is 1 M LiPF₆.



4.3 Potential of zero nucleation energy (U_{PZN})

The potential of zero nucleation energy (U_{PZN}) is the voltage for which the nucleation energy ($\Delta\Omega$) computed from eqn (1) becomes zero. In other words, it is the voltage below which the formation of a Li cluster of a particular size becomes thermodynamically feasible. For each cluster size, the potential of zero nucleation energy is calculated by interpolating the calculated nucleation energy ($\Delta\Omega$) at the three potential values of $U = -0.1, 0.0, 0.1$ V to obtain U_{PZN} where the nucleation energy, $\Delta\Omega = 0$. This potential of zero nucleation energy (U_{PZN}) is plotted in Fig. 3 for all the clusters with 1–65 atoms in all considered situations. The nucleation of a cluster of a particular size (n) occurs when the applied voltage at the graphite electrode is below the value of $U_{PZN}(n)$. Comparing different situations, we see a significant difference between the results in vacuum with

jellium as compared to the electrolyte environment, the latter being more representative of the actual environment in Li-ion batteries. The U_{PZN} is higher in the electrolyte environment than in vacuum, suggesting that the Li nucleation is more likely to occur in the electrolyte environment, than what would be predicted from vacuum calculations. From a comparison between the basal plane and zigzag edge, we see that U_{PZN} is higher near the zigzag edge than on the basal plane, suggesting the nucleation to be more likely on the zigzag edge termination of the graphite electrode than the periodic basal plane. The value of $U_{PZN}(n)$ gives the necessary condition on the voltage for nucleation of a particular cluster, which leads to the next question as to what the critical voltage for cluster growth is.

Potential of Zero Nucleation Energy, U_{PZN} (V)

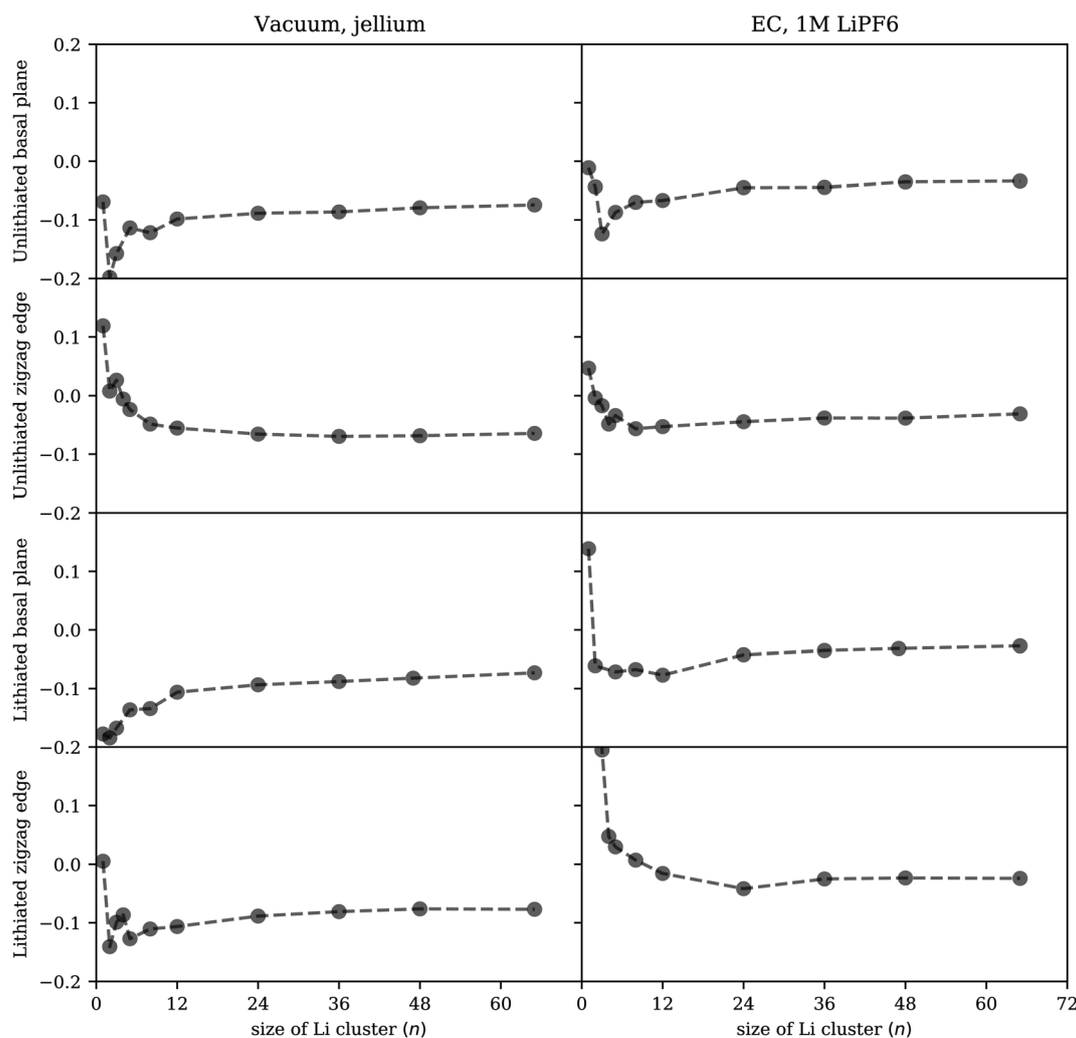


Fig. 3 The potential of zero nucleation energy (U_{PZN}) as a function of the size of the Li cluster (n), obtained from interpolating the nucleation energy computed for potentials ($U = -0.1, 0.0, 0.1$ V) in Fig. 2. The left panel shows the U_{PZN} computed from the grand canonical DFT method in vacuum (gcDFTv), while the right panel shows the same from the grand canonical DFT method in electrolyte (gcDFTe). The electrolyte is 1 M LiPF₆ in EC solvent.



Table 3 Potential for cluster growth (U_{PCG}) on graphite with respect to the Li reference electrode (below which the nucleated clusters start growing unboundedly)

Ensemble		Canonical	Grand canonical		Experiments
Environment		Vacuum (cDFTv)	Vacuum, jellium (gcDFTv)	EC, 1 M LiPF ₆ (gcDFTe)	EC/DEC (1 : 1), 1 M LiPF ₆
System					
Unlithiated graphite	Basal plane	−0.65 V (ref. 23)	−0.20 V	−0.12 V	−0.15 V to
	Zigzag edge	−0.31 V (ref. 23)	−0.07 V	−0.06 V	−0.04 V (ref. 13)
Lithiated graphite	Basal plane	N.A.	−0.18 V	−0.08 V	
	Zigzag edge	−0.47 V (ref. 23)	−0.14 V	−0.04 V	

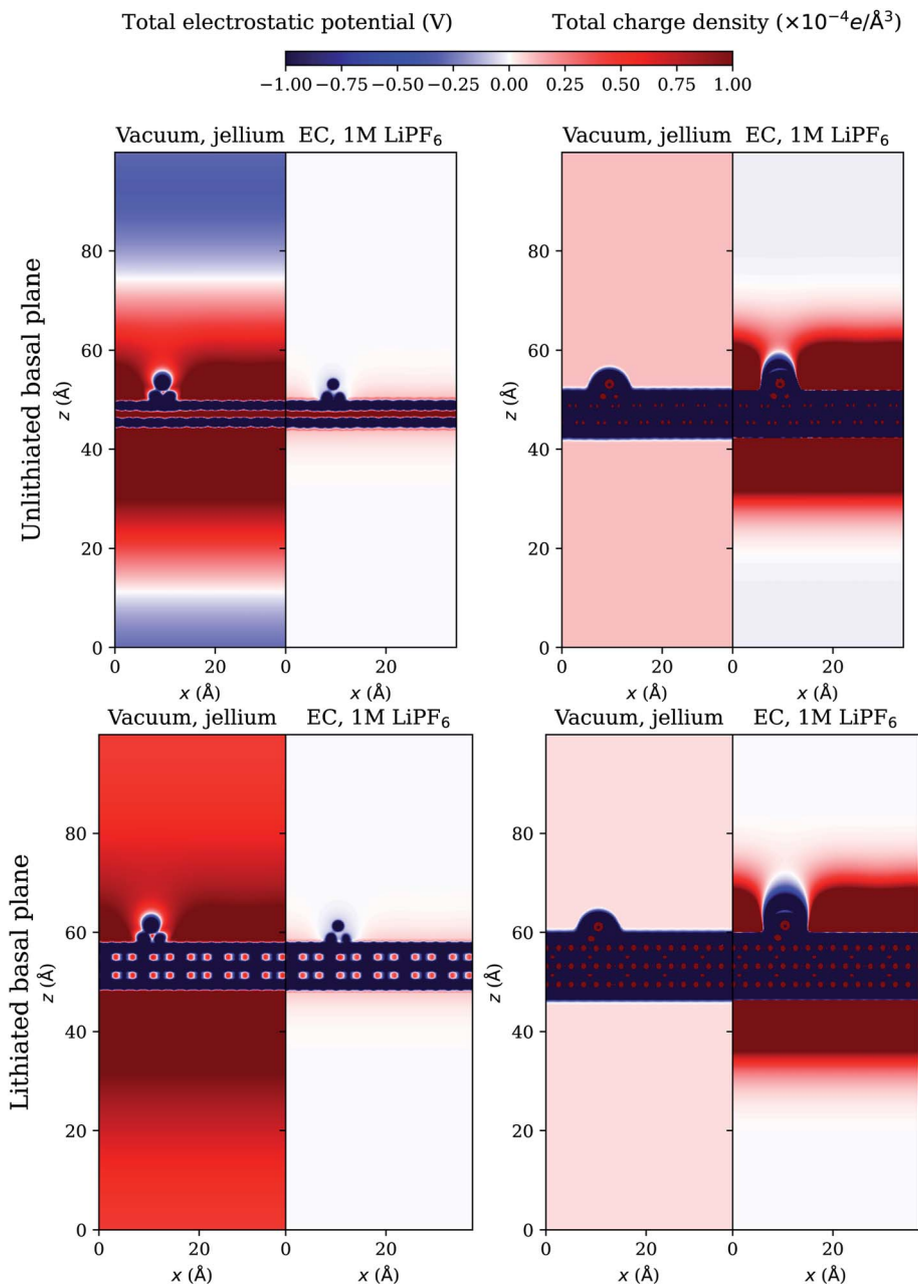


Fig. 4 Total electrostatic potential (left) and charge density (right) in a plane cutting through a 5-atom Li cluster nucleated on the periodic basal plane of the graphite electrode at $U = -0.1$ V, computed using the grand canonical DFT method.



4.4 Potential for cluster growth (U_{PCG})

In each case of Fig. 3, the U_{PZN} values represent the threshold applied potential below which the nucleation of a Li cluster of size n is possible. For $n = 1$, we find this potential close to zero in many cases and even at positive potentials under some conditions. Proceeding to larger n values, we see a decrease of the U_{PZN} with cluster size. This suggests that subsequent cluster growth requires successively larger (more negative) overpotentials until the minimum U_{PZN} is reached. Beyond this cluster size, the U_{PZN} increases so that spontaneous growth can occur without an increased overpotential, and in some cases even at reduced overpotentials. Thus we can define a potential for cluster growth,

$$U_{\text{PCG}} = \min_n U_{\text{PZN}}(n), \quad (3)$$

as the potential for (uncontrolled) cluster growth, that can only be stopped by applying a more positive potential. This potential for cluster growth (U_{PCG}) is shown in Table 3. We also compare our results computed using the grand canonical ensemble DFT method (gcDFT) with our previous work using canonical DFT simulations in vacuum (cDFTv).²³ We see that the magnitude of potential for cluster growth (U_{PCG}) computed with the grand canonical DFT method is much lower than that calculated with the cDFTv approach. This suggests that the growth of clusters on the graphite electrode would occur much earlier than predicted by cDFTv calculations. This signifies the importance of the advanced gcDFT model over the conventional cDFTv models. Now, comparing the gcDFT in different environments, we find that the magnitude of U_{PCG} in the electrolyte

environment is lower than that in vacuum, suggesting that the growth of clusters is much easier in electrolyte than in vacuum. From a comparison of the location of Li nucleation, we observe that cluster growth is easier on a zigzag edge termination than on the periodic basal plane of graphite. From a comparison of the degree of lithiation of graphite, we note that the cluster growth is preferred on lithiated graphite compared to unli-thiated graphite. A recent *in situ* experimental study by Gao *et al.* (which focused more on the kinetic aspects of Li nucleation) also observed distinct minima in the voltage–capacity plots shown in Fig. S7 of ref. 13. Their value of minimum potentials (nucleation barriers) vary from -0.04 V to -0.15 V with respect to the Li reference electrode, which is quite close to the values predicted from gcDFTe.

4.5 Electrostatic potential and charge density

The effect of the environment can be more evident from the plots of electrostatic potential and charge density. We investigate the case of a 5-atom Li cluster on graphite surface at $U = -0.1$ V and plot the total electrostatic potential and charge density in a plane cutting through the Li cluster on the periodic basal plane in Fig. 4 and on the zigzag edge termination of the graphite in Fig. 5. The total electrostatic potential of the system in vacuum with jellium is diffuse and extends all over the simulation cell. In the presence of electrolyte, the total electrostatic potential is effectively screened at the interface so that it decays rapidly as it should in bulk electrolyte. The total charge density is shown in the right panels. In vacuum, the negatively charged electrode system is neutralized by the uniform

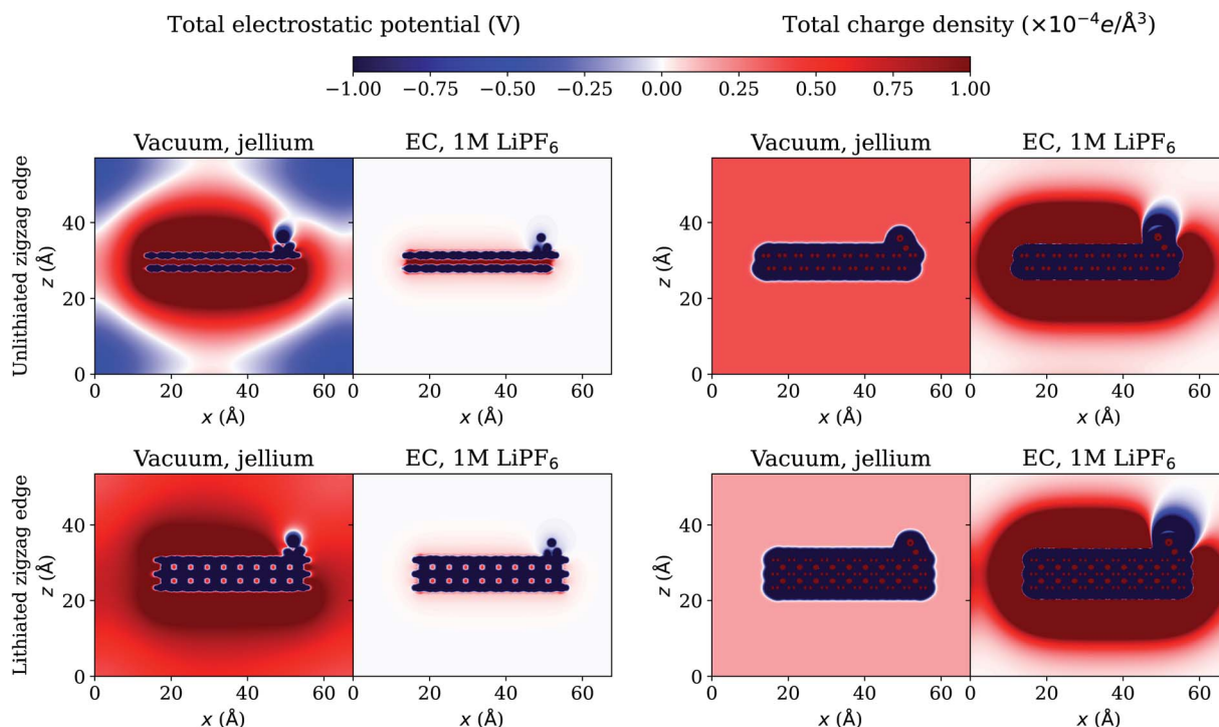


Fig. 5 Total electrostatic potential (left) and charge density (right) in a plane cutting through a 5-atom Li cluster nucleated near the zigzag edge termination of the graphite electrode at $U = -0.1$ V, computed using the grand canonical DFT method.



Table 4 Charge on the electrode system for a of 5-atom Li cluster on graphite at $U = -0.1$ V computed using grand canonical DFT

Environment		Vacuum, jellium	EC, 1 M LiPF ₆
System		(gcDFTv)	(gcDFTv)
Unlithiated graphite	Basal plane	-0.73e	-12.75e
	Zigzag edge	-2.96e	-23.67e
Lithiated graphite	Basal plane	-0.41e	-11.58e
	Zigzag edge	-1.45e	-19.96e

positively charged jellium which extends all over the simulation cell. In electrolyte environment, the negatively charged electrode system is neutralized by the build up of positively charged ions near the interface, forming a double layer and the electrolyte charge density decays to zero far away in bulk electrolyte. The total charge density integrates to zero in the simulation cell in all cases, which is a necessary requirement for the convergence of the electrostatic potential in periodic boundary conditions.²⁵ The net charge on the electrode system is shown in Table 4 for all cases. The net charge on the surrounding medium (jellium or electrolyte) is the opposite of this charge on the electrode system. The magnitude of the charging of the electrode system is much larger in electrolyte environment than what occurs in vacuum with jellium. All these observations highlight the limitations of conventional in-vacuum models based on jellium neutralization and show the realistic description with our advanced electrolyte model.

5 Conclusions

We studied the thermodynamics of the nucleation of Li clusters of 1–65 atoms and their growth on graphite electrodes at an applied voltage with respect to the Li reference electrode, using a combined grand canonical large scale density-functional and Poisson–Boltzmann Theory. We computed the nucleation energy of Li clusters for different values of the applied voltage, $U = -0.1, 0.0, 0.1$ V, with respect to the Li reference electrode. We find that the nucleation energy is positive and monotonically increasing at $U = 0.0$ V and $U = 0.1$ V, while it achieves a negative value beyond a certain cluster size for a negative potential $U = -0.1$ V. A negative value of the nucleation energy would mean an energetically favourable nucleation process. To find the voltage below which the nucleation energy would become negative, we compute the potential of zero nucleation energy (U_{PZN}) for each cluster size (n). The lowest value of $U_{\text{PZN}}(n)$ with respect to the Li reference electrode marks the potential for cluster growth on graphite electrode, below which the Li clusters would grow unbounded on graphite. We have considered the variation with the state of charge, location of nucleation, and the influence of the surrounding electrolyte environment, reaching the following observations:

1. The grand canonical method in electrolyte (gcDFTe) is a significant advancement over conventional canonical DFT in vacuum (cDFTv). For instance, the potential for cluster growth computed with gcDFTe method on the zigzag edge is -0.06 V on

unlithiated graphite and -0.04 V on lithiated graphite. This is close to experimentally observed values of -0.04 V to -0.15 V.¹³ A corresponding value from cDFTv method is -0.31 V on unlithiated graphite and -0.47 on lithiated graphite.²³

2. The consideration of the environment plays an important role in quantifying the thermodynamics of the process. The magnitude of the potential of cluster growth is found to lower be in an electrolyte environment (gcDFTe) compared to vacuum (gcDFTv) for all cases. *E.g.* the potential for cluster growth on the zigzag edge of lithiated graphite in vacuum is -0.14 V as compared to -0.04 V in electrolyte environment. Thus, the nucleation process is more likely to happen in electrolyte environments than what could be predicted from calculations in vacuum.

3. Nucleation is preferred on the zigzag edge termination rather than the basal plane of graphite. *E.g.* the potential for cluster growth on the unlithiated graphite basal plane is -0.12 V while that on the zigzag edge termination is -0.06 V with respect to the Li reference electrode.

4. Nucleation is preferred on the lithiated graphite as compared to the unlithiated graphite. The potential for cluster growth at the zigzag edge is -0.06 V on unlithiated graphite as compared to -0.04 V on the lithiated graphite, while the corresponding value on the basal plane is -0.12 V on unlithiated as compared to -0.08 V on lithiated graphite.

The above observations have profound implications for the occurrence, avoidance and control of dendrite growth in a battery cell through *ab initio* simulations. The resulting thermodynamic parameters will represent the very first step of subsequent kinetic analyses for *in silico* predictions of important results before experimental verification, leading to rapid discovery of new electrolytes and charging procedures to improve safety and lifetime of future cells.

The technologically-important insights obtained from this study to understand the issue of Li nucleation in Li-ion batteries, which is vital for controlling degradation, have been made possible by the development of methods for atomistic simulations of electrodes in an electrolyte environment. The study also demonstrates the usefulness of the grand canonical DFT method in electrolyte (gcDFTe) which is a significant advancement over conventional canonical DFT in vacuum (cDFTv). The gcDFTe method is generally applicable and could be used to model other technologically relevant electrochemical systems such as fuel cells and electrolyzers. The grand canonical method can also be used for molecular dynamics simulations. However, even with a linear-scaling code like ONETEP, which overcomes the length-scale limitations of conventional DFT, we are still limited by the very-short time-scales possible with *ab initio* molecular dynamics. Tackling the time-scale problem would require further developments that could combine more empirical quantum atomistic methods with machine-learning approaches.

Data availability

The data that supports the findings of this study is available within the article.



Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was carried out with the funding from the Faraday Institution (<https://www.Faraday.ac.uk>; EP/S003053/1), grant numbers FIRG003 and FIRG025. The calculations presented in this work were performed on the Iridis5 supercomputer of the University of Southampton, the Michael supercomputer of the Faraday Institution, the Young supercomputer at the UCL and the ARCHER2 UK National Supercomputing Service (<https://www.archer2.ac.uk>; via EPSRC grant: EP/P022030/1). We acknowledge the United Kingdom Materials and Molecular Modelling Hub for computational resources, partially funded by EPSRC (EP/P020194/1).

References

- 1 J. B. Goodenough and K.-S. Park, The Li-Ion Rechargeable Battery: A Perspective, *J. Am. Chem. Soc.*, 2013, **135**, 1167–1176.
- 2 V. Etacheri, R. Marom, R. Elazari, G. Salitra and D. Aurbach, Challenges in the development of advanced Li-ion batteries: a review, *Energy Environ. Sci.*, 2011, **4**, 3243–3262.
- 3 J. B. Goodenough and Y. Kim, Challenges for rechargeable Li batteries, *Chem. Mater.*, 2010, **22**, 587–603.
- 4 H. Zhang, Y. Yang, D. Ren, L. Wang and X. He, Graphite as anode materials: Fundamental mechanism, recent progress and advances, *Energy Storage Mater.*, 2021, **36**, 147–170.
- 5 K. Persson, Y. Hinuma, Y. S. Meng, A. Van der Ven and G. Ceder, Thermodynamic and kinetic properties of the Li-graphite system from first-principles calculations, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2010, **82**, 125416.
- 6 C. Peng, M. P. Mercer, C. K. Skylaris and D. Kramer, Lithium intercalation edge effects and doping implications for graphite anodes, *J. Mater. Chem. A*, 2020, **8**, 7947–7955.
- 7 J. X. Huang, G. Csányi, J. B. Zhao, J. Cheng and V. L. Deringer, First-principles study of alkali-metal intercalation in disordered carbon anode materials, *J. Mater. Chem. A*, 2019, **7**, 19070–19080.
- 8 J. Vetter, P. Novák, M. R. Wagner, C. Veit, K. C. Möller, J. O. Besenhard, M. Winter, M. Wohlfahrt-Mehrens, C. Vogler and A. Hammouche, Ageing mechanisms in lithium-ion batteries, *J. Power Sources*, 2005, **147**, 269–281.
- 9 X. Qin, M. Shao and P. B. Balbuena, Elucidating mechanisms of Li plating on Li anodes of lithium-based batteries, *Electrochim. Acta*, 2018, **284**, 485–494.
- 10 J. Jiao, G. Lai, L. Zhao, J. Lu, Q. Li, X. Xu, Y. Jiang, Y. B. He, C. Ouyang, F. Pan, H. Li and J. Zheng, Self-Healing Mechanism of Lithium in Lithium Metal, *Adv. Sci.*, 2022, **9**(12), 2105574.
- 11 D. Hu, L. Chen, J. Tian, Y. Su, N. Li, G. Chen, Y. Hu, Y. Dou, S. Chen and F. Wu, Research Progress of Lithium Plating on Graphite Anode in Lithium-Ion Batteries, *Chin. J. Chem.*, 2021, **39**, 165–173.
- 12 T. Waldmann, B. I. Hogg and M. Wohlfahrt-Mehrens, Li plating as unwanted side reaction in commercial Li-ion cells – A review, *J. Power Sources*, 2018, **384**, 107–124.
- 13 T. Gao, Y. Han, D. Fragggedakis, S. Das, T. Zhou, C. N. Yeh, S. Xu, W. C. Chueh, J. Li and M. Z. Bazant, Interplay of Lithium Intercalation and Plating on a Single Graphite Particle, *Joule*, 2021, **5**, 393–414.
- 14 P. Arora, M. Doyle and R. E. White, Mathematical Modeling of the Lithium Deposition Overcharge Reaction in Lithium-Ion Batteries Using Carbon-Based Negative Electrodes, *J. Electrochem. Soc.*, 1999, **146**, 3543–3553.
- 15 N. Legrand, B. Knosp, P. Desprez, F. Lopicque and S. Raël, Physical characterization of the charging process of a Li-ion battery and prediction of Li plating by electrochemical modelling, *J. Power Sources*, 2014, **245**, 208–216.
- 16 C. Uhlmann, J. Illig, M. Ender, R. Schuster and E. Ivers-Tiffée, In situ detection of lithium metal plating on graphite in experimental cells, *J. Power Sources*, 2015, **279**, 428–438.
- 17 L. M. Morgan, *et al.*, Pushing the boundaries of lithium battery research with atomistic modelling on different scales, *Prog. Energy*, 2022, **4**, 012002.
- 18 X. Fan, W. T. Zheng, J. L. Kuo and D. J. Singh, Adsorption of single Li and the formation of small Li clusters on graphene for the anode of lithium-ion batteries, *ACS Appl. Mater. Interfaces*, 2013, **5**, 7793–7797.
- 19 M. Liu, A. Kutana, Y. Liu and B. I. Yakobson, First-Principles Studies of Li Nucleation on Graphene, *J. Phys. Chem. Lett.*, 2014, **5**, 1225–1229.
- 20 J. Cui, S. Yao, M. Ihsan-ul haq, J. Wu and J.-k. Kim, Correlation between Li Plating Behavior and Surface Characteristics of Carbon Matrix toward Stable Li Metal Anodes, *Adv. Energy Mater.*, 2019, **9**(1), 1802777.
- 21 X. Chen, X. R. Chen, T. Z. Hou, B. Q. Li, X. B. Cheng, R. Zhang and Q. Zhang, Lithiophilicity chemistry of heteroatom-doped carbon to guide uniform lithium nucleation in lithium metal anodes, *Sci. Adv.*, 2019, **5**(2), eaau7728.
- 22 E. G. Leggesse, C.-L. Chen and J.-C. Jiang, Lithium diffusion in graphene and graphite: Effect of edge morphology, *Carbon*, 2016, **103**, 209–216.
- 23 C. Peng, A. Bhandari, J. Dziedzic, J. R. Owen, C.-K. Skylaris and D. Kramer, Mechanism of Li nucleation at graphite anodes and mitigation strategies, *J. Mater. Chem. A*, 2021, **9**, 16798–16804.
- 24 A. Bhandari, C. Peng, J. Dziedzic, L. Anton, J. R. Owen, D. Kramer and C.-K. Skylaris, Electrochemistry from first-principles in the grand canonical ensemble, *J. Chem. Phys.*, 2021, **155**, 024114.
- 25 A. Bhandari, L. Anton, J. Dziedzic, C. Peng, D. Kramer and C.-K. Skylaris, Electronic structure calculations in electrolyte solutions: Methods for neutralization of extended charged interfaces, *J. Chem. Phys.*, 2020, **153**, 124101.
- 26 J. C. Womack, L. Anton, J. Dziedzic, P. J. Hasnip, M. I. J. Probert and C.-K. D. L. M. G. Skylaris, A Parallel Multigrid Poisson and Poisson-Boltzmann Solver for



- Electronic Structure Calculations in Vacuum and Solution, *J. Chem. Theory Comput.*, 2018, **14**, 1412–1432.
- 27 <http://www.dlmg.org>.
- 28 J. C. A. Prentice, *et al.*, The ONETEP linear-scaling density functional theory program, *J. Chem. Phys.*, 2020, **152**, 174111.
- 29 J. Dziedzic, A. Bhandari, L. Anton, C. Peng, J. C. Womack, M. Famili, D. Kramer and C.-K. Skylaris, Practical Approach to Large-Scale Electronic Structure Calculations in Electrolyte Solutions *via* Continuum-Embedded Linear-Scaling Density Functional Theory, *J. Phys. Chem. C*, 2020, **124**, 7860–7872.
- 30 C.-K. Skylaris, P. D. Haynes, A. A. Mostofi and M. C. Payne, Introducing ONETEP: Linear-scaling density functional simulations on parallel computers, *J. Chem. Phys.*, 2005, **122**, 084119.
- 31 A. A. Mostofi, P. D. Haynes, C.-K. Skylaris and M. C. Payne, Preconditioned iterative minimization for linear-scaling electronic structure calculations, *J. Chem. Phys.*, 2003, **119**, 8842–8848.
- 32 C.-K. Skylaris, A. A. Mostofi, P. D. Haynes, O. Diéguez and M. C. Payne, Nonorthogonal generalized Wannier function pseudopotential plane-wave method, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2002, **66**, 035119.
- 33 S. Grimme, Semiempirical GGA-type density functional constructed with a long-range dispersion correction, *J. Comput. Chem.*, 2006, **27**, 1787–1799.
- 34 G. Bramley, M.-T. Nguyen, V.-A. Glezakou, R. Rousseau and C.-K. Skylaris, Reconciling Work Functions and Adsorption Enthalpies for Implicit Solvent Models: A Pt (111)/Water Interface Case Study, *J. Chem. Theory Comput.*, 2020, **16**, 2703–2715.
- 35 D. S. Hall, J. Self and J. R. Dahn, Dielectric Constants for Quantum Chemistry and Li-Ion Batteries: Solvent Blends of Ethylene Carbonate and Ethyl Methyl Carbonate, *J. Phys. Chem. C*, 2015, **119**, 22322–22330.
- 36 N. Yao, X. Chen, X. Shen, R. Zhang, Z. H. Fu, X. X. Ma, X. Q. Zhang, B. Q. Li and Q. Zhang, An Atomic Insight into the Chemical Origin and Variation of the Dielectric Constant in Liquid Electrolytes, *Angew. Chem., Int. Ed.*, 2021, **60**, 21473–21478.
- 37 J. Dziedzic, H. H. Helal, C.-K. K. Skylaris, A. A. Mostofi and M. C. Payne, Minimal parameter implicit solvent model for *ab initio* electronic-structure calculations, *Epl*, 2011, **95**, 43001.
- 38 R. Naejus, C. Damas, D. Lemordant, R. Coudert and P. Willmann, Excess thermodynamic properties of the ethylene carbonate-trifluoroethyl methyl carbonate and propylene carbonate-trifluoroethyl methyl carbonate systems at $T = (298.15 \text{ or } 315.15) \text{ K}$, *J. Chem. Thermodyn.*, 2002, **34**, 795–806.

