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## Search for active and inactive ion insertion sites in organic crystalline materials

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The position of mobile active and inactive ions, specifically ion insertion sites, within organic crystals, significantly affects the properties of organic materials used for energy storage and ionic transport. Identifying the positions of these atomic (and ionic) sites in an organic crystal is challenging, especially when the element has low X-ray scattering power, such as lithium (Li) and hydrogen, which are difficult to detect by powder X-ray diffraction. First-principles calculations, exemplified by density functional theory (DFT), are very practical for confirming the relative stability of ion positions in materials. However, the lack of effective strategies to identify ion sites in these organic crystalline frameworks renders this task extremely challenging. This work presents two algorithms: the (i) efficient location of ion insertion sites from extrema in electrostatic local potential and charge density (ELIISE), and the (ii) ElectRostatic InsertioN (ERIN), which leverage charge density and electrostatic potential fields accessed from first-principles calculations, combined with the Simultaneous Ion Insertion and Evaluation (SIE) workflow. SIE inserts all ions simultaneously—to determine ion positions in organic crystals. We demonstrate that these methods accurately reproduce known ion positions in 16 organic materials and identify previously overlooked low-energy sites in tetralithium 2,6-naphthalenedicarboxylate (Li<sub>4</sub>NDC), an organic electrode material, highlighting the importance of inserting all ions simultaneously, as in the SIE workflow. These algorithms are also integrated with off-the-shelf machine learning potentials, yielding promising results comparable to first-principles findings.

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## 1 Introduction

Understanding the structure of inorganic and organic-based materials containing electroactive elements, such as lithium (Li), sodium (Na), and zinc (Zn)<sup>1–14</sup> is relevant for the development of the next generation of rechargeable battery technologies, which are prevalent in vehicular transportation, heavy-duty applications, and the supporting electrical grids.

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With the advent of the Rietveld methods,<sup>15–18</sup> diffraction-based techniques using X-ray or neutron sources are commonly employed to determine the structures of energy materials. However, structural determinations through diffraction methods can be hindered by the weak scattering of X-rays by light elements, such as hydrogen and Li, which can partially occupy multiple crystallographic sites because of their high intrinsic mobility. In some cases, significant ion mobilities can cause the blurring of Bragg intensities.<sup>19,20</sup> While neutron diffraction (ND) experiments enhance sensitivity to specific elements, *e.g.*, light elements H and Li, ND experiments are significantly less available than X-ray diffraction analogues. ND still face challenges related to disorder, defects, and temperature-dependent site mixing.<sup>21</sup>

Organic electrode materials (OEMs) are promising for inexpensive and high-energy-density rechargeable batteries. Indeed, OEMs can provide three main benefits: (i) high material-level energy densities of 900–1000 kWh kg<sup>–1</sup>, (ii) a growing range of designs as shown by the variety of molecules studied so far, and (iii) potential benefits in sustainability and supply chains.<sup>6,22–28</sup> One of the main challenges hindering the practical implementation of OEMs is a poor understanding of their structure–property relationships. The electrochemical reaction



pathways of OEMs are typically interpreted based on simplistic molecular models, rather than describing these processes at the material level. These molecular-type OEM models are often paired with *ex situ* or *operando* XRD experiments, which frequently cannot reveal their underlying crystal structures and potential phase transformations.<sup>29–32</sup> To date, only a handful of studies have resolved structures at both the charged and discharged (ion-containing) states of specific OEMs.<sup>23,33–37</sup> For example, 1,4-benzoquinone (BQ) undergoes a phase transition from monoclinic ( $P2_1/c$ ) to orthorhombic ( $P4_2/ncm$ ) upon lithiation or sodiation, forming  $\text{Li}_2\text{BQ}$  or  $\text{Na}_2\text{BQ}$ ,<sup>23,38</sup> indicative of a biphasic reaction with substantial change in the molecular orientation, symmetry and the lattice parameters. By contrast, dilithium 2,6-naphthalene dicarboxylate ( $\text{Li}_2\text{NDC}$ ) retains a monoclinic ( $P2_1/c$ ) structure upon complete lithiation ( $\text{Li}_4\text{NDC}$ ),<sup>34,35,39</sup> following a biphasic intercalation reaction, with little change in molecular orientation and lattice parameters. However, distinguishing specific reaction mechanisms remains poorly understood in the OEM literature, mainly because structural data for the discharged state are scarce. In contrast, structural models for the charged state are often available.

A comprehensive understanding of OEM structures and their behavior across different ion content (*i.e.*, different states of charge) remains crucial for bridging the gap between molecular-scale insights and crystalline materials with intrinsic periodicity. In general, this knowledge is essential for elucidating the physicochemical processes that govern electrochemical behavior, including phase transitions,<sup>23</sup> voltage profiles,<sup>37</sup> and mechanical degradation phenomena.<sup>36,40</sup> However, accurate identification of the positions of mobile ions such as  $\text{Li}^+$ ,  $\text{Na}^+$ ,  $\text{K}^+$ ,  $\text{Zn}^{2+}$ , and  $\text{Mg}^{2+}$ , in organic crystalline frameworks, remains a significant experimental and computational challenge.<sup>29,33,35,36,41–49</sup> This challenge arises primarily from the ample intermolecular space (void within the crystal framework) caused by weak intermolecular interactions, which are responsible for molecular assembly into organic molecular crystals.<sup>50,51</sup> This results in numerous possible cation arrangements within the organic framework. Therefore, developing computational tools that accurately predict ion positions within the crystalline structures of organic materials is essential for breakthroughs and for unlocking hidden potential for rational materials design at the crystal level. While such approaches have been established for inorganic electrode materials,<sup>52–55</sup> predictive search for ion sites in OEMs remains largely confined to the molecular scale and,<sup>46,56</sup> to the best of our knowledge, has not yet been demonstrated for crystalline organic frameworks.

In this paper, we present two complementary computational algorithms and an ion insertion workflow for identifying the optimal active and inactive ion insertion sites within organic frameworks: (i) the Efficient Location of Ion Insertion Sites from Extrema (ELIISE) in electrostatic local potential and charge density, which identifies electrostatically stable ion positions by locating local extrema in electrostatic local potential and electronic charge density (CD) fields. (ii) The ElectRostatic InsertioN (ERIN), which, by following the electrostatic potential, iteratively places ions into electrostatically favorable regions of the unit cell until the entire cell volume is sampled. (iii) The

Simultaneous Ion Insertion and Evaluation (SIIE), which inserts the ions simultaneously and evaluates to determine the correct structure. ELIISE and ERIN, in conjunction with SIIE, are applied with a library of 16 representative organic structures, illustrating their ability to automatically identify candidate ion sites within complex, flexible organic frameworks using limited computational resources. These predictions are subsequently refined using agnostic first-principles calculations, resulting in high-confidence ion positions, which provide crucial insights into ion coordination environments and other structural evidence. These predictive workflows establish a powerful toolkit for the systematic exploration of electroactive structures and their ion transport pathways, elucidating reaction mechanisms and guiding the rational design of materials for the development of competitive organic-based electrochemical devices, as well as the exploration of soft materials for other energy-storage applications.

## 2 Methods

### 2.1. Finding candidate ion insertion sites in organic crystals: ELIISE and ERIN

Fig. 1 shows the identification of candidate ion sites using ELIISE and ERIN, which rely on the electronic charge density

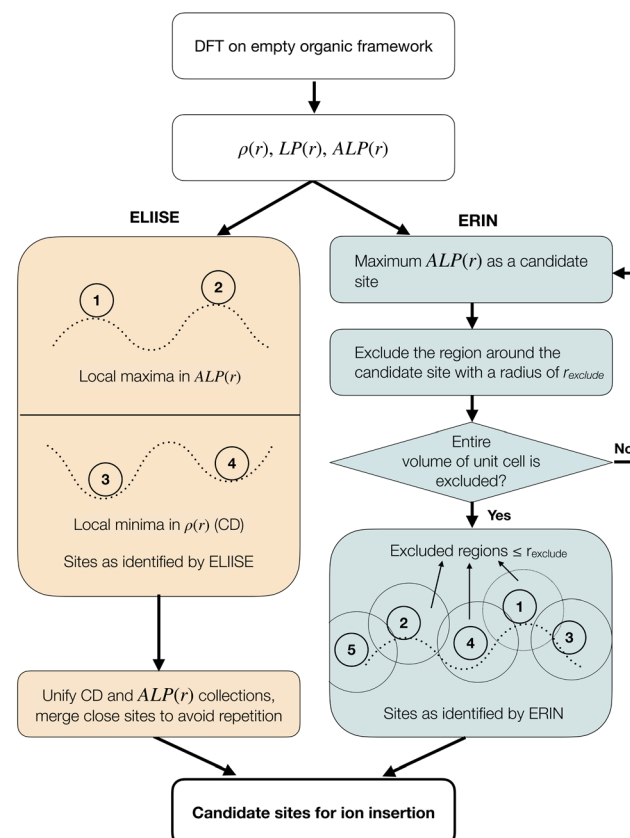


Fig. 1 Workflow and schematic illustration of the ELIISE (left) and ERIN (right) algorithms. ELIISE: process for identifying potential ion sites based on local potential and charge density. ERIN: workflow illustrating the iterative process used to identify potential ion sites based on electrostatic stability systematically.



(CD, or  $\rho(r)$ ) and the local electrostatic potential  $LP(r)$  of eqn (1), and the spherically averaged local potential,  $ALP(r)$ , of eqn (2).

$$LP(r) = V_{\text{hartree}}(r) + V_{\text{ionic}}(r); \quad (1)$$

where  $V_{\text{hartree}}(r)$  is the Coulomb potential set by  $\rho(r)$ , and  $V_{\text{ionic}}(r)$  is the Coulomb potential originating from the ionic core of all atoms. From  $LP(r)$ , the  $ALP(r)$  is defined as:

$$ALP(r) = \frac{\int_{V_s} LP(r + r') d^3r'}{\frac{4}{3}\pi r_s^3}. \quad (2)$$

where  $V_s$  represents the spherical region within radius  $r_s$  that is averaged at each point  $r$ , and  $r'$  is a dummy variable over which the averaging occurs. Note, in eqn (2),  $ALP(r)$  is the spherically averaged electrostatic potential at each point in the organic crystal framework, with constant radii ( $r_s$ ) set by the user (here, fixed to the empirical covalent radii of the atom that is inserted).

Both methodologies rely on an initial organic framework to compute  $LP(r)$  and CD. The choice of the initial organic framework is trivial when the organic framework of the discharged (ion-inserted) structure—*i.e.*, the lattice parameters, the symmetry, and the atomic positions—is known. In such cases, only the precise determination of ion positions is necessary, as is often the case with lithium; here, ELIISE serves as an appropriate tool. In contrast, the more common scenario is when only the organic framework of the charged (ion-removed) phase is known. Under these circumstances, ERIN is recommended (see Discussion section for more details) as employing ELLISE may result in unreliable outcomes.

## 2.2. The efficient location of ion insertion sites from extrema in electrostatic local potential and charge density (ELIISE) method

The ELIISE method identifies candidate ion sites using both the CD and the spherically averaged local potential  $ALP(r)$ , defined in eqn (2), within the empty (*i.e.*, without any mobile species) organic framework from the discharged state.

Following the ELIISE workflow in the left part of Fig. 1, the local potential and electronic CD are initially computed by performing a first-principles calculation without changing the atom positions (or changing the volume or cell shape of the material) of the empty organic framework. Subsequently, ELIISE identifies candidate active sites in an organic framework by identifying the local maxima and the local minima in the  $ALP(r)$  the CD fields, respectively. Fig. 1 shows a schematic diagram of the local minima in CD and local maxima in the  $ALP(r)$ , which are used to identify candidate sites. We have concluded that a spherical average of the  $LP(r)$  provides the best predictions in terms of candidate ion sites, as it accounts for the finite size of inserted ions. Here, the empirical covalent radii of atoms by Slater<sup>57</sup> appear to be a reliable option for defining this ion size. When different types of active species are involved in ion insertion, we imposed the empirical radius of the smaller atom for spherical averaging of the local potential. The candidate sites identified by both the CD and the  $ALP(r)$  descriptors are

combined into a single collection. To avoid repetition of sites, spatially close sites are merged by calculating the geometric mean of their coordinates, as indicated in Fig. 1. The resulting sites are then ordered by their electrostatic stability, from most to least favorable, as determined by the  $ALP(r)$ .

Large values of  $ALP(r)$  indicate regions that are electrostatically favorable for positively charged ions, typically near more electronegative (usually negatively charged) atoms within the crystal structure. Therefore, local maxima in  $ALP(r)$  naturally point to potential cation sites. Similarly, local minima in the CD usually occur in voids near more electronegative atoms. It is reasonable to assume that these electronegative atoms strongly attract electron density, leading to electron-deficient regions nearby, which are likely to form local minima in the charge density. Therefore, these minima in CD often correspond to favorable coordination environments for cations. An exception to this trend becomes evident when local minima occur within large structural voids, which are common in organic crystals and nanoporous materials. These voids may be too far from the organic framework to provide any stabilizing interactions for inserted ions. These situations can lead to surprisingly high Coulomb energies. To address this potential issue during the site search in both ELIISE and ERIN, we set cutoff distance thresholds to exclude regions that are either closer than the specified distance (*e.g.*, 1.5 Å) or farther than ~3 Å from atoms in the organic framework.

## 2.3. The ElectRostatic InsertioN (ERIN) method

In contrast to ELIISE, ERIN only requires the  $LP(r)$  (eqn (1)) of the organic framework, obtained from first-principles calculations. The ERIN approach in the right part of Fig. 1 only uses the spherically averaged local potential  $ALP(r)$  (eqn (2)) to explore electrostatic energy landscapes and systematically generates sparsely spaced candidate ion sites in a decreasing order of electrostatic favorability within the given unit cell. ERIN identifies and selects the site with the maximum  $ALP(r)$  value. Subsequently, ERIN excludes a spherical region within a user-defined radius,  $r_{\text{exclude}}$  (*e.g.*, 2 to 3 Å), around this site from future searches. Then, ERIN repeats the search for the site with the next highest value of  $ALP(r)$  to select as a candidate. This process is repeated iteratively until the entire volume of the unit cell is scanned systematically and excluded. The site search is performed independently within each symmetry-distinct region of the unit cell to identify candidate sites in all possible Wyckoff positions.

## 2.4. Simultaneous ion insertion and evaluation (SIEE) workflow for candidate site insertion

Once ELIISE or ERIN identifies the candidate sites, the next step is to determine the optimal combination of sites that yield the expected stoichiometry of the system and is energetically viable. For this purpose, we introduce the Simultaneous Ion Insertion and Evaluation (SIEE) workflow, in which ions are introduced simultaneously within the organic framework to achieve the desired stoichiometry, as opposed to the sequential insertion method, where ions are inserted one at a time.<sup>52,58</sup> In the SIEE



workflow, if the organic framework of the discharged structure is known, its atoms are kept fixed during ion insertion, with only inserted ions and decorating hydrogen atoms (whose positions are typically unknown from standard XRD experiments) allowed to change. Otherwise, if only the organic framework of the charged phase is known, then both the ions and the organic framework are allowed to relax completely.

As shown in Fig. 2a, SIIIE uses the candidate ion insertion sites from the ELIISE or ERIN methods discussed in the previous sections as inputs. While ELIISE and ERIN provide sets of candidate ion positions, they do not determine the final configuration of ions in the discharged structure. In real systems, multiple symmetry-distinct sites may be occupied simultaneously, and therefore, the correct discharged state often results from a specific combination of these candidate sites.

Finding the right combination of candidate sites is challenging due to the large number of unique possibilities. Note that the number of possible combinations of predicted ion sites that produce the target ion content can grow combinatorially, in the range  $10$  to  $10^6$ , which typically makes direct DFT evaluation of all combinations computationally infeasible.

To address this challenging task, we incorporate a machine learning interatomic potential (MLIP), such as MACE or Orb-v3,<sup>59–61</sup> to screen thousands of configurations efficiently. Here, we use MACE to identify and remove high-energy structures. More details on how MACE and DFT are combined in the SIIIE workflow are provided later (see Computational details section).

## 3 Results

### 3.1. Case studies: implementing the suggested workflows on example systems

To demonstrate the effectiveness of these algorithms, we showcase the application of the ELIISE (ERIN) + SIIIE workflow to several model organic compounds. The accuracy of ELIISE +

SIIIE is verified by “rediscovering” the Na-ion sites in two known fully sodiated compounds: (i) the disodium 1,4-benzoquinone ( $P4_2/ncm$ ,  $Na_2BQ$ ),<sup>38</sup> and (ii) the tetra-sodium rhodizonate with formula  $Na_4C_6O_6$  ( $C2/m$ ).<sup>62</sup> Similarly, starting from the known dilithium 2,6-naphthalenedicarboxylate ( $P2_1/c$ ,  $Li_2NDC$ ), using ERIN + SIIIE, we predict Li positions in  $Li_4NDC$  (tetra-lithium 2,6-naphthalenedicarboxylate),<sup>35,63</sup> which contains two additional Li sites per formula unit (f.u.).

$Na_4C_6O_6$  ( $C2/m$ )<sup>62</sup> is an oxocarbon-salt-based layered compound; it is a planar conjugated ring with six carbonyl oxygens, known to undergo a two-to-four electron reduction with Na.<sup>36</sup> To test the ELIISE method, we first remove all Na atoms from  $Na_4C_6O_6$  and perform a static DFT calculation, at the volume and geometry available in literature, to calculate the electronic charge density and local electrostatic potential fields, and subsequently apply the ELIISE algorithm to find the possible sites for Na-ions (as described in Methods section 2.2).

The ELIISE algorithm yields nine symmetry-distinct Na sites with different multiplicities (Table S2). These candidate sites from ELIISE are then “funneled” into the SIIIE workflow. We examine all possible combinations of sites from these nine symmetry-distinct candidates that produce the desired  $Na_4C_6O_6$  stoichiometry. This yields only 59 unique sodium configurations, for which we perform DFT and MACE optimizations to identify the most plausible Na-atom positions. To quantify the prediction accuracy of ELIISE + SIIIE, we use a metric called the maximum atom displacement error (MADE), which is defined in eqn (3):

$$MADE = \max \{ |r_i - r_i^{\text{ref}}|; i \in \text{all active, inactive species} \} \quad (3)$$

where  $r_i$  is the predicted ion position of the  $i$ th ion, and  $r_i^{\text{ref}}$  is the position of the corresponding ion from the literature (*i.e.*, from experimentally determined structures). The predicted and literature ion pairs are matched based on the pairing that

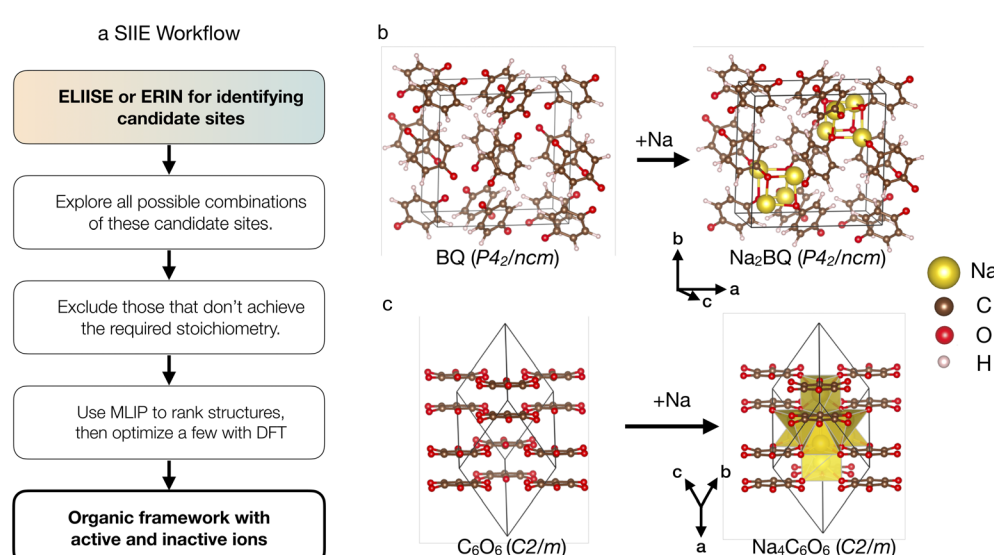


Fig. 2 Workflow for the Simultaneous Ion Insertion and Evaluation (SIIIE) method (a) and its application in two OEMs (b) disodium 1,4-benzoquinone ( $Na_2BQ$ ) and (c) tetrasodium rhodizonate ( $Na_4C_6O_6$ ). Combining SIIIE with ELIISE, the Na-ion positions are located in two exemplary OEMs.<sup>38,62</sup>



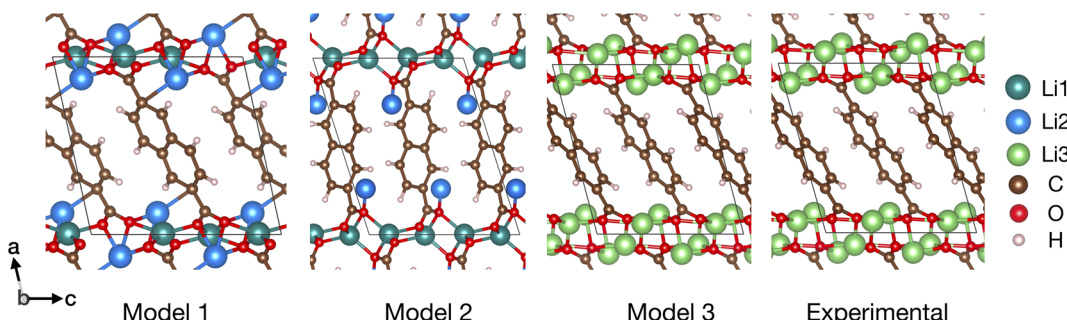


Fig. 3 Li<sub>4</sub>NDC structures: model 1, model 2, model 3, and experimentally refined from single-crystal XRD.<sup>35</sup> Different crystallographic Li sites are shown.

minimizes the sum of their absolute deviations. Thus, MADE (eqn (3)) is the maximum displacement between any predicted ion position and its corresponding literature reference, *i.e.*, the actual atom positions reported in the literature.<sup>62</sup> MADE is defined when the organic framework of the discharged structure is known and remains fixed during the optimization of ionic positions. MADE assesses the accuracy of predicted sites by quantifying the extent to which ions deviate from the positions reported in the literature.

In practice, we interpret MADE values  $\leq \sim 0.4$  Å as indicating correct predictions for all ions in the structure, while MADE values  $\geq \sim 0.4$  Å suggest that at least one ion site has been predicted incorrectly. Among the predicted structures for Na<sub>4</sub>C<sub>6</sub>O<sub>6</sub>, the lowest-energy structure of Fig. 2c displays a MADE of  $\sim 0.026$  Å, indicating excellent agreement with the structure previously reported in the literature.<sup>62</sup>

In the second example, we investigate the structure of di-sodium 1,4-benzoquinone, Na<sub>2</sub>BQ<sup>38</sup> (Fig. 2b), and the fully reduced (sodiated) form of a *para*-benzoquinone, *i.e.*, BQ. In Na<sub>2</sub>BQ, the Na atom positions are available from the literature.<sup>38</sup> In Fig. 2b, the Na<sub>2</sub>BQ features a highly open organic framework with large voids, making a brute-force search of the potential Na positions computationally expensive. Using ELIISE, we identify four symmetrically distinct candidate sites. Applying the SIEE workflow, we evaluate all combinations of these four symmetry-distinct ions that produce the stoichiometry of Na<sub>2</sub>BQ. This step yields only three valid configurations, which we used for further DFT structural relaxations (keeping the volume and shape fixed to the experimental structure of Na<sub>2</sub>BQ). The most stable structure with the lowest DFT total energy singled out with ELIISE + SIEE, shown in Fig. 2b, exhibits a MADE of  $\sim 0.024$  Å, demonstrating excellent agreement with the experimental Na positions.<sup>38</sup>

We tested our methods on a third structure, Li<sub>4</sub>NDC, which shares structural similarities with its precursor, Li<sub>2</sub>NDC,<sup>64</sup> with 2 Li atoms per molecule. Prior experimental studies indicated that the NDC organic framework largely remains unchanged upon reaction with Li. Two structural arrangements of Li species, model 1 and model 2 (Fig. 3), have been proposed for Li<sub>4</sub>NDC.<sup>63</sup> Models 1 and 2 preserve Li-ions in tetrahedral coordination sites as observed in Li<sub>2</sub>NDC<sup>64</sup> (shown as Li1 in Fig. 3)

and introduce new Li positions (shown as Li2 in Fig. 3), approaching composition Li<sub>4</sub>NDC (four Li atoms per molecule).

Using ERIN with SIEE workflow, we predict the Li positions in Li<sub>4</sub>NDC based on the organic framework of Li<sub>2</sub>NDC as input. We recovered models 1 and 2, described in the literature. A close inspection of model 1 and model 2 reveals that these two structures reported in ref. 63 are different in the relative orientation of NDC molecules, as displayed in Fig. 3. However, this striking contrast is not highlighted in ref. 63, which suggests that model 1 and model 2 differ only in Li distribution but share the same molecular framework.

In addition to models 1 and 2, we identified a third model, model 3, as the lowest-energy structure, as determined by DFT minimization (and MACE). In model 3, the tetrahedral sites reported for Li<sub>2</sub>NDC are unoccupied, and all the Li atoms assume new crystallographic positions (collectively indicated as Li3 in Fig. 3), which are localized near the inorganic layer of the organic framework, consisting of oxygen atoms in NDC. These newly identified Li-atom positions in model 3 yield an electrostatically favorable arrangement of Li atoms relative to models 1 and 2 at the same composition. Quantitatively, DFT total-energy comparisons indicate that model 3 is energetically favored over models 1 and 2 by  $\sim 100$  meV per Li<sub>4</sub>H<sub>6</sub>C<sub>12</sub>O<sub>4</sub> and  $\sim 440$  meV per Li<sub>4</sub>H<sub>6</sub>C<sub>12</sub>O<sub>4</sub>, respectively.

Indeed, model 3 is in excellent agreement with a recent study (appeared while drafting this paper) that used single-crystal XRD to refine the structure of Li<sub>4</sub>NDC.<sup>35</sup> Because ERIN uses the organic framework of the charged state as input and optimizes all atomic positions, the MADE metric—measuring the prediction error in active species positions—appears unsuitable. Instead, a simple qualitative measure of similarity between two structures can be obtained by using the StructureMatcher (with the default parameters) as implemented in the Pymatgen library.<sup>65</sup> Here, StructureMatcher indicated that model 3 qualitatively matches the recently reported experimental structure of Li<sub>4</sub>NDC.<sup>35</sup> Alternatively, Table S4 shows a close match between the lattice parameters and atomic positions of experimental and predicted structures.

These three case studies, discussed in the previous paragraphs, collectively demonstrate the robustness of the ELIISE and ERIN methods when integrated with the SIEE workflow in



accurately identifying ion positions across various organic electrode materials.

### 3.2. Assessing the accuracy of the ELIISE + SIIIE approach

To assess the accuracy of the ELIISE + SIIIE approach, we apply it to a diverse set of organic materials (16 in total) for which the organic framework and ion positions of the crystal structure are known from literature. Using MADE (eqn (3)) as a similarity metric we evaluate the performance of the ELIISE method with the SIIIE workflow across 16 organic systems. These structure organic electrode materials, such as  $\text{Li}_2\text{NDC}$  and  $\text{Li}_4\text{NDC}$  (di and tetra lithium 2,6-naphthalene dicarboxylate),<sup>64</sup>  $\text{Li}_2\text{BDC}$  and  $\text{Li}_4\text{BDC}$  (di and tetra lithium 1,4-benzene dicarboxylate),<sup>66</sup>  $\text{Li}_2\text{BPDC}$  (dilithium biphenyl dicarboxylate),<sup>67</sup>  $\text{Li}_2\text{Mg-}p\text{-DHT}$  (magnesium(2,5-dilithium-oxy)-terephthalate),<sup>33</sup>  $\text{Na}_2\text{BQ}$  (di-sodium hydroquinone),<sup>38</sup>  $\text{Na}_2\text{C}_6\text{O}_6$  (disodium rhodizonate),<sup>68</sup> and  $\text{Na}_4\text{C}_6\text{O}_6$  (tetra sodium rhodizonate),<sup>62</sup> as well as other selected structures (see Table S1) were obtained from crystallographic databases, the Cambridge Crystallographic Data Centre (CCDC)<sup>69</sup> and the Inorganic Crystal Structure Database (ICSD).<sup>70</sup>

As shown in Fig. 4a, the lowest-energy structure identified by the SIIIE workflow using ELIISE correctly reproduces the known ion positions in all tested cases, as measured by MADE, which is always  $\leq 0.4$  Å. Furthermore, Fig. 4b demonstrates that using a pre-trained machine-learning interatomic potential (MLIP), specifically MACE (used only to optimize the ion positions), yields correct predictions of ion positions, with MADE values consistently below  $\sim 0.4$  Å. This suggests that DFT-level structural relaxation may not be strictly necessary to determine the general location of inserted ions, thereby substantially reducing computational costs. Fig. S1 presents a similar assessment, but with predictions generated using ERIN + SIIIE instead of ELIISE + SIIIE, and only MACE was employed for optimizing ion

positions. The MADE values in Fig. S1 show that ERIN + SIIIE also accurately predicted the positions of all inserted ion sites for all 16 tested organic systems.

## 4 Discussion

Predicting ion positions in “soft” crystalline materials, such as organic electrode materials, remains a significant challenge due to the combination of large free volumes, weak intermolecular forces, and the dynamic nature of ion coordination. In this paper, we have developed two new methodologies, ELIISE and ERIN, and an ion insertion workflow, SIIIE, designed to identify ion-insertion sites in organic structures. These include metal atoms, such as Na, Zn, and Mg, as well as light elements, for example, Li. Here, we discuss the advantages and disadvantages of implementing the ELIISE and ERIN algorithms to resolve unknown ion positions in crystal structures, where traditional structural techniques fail.

ELIISE uses both the electronic charge density (CD) and the local electrostatic potential  $\text{LP}(r)$  derived from first-principles calculations of the empty organic framework. In combination, the CD and the  $\text{LP}(r)$  are used to find candidate crystallographic sites for light elements in these materials. However, since the CD and the  $\text{LP}(r)$  are intrinsically linked (as the latter is derived from atomic positions and the electronic charge density distribution), the independent use of the CD and the  $\text{LP}(r)$  often arrives at similar predictions for candidate ion sites. Since ELIISE directly identifies coordinating environments—regions surrounded by electronegative atoms in the organic frameworks—ELIISE appears most effective for systems in which the organic framework structure of the discharged (ion-inserted) phase is known. This signifies that prior experiments determining the space group, lattice constants, and atom positions of the organic framework (excluding the mobile ions) must be

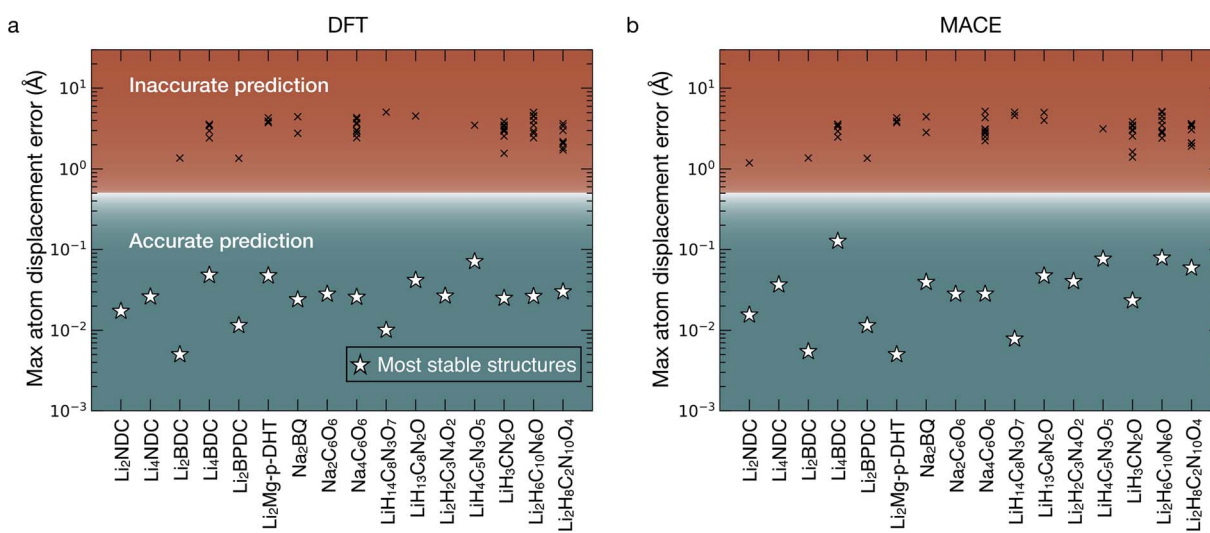


Fig. 4 Accuracy of the predicted active ion positions with the proposed SIIIE workflow and ELIISE. MADE (eqn (3)) values determined after optimization of ionic positions with (a) first-principles DFT and (b) the machine learned potential MACE. Stars indicate the most stable (lowest total energy) ion arrangements in each organic structure, and crosses indicate the thermodynamically unfavorable (high energy) structures. Details of reference structures reported in the literature are in Table S1.



available. In some sense, this limits the capabilities of the ELIISE strategy. For example, if only the charged-phase structure of an organic system is available and significant structural changes and/or phase transitions are expected upon electrochemical reactions, using ELIISE may yield unreliable results.

To address the apparent limitation of ELIISE, we have developed an alternative method called ERIN. ERIN generates a sparse set of points that cover all possible Wyckoff sites, ranked from the electrostatically lowest-energy sites to the highest-energy sites. Therefore, ERIN should be applied when the structure of the organic framework, whose ion-inserted/discharged phases remain unknown. In such cases, the charged (empty) structure of the materials can be used as a starting point, assuming the unit cell of the pristine organic crystal shape and most of the symmetry remains unchanged or structural modifications occur gradually (for example, a displacive phase transition) upon reaction with Li (or sodium), which is typical of intercalation materials.

ELIISE identifies stationary sites (*i.e.*, local minima) by populating the energy landscape; these sites are expected to be stationary points for ions in the potential created by the anion framework. In contrast, in ERIN, the identified sites are not necessarily located at energy minima – non-stationary sites, such as saddle points, on the potential energy surface, and may be mobile during structural relaxation. These non-stationary sites can perturb the organic framework during relaxation and promote structural changes that lead to a more stable discharged structure.

Even if the discharge phase does not involve phase transitions, the specific coordination environments and lattice parameters required to accommodate mobile metal atoms in the discharged organic framework may not be available in the charged organic framework. ERIN systematic electrostatic screening helps identify plausible insertion sites that would be hard to locate with ELIISE alone. Note, however, that ERIN predicts more candidate sites than ELIISE, resulting in a greater number of possible ion-vacancy arrangements to evaluate in the SIEE workflow.

Our study has focused on identifying plausible sites for cation insertion; however, with suitable modifications, these algorithms can also be used to locate anion positions in a crystal structure. For example, in the case of the ELIISE method, local minima in the  $\text{ALP}(r)$  should be used instead of local maxima. The approach remains the same when using CD. In ERIN, instead of selecting the maximum in the  $\text{ALP}(r)$  as a candidate site, the minimum in the  $\text{ALP}(r)$  should be chosen.

At its core, ERIN uses the Coulomb electrostatic potential to model the energy landscape and identify sites generated from electronic charge densities obtained from first-principles calculations. Other methods, such as machine learning interatomic potentials<sup>61,71,72</sup> or bond valence sums<sup>73</sup> appear viable for modeling and developing the potential energy landscape for ion insertion.

It is important to comment on some choices made in developing the averaged local potential,  $\text{ALP}(r)$ , which uses a spherical averaging radius set in advance. We have observed that ion positions predicted from  $\text{ALP}(r)$  are generally

insensitive to the precise value of this radius. However, because of the radius ionic size of the predicted sites, even a  $\sim 10\%$  change can modify the number of local extrema in  $\text{ALP}(r)$  and, consequently, the set of candidate sites identified by ELIISE. For ERIN, changes in the spherical averaging radius directly affect the predicted site positions by altering the underlying energy landscape. This sensitivity is not problematic, as ERIN is intended to generate a sparse set of representative ion sites rather than an exhaustive list. The exclusion radius in ERIN determines the total number of predicted ion sites. Increasing the radius reduces the number of distinct sites by merging nearby candidates. Thus, tuning the radius minimizes the number of ion combinations, increasing computational efficiency in the search. However, because of the MACE and DFT optimization steps, minor deviations from reasonable values are unlikely to affect the accuracy of ion-position predictions.

In general, when testing the ELIISE and ERIN algorithms on 16 previously known structures (Fig. 4 and S1), we demonstrated that these algorithms, in combination with the SIEE workflow, provide a reliable and systematic approach for accurately predicting unknown ion positions in organic frameworks.

Another important aspect is the integration of machine learning interatomic potentials (MLIPs), such as MACE, into the SIEE workflow. In principle, ERIN and ELIISE are applicable together with any foundational MLIP model, including M3GN<sub>ET</sub>, CHGNET, and ORB-3, among others.<sup>61,71,72</sup> This hybrid MLIP-DFT approach appears to be viable, significantly reducing the computational burden of exploring vast combinatorial site occupancies while providing sufficient accuracy in predicting ion positions. Furthermore, the ability of MLIPs to screen thousands of candidate configurations before subsequent, more accurate but computationally intensive refinements with first-principles calculations highlights a scalable pathway for high-throughput exploration of ion-insertion chemistries in soft systems, such as organic crystals, porous systems, metal-organic frameworks, and larger molecules with biological relevance.

In this vein, we have tested MACE predictions against DFT predictions for the same systems shown in Fig. 4. We have observed that, when the organic framework is fixed at the known discharged structure, identifying the structure with the lowest energy and correct ion arrangements using (MADE) with a prediction error within a tolerable range of 0.4 does not require DFT; foundational out-of-the-shelf MLIPs appear suited for this task. Here, we have only tested MACE.<sup>59,60</sup> However, fine-tuning the foundational MLIPs with system-specific DFT data can provide even more accurate results.

The SIEE workflow appears limited when dealing with either amorphous structures or extremely large unit cells with low or no symmetry. In these cases, the number of possible ion combinations can be in the order of  $10^4$  to  $10^6$ , requiring significant computational resources to evaluate total energies across all configurations, even with MACE. In this context, we had relied on the fact that candidate sites are ranked in decreasing order of electrostatic favorability, as indicated by the averaged local potential. Unfavorable sites are systematically excluded during the generation of ion combinations; this



removal process continues until the number of valid ion combinations falls below the preset threshold ( $\sim 10^3$ ). Developing more efficient yet accurate machine-learning potentials is a promising approach to address this issue in amorphous compositions, and considerable effort is currently devoted to this area.<sup>74–77</sup>

Aside from ELIISE and ERIN, the simultaneous insertion of ions, implemented in the SIE workflow, is also essential for accurately finding discharged structures, as illustrated by the case study of Li<sub>4</sub>NDC. In Li<sub>4</sub>NDC, we proposed model 3, which yields the lowest DFT total energy (Fig. 3), and is lower than that of the previously proposed models 1 and 2.<sup>63</sup> Before this work,<sup>55,63</sup> other strategies, such as sequential insertion methods, have been proposed.<sup>52,58</sup> In the sequential approach, a new model is created by inserting an atom at a new position within the unit cell, and the structure is then optimized. This process is repeated until the desired stoichiometry is achieved, thereby identifying a representative global minimum. The sequential approach can lead to incorrect selection of stable candidates because ions may become trapped in local minima of the potential energy surface.

Organic molecular crystals often exhibit polymorphism, resulting from different packing arrangements of the same molecule, including metastable ones.<sup>78,79</sup> However, once the organic framework is established, ions are expected to occupy the most energetically favorable positions, as observed in all cases shown in Fig. 4, thus partially ignoring metastable effects. Therefore, the proposed workflows are effective when the discharged organic frameworks are known. When the discharge state is unknown, as in Li<sub>4</sub>NDC (where only the charged phase is known) lattice constants, crystal shape, and positions are allowed to vary during ion insertion. This can cause significant structural changes and different packing arrangements, potentially leading to the prediction of metastable or energetically favorable phases.

Current methods are limited because they either rely on the organic framework of the discharged structure or assume that the symmetry and unit cell of the discharged phase match those of the charged phase. In the case of Li<sub>4</sub>NDC, determining the optimal ion positions, starting from Li<sub>2</sub>NDC, was straightforward because the organic framework remained essentially unchanged during lithiation, *i.e.*, the symmetry and unit cell of Li<sub>4</sub>NDC were commensurate with those of Li<sub>2</sub>NDC.

However, this is not the case for most systems. For example, when Li<sub>2</sub>BDC is lithiated to form Li<sub>4</sub>BDC, a symmetry reduction is observed.<sup>35</sup> It involves doubling the unit cell along the stacking direction of the organic layers, accompanied by changes in the organic framework.<sup>35</sup> Consequently, attempting to predict ion positions in Li<sub>4</sub>BDC by starting directly from the Li<sub>2</sub>BDC structure may produce an incorrect model structure. Na<sub>2</sub>-BQ and Na<sub>4</sub>C<sub>6</sub>O<sub>6</sub> exemplify this type of structural change, undergoing phase transformations with altered unit cells (and symmetries) during discharge. Beyond the organic molecular crystals studied here, ion insertion has been shown to induce additional distortions, including octahedral tilting, phase separation, cell collapse, loss of crystallinity, and even phase

changes, as observed in hybrid organic–inorganic metal halides and redox-active metal–organic frameworks.<sup>80–85</sup>

These cases highlight a broader challenge: it is often unclear whether a significant phase transition occurs during lithiation (or sodiation). When this uncertainty arises, a practical approach is first to predict ion positions based on the charged structure (*i.e.*, the host in its oxidized or delithiated form). The resulting model can then be compared with experimental data, such as XRD or ND; if the simulated patterns are similar to the experimental patterns, the predicted structure can serve as a solid starting point for refinement.

However, if the comparison shows significant differences, it becomes necessary to explore alternative structural hypotheses. In such cases, one must generate and assess a set of candidate organic framework motifs to serve as initial models in the ERIN + SIE workflow. Developing systematic methods for constructing and testing these organic framework variations appears crucial for reliably understanding the structural complexity of ion insertion in molecular crystals.

The issue of significant structural rearrangement or symmetry breaking is directly related to the problem of locating the global minimum on a structure's potential energy surface; for this, all possible structures must be examined, which entails solving global optimization problems. This goes beyond the simple task of determining ion positions discussed in the current work. Several strategies for identifying global minima in materials space have been proposed, including genetic algorithms, as implemented in USPEX,<sup>86–88</sup> and particle swarm optimization, as implemented in CALYPSO.<sup>89–91</sup> Recently, generative models aided by machine learning have emerged as an effective avenue for efficiently exploring complex potential energy landscapes and chemical spaces, such as those of organic/molecular crystals.<sup>92–95</sup> These strategies are claimed to predict the positions of the organic framework and ions, thereby enabling a complete materials structure model.

For systems that amorphize during discharge, a realistic description of ion positions will ultimately require combining our structure-generation strategy with total-scattering measurements, melt-quench-type molecular dynamics (with machine learning), and possibly reverse Monte Carlo approaches to produce chemically meaningful, though not uniquely determined, models of the disordered Li environments.<sup>96–103</sup>

Besides XRD and ND for determining the structures of organic materials, solid-state nuclear magnetic resonance (ss-NMR) crystallography is another valuable tool that offers structural insights.<sup>104,105</sup> The nascent application of ss-NMR in conjunction with first-principles calculations, termed NMR crystallography, has been used to establish the structure of functional materials.<sup>106</sup>

## 5 Conclusions

In summary, we have developed a computational framework for accurately predicting ion positions in organic materials, utilizing DFT-derived electrostatic and charge density fields with the ELIISE and ERIN algorithms and the SIE workflow.



Through case studies of  $\text{Na}_4\text{C}_6\text{O}_6$ ,  $\text{Na}_2\text{BQ}$ , and  $\text{Li}_4\text{NDC}$ , we showed that ELIISE and ERIN methodologies can reproduce ion positions known from the literature with sub-angstrom accuracy. Additionally, we demonstrate that these methodologies can uncover new, energetically favorable structural models that can improve our understanding of experimental observations.

We have demonstrated that computationally intensive DFT predictions are not a strong requirement for accurately identifying ion positions, and we have shown that machine-learned potentials provide sufficient accuracy for large-scale screening tasks.

Our results demonstrate that ELIISE and ERIN, combined with SIEE, are powerful and can significantly improve, if not augment, the limits of traditional structural characterization techniques. The framework proposed in this paper offers a pathway toward deeper mechanistic insights and the rational design of organic materials for next-generation functional materials. Moreover, MLIPs can efficiently screen thousands of candidate configurations initially, before performing more precise but computationally expensive refinements with first-principles methods. This approach offers a scalable pathway for high-throughput exploration of ion-insertion chemistries in soft systems, including organic crystals, porous materials, metal-organic frameworks, and larger biologically relevant molecules.

## 6 Computational details

All first-principles calculations presented in this work were performed using the density functional theory formalism, as implemented in the Vienna *Ab initio* Simulation Package (VASP).<sup>107–109</sup> The PAW potentials describe the core electrons. Perdew, Burke, and Ernzerhof (PBE)<sup>110</sup> was used to approximate the unknown DFT exchange and correlation XC functional. To account for van der Waals interactions, the empirical D3 method proposed by Grimme and collaborators, with Becke-Johnson (BJ) damping, was used.<sup>111,112</sup>

VASP inputs prepared for geometry optimization and energy calculations closely followed the MITRelaxSet<sup>113</sup> as available in pymatgen.<sup>65</sup> The kinetic energy cutoff for the plane waves was set to 520 eV, and the total energy was converged to  $10^{-5}$  eV per cell. Geometries (coordinates, volumes, and cell shapes) were considered converged when the forces on all atoms are lower than  $0.05 \text{ eV } \text{\AA}^{-1}$ . A  $\Gamma$ -centered Monkhorst-Pack<sup>114</sup> grid with a density of 25  $k$ -points per  $\text{\AA}^{-1}$  for all systems. With these DFT settings, local potentials are computed as the sum of Ewald and Hartree parts, excluding the Exchange and Correlation (XC) contributions.

To reduce the number of structures for DFT optimization, MACE-MATPES-PBE-0 a machine-learned foundational model, based on MACE,<sup>59,60</sup> with PFP-based PyTorch implementation of DFT-D3,<sup>111,112,115</sup> was used to evaluate static total energy and/or optimize ion positions. Structural relaxations are carried out until the forces are converged below  $0.03 \text{ eV } \text{\AA}^{-1}$ .

When MACE is implemented in the SIEE workflow, it is used as follows:

(1) Generate all possible ion combinations. If the number of combinations exceeds a preset limit ( $\sim 10^3$ ), they can be reduced systematically by excluding electrostatically unfavorable sites in ELIISE or ERIN before generating possible ion combinations.

(2) Use MACE to evaluate all the generated combinations.

(3) Select approximately  $\sim 10^2$  of the lowest-energy candidate structures for optimization of ion positions using MACE.

(4) From these structures, we select approximately 10 distinct lowest-energy structures for subsequent accurate DFT optimization of the atomic positions.

This hybrid approach, which combines the computational speed of MACE-based MLIPs with the accuracy of DFT, enables us to successfully identify the correct atomic configurations in organic frameworks with substantial structural complexity.

## Author contributions

H. R. G.: conceptualization, data curation, formal analysis, investigation, methodology, validation, visualization, writing – original draft, writing – review & editing. A. E. L.: data curation, discussion, writing – original draft, writing – review & editing. A. A. P.: discussion, writing – review & editing. Y. C.: discussion, writing – review & editing. V. S. C. K.: discussion, writing – review & editing. L. J.: discussion, writing – review & editing. J. W.: discussion, writing – review & editing. Y. C.: discussion, writing – review & editing. K. W.: discussion, writing – review & editing. M. K. Y. C.: discussion, writing – review & editing. Y. Y.: funding acquisition, supervision, discussion, writing – review & editing. P. C.: conceptualization, funding acquisition, investigation, project administration, resources, supervision, visualization, writing – original draft, writing – review & editing.

## Conflicts of interest

Y. Y. has equity interests in LiBeyond, LLC and Solid Design Instruments, LLC. The University of Houston reviewed and approved his relationship in compliance with its conflict-of-interest policy. The remaining authors declare no competing interests.

## Data availability

Computational data related to this study, including Python scripts, input, and output files of the simulations, are available on GitHub at [https://github.com/caneparesearch/data\\_site\\_finder](https://github.com/caneparesearch/data_site_finder).

Supplementary information (SI) is available. See DOI: <https://doi.org/10.1039/d5sc07602a>.

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