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## Solid–electrolyte–interphase design in constrained ensemble for solid-state batteries†

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Solid-state-batteries (SSBs) represent one of the most promising directions in the energy-storage field. The development of SSBs, however, is currently limited by the complex [electro-]chemical reactions that inevitably occur at the interface of solid-state electrolyte (SSE) particles. Moreover, given the material complexity of such systems, there is no straightforward methodology for addressing these interface instabilities. In this work, a combined high-throughput *ab initio* computation and machine learning approach is used to study and design solid-state solid–electrolyte–interphase (SEI) with tunable electrochemical stabilities using our unique constrained ensemble description. Machine learning reveals that the ability of a solid-state SEI to be stabilized by the mechanical constriction effect is a nonconvex and nonlinear, but deterministic none-the-less, function of composition. The power of this approach is demonstrated using the interface of glass and ceramic sulfide families of solid-electrolytes. Finally, it is experimentally verified that the designed interfaces, in fact, decompose and electrochemically passivate based on our predictions.

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### Broader context

Solid-state batteries are widely seen as the next great step in energy storage technology to displace lithium-ion batteries as the gold standard of secondary batteries. However, the development of solid state batteries has remained slow, largely due to complicated [electro-]chemical reactions that occur at interfaces between solid phases. Thus, there is an urgent need to develop a more effective approach to handle this major design hurdle. In this work, we introduce a comprehensive method to systematically study and, ultimately, design solid–electrolyte–interphases for such an all-solid electrochemical system. Within the general framework of mechanical constriction design, a new concept of critical effective modulus of interphase is proposed and quantified. Any local mechanical constriction level below this critical value will allow decompositions to happen. The decomposition, however, will fill local voids and cracks, and eventually increase the local mechanical constriction beyond the critical value to suppress further decompositions through metastability. Therefore, from this unique perspective of interphase electrochemical evolution, designing the interphase with a reduced critical effective modulus becomes an effective approach to increase dynamic interface stabilities. The method can be applied to the interface design of broad types of material systems, including glass, ceramic, polymer and their composites.

The rapid charging, high energy density, and non-flammability of SSBs<sup>1–8</sup> are critical for the ongoing push to achieve 100% vehicle electrification – a necessity for meaningful progress on climate change. Despite this promise, there remain many technical barriers in the way of broadly adoptable SSBs. One of the greatest of which is the electrochemical instability often found at the various interfaces in SSB cells.<sup>9–13</sup>

In our previous work, we have detailed how the fundamental thermodynamics of materials deviate in SSBs as compared to conventional liquid-cells.<sup>11–13</sup> In short, liquid-cells operate in

accordance with the well-studied state variables of pressure ( $p$ ), temperature ( $T$ ), and voltage ( $\phi$ ). In other words, the operator of a conventional cell sets these state variables experimentally and the material phases equilibrate (by, for example, [de]lithiation). Thus, the conventional liquid-cell can be well described by the  $pT\phi$  thermodynamic ensemble, which we term the “unconstrained” ensemble for contrast with the “constrained” ensemble for all-solid-state systems, described below.

Our previous work has shown that, unlike liquid systems, pressure is not a valid state variable in SSBs. In other words, the experimenter does not have direct control of the pressure within a working cell. When chemical decompositions occur within SSBs, strong local pressure gradients can form as the pressure is a function of the phases which are present locally. In liquid-cells, the liquid–electrolyte would flow to equilibrate this local pressure in response to any external pressure applied

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at the cell level. This cannot occur in an SSB, where the SSE maintains a pressure gradient. Thus, an operator has no direct control of the local pressure within an SSB. As a result, the electrochemical stability of solid-electrolytes varies considerably depending on cell and material design.

To describe these profound thermodynamic differences in SSBs, we introduce the concept of “constrained” ensemble, where the pressure is no-longer a state variable but is a function of the mole-fraction of local phases ( $\{x_i\}$ ). There are two critical insights for understanding this change of state variables. The first is the observation that many solid-electrolytes tend to decompose to products that are larger in total volume than the original electrolyte. The second is that, unlike liquid batteries, this volume expansion induced by decomposition has a profound impact on the underlying thermodynamics of the system.

With regards to the volume expansion, consider the electrolyte  $\text{Li}_{10}\text{GeP}_2\text{S}_{12}$  (LGPS). When LGPS chemically decays, it forms the products  $\text{Li}_4\text{GeS}_4$  (LGS) and  $\text{Li}_3\text{PS}_4$  (LPS). These products maintain a volume that is 2% larger than the original LGPS. Moreover, the volume expansion from electrochemical decompositions of LGPS when charged *vs.* lithium metal is even larger. In fact, the volume increases can reach values in excess of 30% at the complete oxidation (at high voltage) and reduction (at low voltage) limits.<sup>11–13</sup> This increase in volume is referred to as the reaction strain (or reaction dilation) and is denoted  $\epsilon_{\text{RXN}}$ . It is important to note that the reaction strain is a stress-free strain, meaning that unlike conventional strain, it does not result from an applied stress – it is solely the result of the [electro-]chemical decomposition that forms products having a different volume than the initial reactant (*i.e.* the electrolyte).

However, under an applied mechanical constriction, like the typical conditions of SSBs, the reaction strain can result in local stresses. That is to say, although the reaction strain is not caused by mechanical stress, it can cause new local mechanical stresses to form within the battery. These local stresses are the result of trying to fit a larger phase (the decomposed products) into the space that was previously occupied by a smaller phase (the electrolyte). This phenomenon is generalized by the formula  $p = K_{\text{eff}}\epsilon_{\text{RXN}}$ , where the pressure ( $p$ ) is the average compressive stress on a solid volume unit and  $K_{\text{eff}}$  is the local effective mechanical modulus. This formula quantifies that for a solid system, characterized by the modulus  $K_{\text{eff}}$ , to accommodate some expanded decomposition products, a local pressure of  $p$  must be generated. The reaction strain is directly a function of the mole fraction of the present phases. For example, when the respective mole fractions of LGPS, LGS, and LPS are 1, 0, 0, then the reaction strain is 0%. Conversely, when the respective mole fractions are 0, 1/3, 2/3, the reaction strain is 2%. As a result of pressure being a function of  $\epsilon_{\text{RXN}}$  and  $\epsilon_{\text{RXN}}$  being a function of the present mole fractions, it is seen that pressure is now no-longer a state variable but instead a function of the present mole fractions.

This dependency of the mechanical stress distribution on the local mole fractions is what causes such strong changes in

the thermodynamics of solid-state *vs.* liquid batteries. To illustrate this, consider a thought experiment in which a small volume fraction of the total electrolyte, located at  $\vec{r}_0$ , has decomposed. The decomposed [volume] fraction ( $x_{\text{D}}$ ) is now a piecewise function of the form:

$$x_{\text{D}}(\vec{r}) = \begin{cases} 1 & \text{when } \vec{r} = \vec{r}_0 \\ 0 & \text{when } \vec{r} \neq \vec{r}_0 \end{cases}$$

In this case, the reaction strain will be non-zero at  $\vec{r}_0$  and zero everywhere else. As a result, the pressure caused by this reaction strain will also be non-zero only in the neighborhood of  $\vec{r}_0$ . In a liquid battery, such a pressure gradient would not be in equilibrium as fluids must have equal pressures at all point. As a result, the decomposed products would flow within the liquid electrolytes until the pressure comes to equilibrium with the environment (*i.e.* 1 atm for non-pressurized battery cells). In a solid-state system, however, this flow of the decomposed products will not occur because solids can maintain pressure gradients. The result is that a fluid battery will maintain a constant pressure equal to the environment regardless of the decomposed fractions whereas a solid-state battery can potentially have significant localized pressure that is not controlled by the external environment. That is, pressure is a valid state variable of liquid batteries but not one of solid-state batteries.

Consideration of the functional form of  $p(\{x_i\})$  has shown to have profound impact on the intrinsic electrochemical stability of ceramic-sulfide SSEs. When the functional form is taken to be constant (*i.e.* isobaric conditions), the traditional  $pT\phi$  (grand canonical) ensemble is recovered, and the electrochemical stability is seen to be on the order of 1.7–2.1 V *vs.* lithium for LGPS.<sup>14,15</sup> In contrast, when the pressure is taken to be proportional to the local strain, the (meta)stability window can greatly expand to approximately 1–4 V,<sup>1,12,13</sup> and kinetic stability can reach down to 0 V and up to 10 V.<sup>1,4</sup>

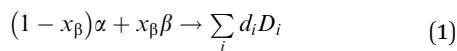
We then for the first time apply the constrained ensemble to explicitly treat the calculation of interface stability, and articulate quantitatively one most critical aspect regarding interface stabilities in SSBs in electrochemical cycling, through the implementation of high-throughput *ab initio* computations and machine learning. We find that when the effective local modulus at the interface ( $K_{\text{eff}}$ ) is above some critical threshold decided by interface materials (denoted  $K_{\text{crit}}$ ), the interface will remain stable throughout cycling. When the effective modulus is lower than this threshold, the interface is unstable and will grow with cycling and thus limit the battery performance. Finally, we utilize this computational approach to develop a method for engineering solid–electrolyte-interphases (SEI's) which are stabilized through electrochemical evolution within a prescribed voltage window.

## Modelling the interfaces within the constrained ensemble

The free energy picture of interface decomposition is depicted in Fig. 1(a). The interface is initially composed of two phases

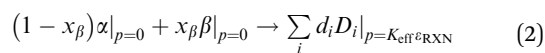


( $\alpha$  and  $\beta$ ) and is seen to decompose with a reaction energy magnitude of  $G_{\text{hull}}$  via some reaction:

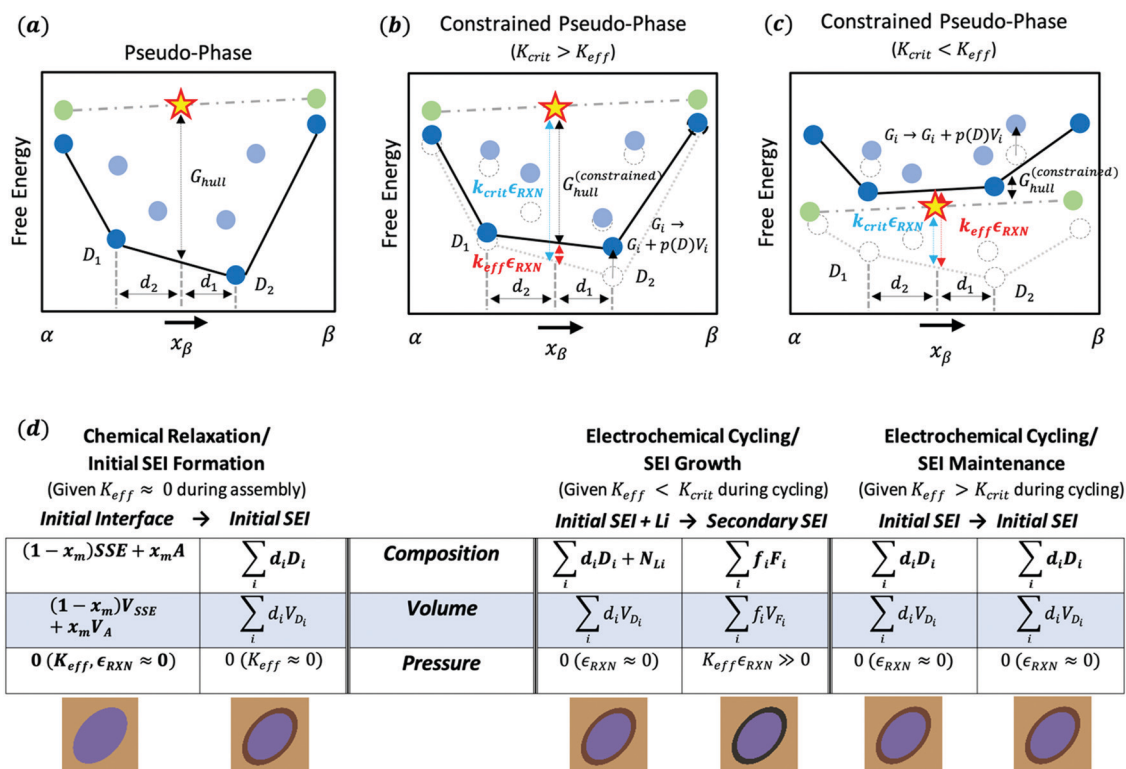


Where  $d_i$ ,  $D_i$  are the decomposed fractions and phases, respectively. The value of  $x_{\beta}$  that maximizes  $G_{\text{hull}}$  is denoted  $x_m$  and defines the “pseudo-phase” as  $(1 - x_m)\alpha + x_m\beta$ .<sup>16</sup> This pseudo-phase is the most kinetically driven decomposition by definition. The value of the pseudo-phase for modelling the interface is that we can computationally represent the interface as a single phase with composition  $(1 - x_m)\alpha + x_m\beta$  and energy  $(1 - x_m)G_{\alpha} + x_m G_{\beta}$  where  $G_i$  is the free energy of phase  $i$ . Thus, *ab initio* computations can be used to predict interface phenomena. Detailed discussion of pseudo-phase methods can be found in ref. 16.

In liquid-cells ( $pT\phi$  ensemble), both the initial interface and decomposed interface exist at the same pressure, which is controlled by the operator. However, analogously to the effects of constraint on bulk SSE's, the pseudo-phase decomposition in solid-state reactions takes the form:



Eqn (1) and (2) differ principally in the products  $D_i$  of eqn (2) existing in a state of elevated pressure given by the local strain due to the decomposition reaction ( $\epsilon_{\text{RXN}}$ ) and some effective modulus ( $K_{\text{eff}}$ ). This additional pressure term perturbs the reaction energy accompanying eqn (1) by a factor of  $K_{\text{eff}}\epsilon_{\text{RXN}}V^0$ , where  $V^0$  is the reference volume. Fig. 1(b) and (c) introduce using the convex hull illustration the parameter  $K_{\text{crit}}$  as a metric for how much mechanical constriction (measured by  $K_{\text{eff}}$ ) is



**Fig. 1** Understanding the critical effective modulus  $K_{\text{crit}}$  within the constrained ensemble description. (a) Convex hull illustration for pseudo-phase models of the solid SEI. The yellow star represents the interphase composition ( $x$ -axis) and energy ( $y$ -axis) of the two reactive phases represented by green dots. The dark blue dots represent phases that are thermodynamically favorable and the black line represents a linear combination of these phases (*i.e.* the convex hull). The phase labels  $D_1$  and  $D_2$  and the lengths  $d_2$  and  $d_1$  represent the well-known “lever rule” for interpreting phase diagrams. The height of the interphase (yellow star) over the convex hull,  $G_{\text{hull}}$ , measures the magnitude of the decomposition energy. For a more detailed understanding of convex-hull pseudo-phase methods, the reader is directed to ref. 9 and 10. (b) An illustration of how a weak mechanical constriction ( $K_{\text{eff}} < K_{\text{crit}}$ ) impacts the interphase convex hull. In this case, each of the potential decomposition phases experiences an increase in free energy corresponding to the pressurization of the phase equal to the generate pressure times the phase’s volume. Under these weak constriction conditions the shift in energy is slight and doesn’t impact which phases are thermodynamically preferred. (c) Same as (b) but now with a strong mechanical constriction ( $K_{\text{eff}} > K_{\text{crit}}$ ). As such, the interphase is now lower in free energy than the prior decomposition products. (d) Comparison of the different chemical and electrochemical reactions that can occur during battery assembly (left) and cycling (right) within the constrained ensemble framework. Generally, mechanical constriction cannot be applied throughout the entire assembly process, so initial SEI formation will likely occur in an unconstrained ensemble. In contrast, during battery cycling the battery may or may not be constrained, which can change the nature of the SEI to propagate and grow with cycling. The illustrations below the table in (d) illustrate how the electrolyte (light brown) and active material (blue) react to form a stable or unstable SEI depending on mechanical constriction. The initial SEI is in dark brown. If the battery is cycled with weak constriction ( $K_{\text{eff}} < K_{\text{crit}}$ ), this initial SEI will further react to a new SEI (black) and likely grow. If the battery is cycled with a strong constriction ( $K_{\text{eff}} > K_{\text{crit}}$ ), the initial SEI will remain in place.



required for the perturbed reaction to cease being thermodynamically favorable. Eqn (3) describes  $K_{\text{crit}}$  mathematically as the value of  $K_{\text{eff}}$  such that the energy cost of local pressurization exactly cancels the driving reaction energy ( $\Delta G_{\text{RXN}}^0$ ). Thus, the interface reaction is expected to be thermodynamically favorable if and only if  $K_{\text{eff}} < K_{\text{crit}}$ .

$$\Delta G_{\text{RXN}}^0 + K_{\text{crit}} \epsilon_{\text{RXN}} V^0 = 0 \quad (3)$$

In light of the effects of Fig. 1(a)–(c), Fig. 1(d) catalogs expected reactions involving SEI formation in solid-state batteries. The initial interfacing materials are taken to be the SSE and an arbitrary material “A”. During battery assembly, there exists virtually no applied mechanical constriction ( $K_{\text{eff}} \approx 0$  GPa) so the interfacial chemical reaction proceeds as in a standard canonical form (*i.e.* the system is not yet open to lithium, hence it is not grand canonical). Following pseudo-phase conventions, the fraction of A that produces the maximum thermodynamic driving force is denoted  $x_m$ .

Deviation from grand canonical behavior is possible after the battery cell is fully assembled, at which point mechanical constriction may (or may not) be applied. In the absence of mechanical constriction, or when the local mechanical constriction is weak ( $K_{\text{eff}} \ll K_{\text{crit}}$ ), the initially formed SEI will be able to further decay once the battery is assembled and the system becomes open to lithium. This is represented in Fig. 1(d) as SEI growth. As the battery is cycled, the voltage will likely pass in and out of the SEI’s electrochemical stability window, causing the initial SEI to continuously decay into secondary SEIs allowing continued growth. In contrast, if the local mechanical constriction is sufficiently rigid ( $K_{\text{eff}} > K_{\text{crit}}$ ), either initially or after certain cycling steps, the following cycling of the voltage in and out of the initial SEI’s stability window will not lead to further decompositions. This process is described as “SEI maintenance” through the local stability gained from the electrochemical evolution that could contain

processes of both the initial decomposition ( $K_{\text{eff}} < K_{\text{crit}}$ ) and the further stabilization ( $K_{\text{eff}} > K_{\text{crit}}$ ).

## Towards SEI maintenance: $K_{\text{crit}}$ distribution and elemental correlation

From a functional perspective, given the inevitable formation of an initial solid–electrolyte–interphase (SEI) during battery assembly, the goal of solid-state battery design should be to focus on designing the initial SEI’s such that they don’t grow during cycling. Fig. 2 illustrates a computational process to determine such ideal initial SEI’s. In short, an ensemble of 50 randomly initialized decision trees is trained to predict  $K_{\text{crit}}$  of the ground state phase at any input composition. The ensemble is then used to fine-tune the composition of known parent glass phase sulfides with good lithium-ion conductivity. The resulting child compositions should be minimally changed from the parent compositions but modulated to decrease the interface  $K_{\text{crit}}$  to as low of values as is possible. In this paper, we illustrate the computational method using only the interface between various glass coatings and four well known ceramic-sulfides. However, such a process is readily generalizable to the interface between the glass coatings and the active electrode materials, between ceramic electrolyte and uncoated electrode materials, as well as coatings beyond glasses, *i.e.*, (nano)crystalline or polymer phases.

Over 80 000 SEI’s were simulated using pseudo-binary calculation of over 20 000 electronic insulators from the Materials Project with four well known ceramic-sulfide solid electrolytes –  $\text{Li}_{10}\text{GeP}_2\text{S}_{12}$  (LGPS),  $\text{Li}_{10}\text{SiP}_2\text{S}_{12}$  (LSPS),  $\text{Li}_7\text{P}_3\text{S}_{11}$  (LPS), and  $\text{Na}_7\text{P}_3\text{S}_{11}$  (NPS). Each data point consisted of the initial SEI composition, formed by simulating a chemical reaction between the SSE and the coating, and the  $K_{\text{crit}}$  corresponding to stabilizing the initial SEI at a cathode voltage (*e.g.*, from 2 V to 4 V) vs. lithium

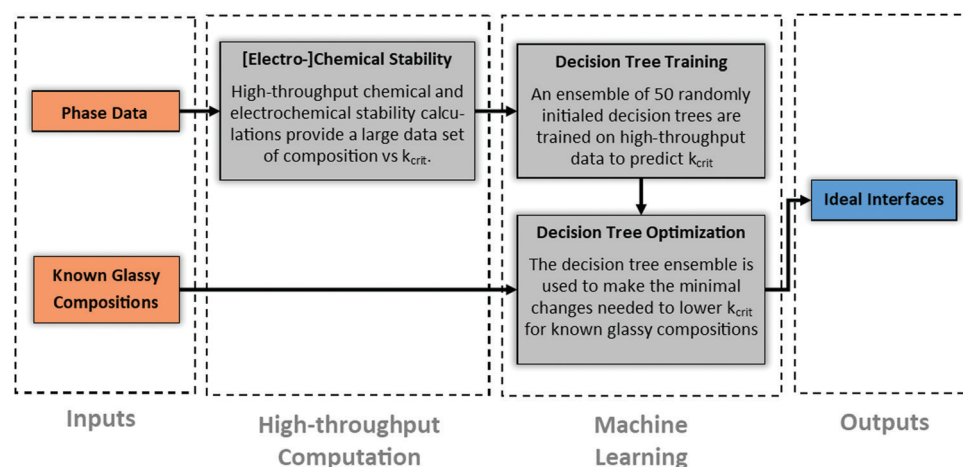


Fig. 2 Overview of computational approach. High throughput computation is performed to obtain datapoints on how  $K_{\text{crit}}$  varies with composition. These datapoints are then leveraged to train an ensemble of 50 decision trees to predict  $K_{\text{crit}}$  based on composition. Finally, using known glassy compounds with good lithium-ion conductivity, the decision tree is used to make minimal compositional changes to lower  $K_{\text{crit}}$ . The resulting compounds should have significantly improved response to mechanical constriction. These represent ideal interfaces for SSEs as they are both ionically conductive and mechanically metastable.



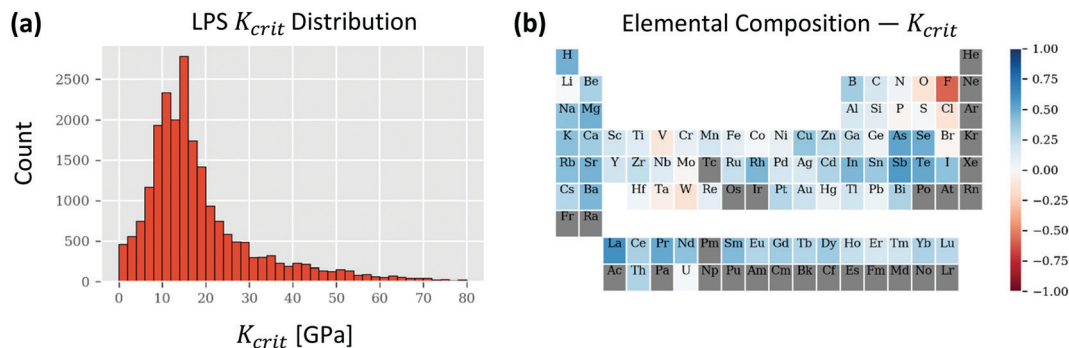


Fig. 3 Statistical behavior of interface  $K_{crit}$  at 4 V. (a) The distribution of  $K_{crit}$  for binary pseudo-phases where one of the interfacial materials is LPS. (b) The correlation of elemental composition and  $K_{crit}$  for all pseudo-phases considered (LPS, LGPS, LSPS, NPS). Red elements are those that lower  $K_{crit}$  [ideal], whereas those elements that are blue raise  $K_{crit}$ .

metal. In other words,  $K_{crit}$  indicates how much local mechanical constriction is required to stop the initial SEI from decomposing when cycled to the cathode voltage region. A lower  $K_{crit}$  thus also suggests that the interface is more likely to be stabilized in further cycling steps even if the initial local  $K_{eff}$  is less than  $K_{crit}$ , as the electrochemical evolution will more easily increase the local  $K_{eff}$  beyond  $K_{crit}$ .

Fig. 3(a) shows the distribution of  $K_{crit}$  determined for interfaces of LPS paired with the various coating materials. The distribution has a mean, median, and standard deviation of 18.5, 14.8, and 13.0 GPa, respectively. The large standard deviation indicates that the value of  $K_{crit}$  of an initial interface can vary considerably with composition and, thus, allows for substantial engineering to optimize the interface's susceptibility to mechanical constriction within the constrained ensemble framework. Similarly, Fig. S1(a–c) (ESI<sup>†</sup>) show the  $K_{crit}$  distribution for interfaces with LGPS, LSPS, and NPS. Fig. S1(d) (ESI<sup>†</sup>) depicts these statistical figures of merit (mean, median, standard deviation) for each SSE interface distribution. It remains true for each SSE that the distribution is quite broad, implying there is ample opportunity for engineering ideal interfaces with low  $K_{crit}$  values.

The correlation between the elemental composition of SEIs and  $K_{crit}$  is depicted in Fig. 3(b). Those atomic species with negative correlation (red) tend to lower  $K_{crit}$  as their atomic fraction is increased. Conversely, those atomic species with positive correlation (blue) tend to raise  $K_{crit}$ . The highly electronegative oxygen, fluorine, and chlorine are seen to be amongst the most beneficial atomic species for maintaining a low  $K_{crit}$  at the cathode voltage. This is likely attributed to these anions having high voltage redox potential. In other words, SEIs with high levels of oxygen, fluorine, and chlorine are less prone to delithiation in the considered voltage ranges. Additionally, V, Mo, and W are also seen to improve the stability of high voltage SEIs, presumably indicating that these atomic species are also unlikely to undergo reactions below 4 V.

## Designing SEIs with low $K_{crit}$

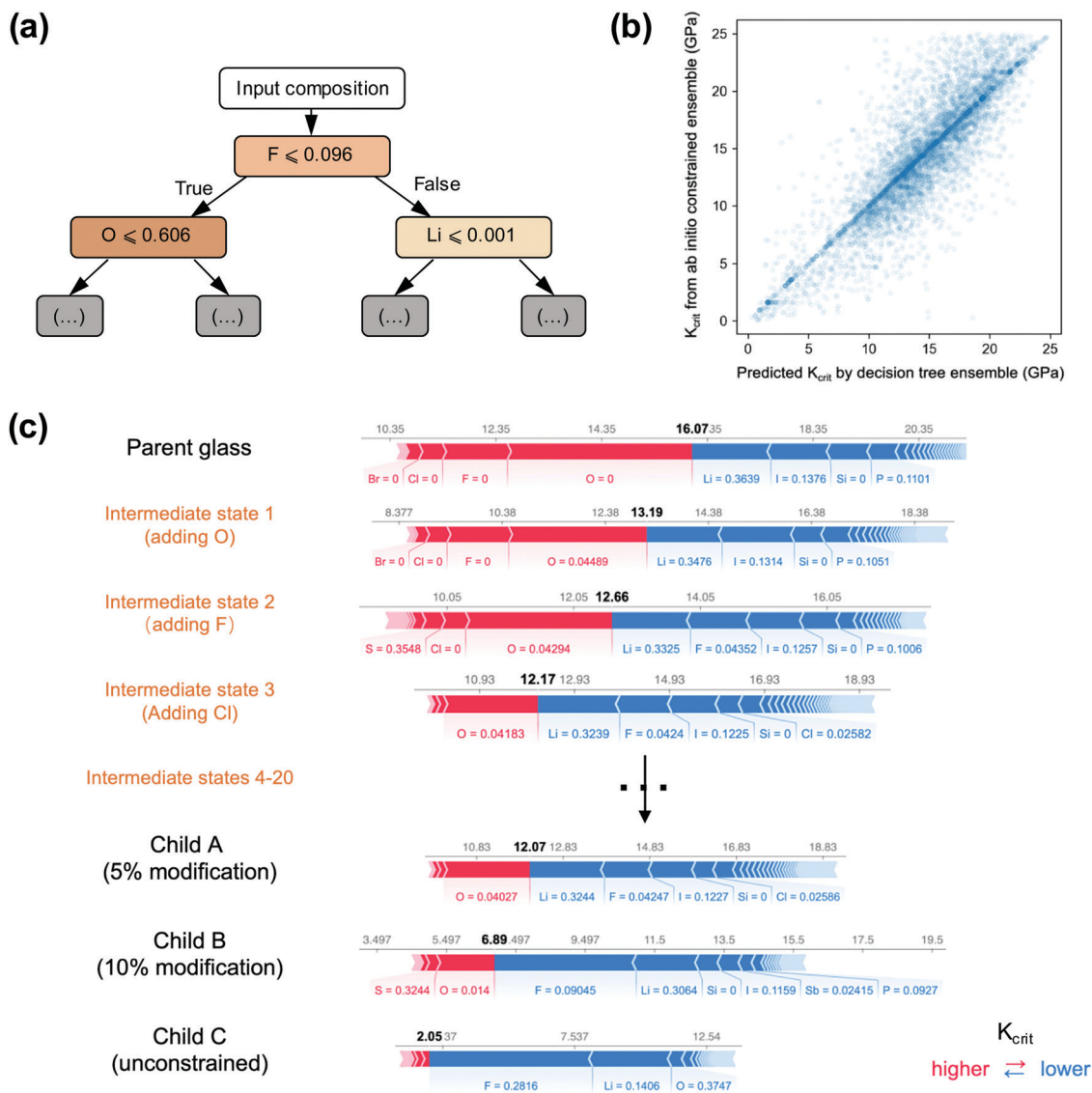
In light of these results, it is apparent that initial SEIs could be designed to minimize  $K_{crit}$  and hence decrease growth of the

SEI upon cycling. In evaluating interfaces that are likely to occur in solid-state cathode, such as between LPS and common cathode materials, Fig. S2 (ESI<sup>†</sup>) shows that the  $K_{crit}$  tends to be non-linear, oscillatory, with multiple local minima vs. composition. This highly nonlinear relationship between  $K_{crit}$  and composition suggests that optimizing  $K_{crit}$  directly is difficult, considering that the composition space is very high dimensional.

To computationally capture the nonlinear effects of composition on  $K_{crit}$ , we utilized an ensemble model of decision trees. The model was trained on the previously determined SEI data to predict  $K_{crit}$ . The conditional statements of each node are learned by the tree to most accurately reflect the underlying composition dependence of  $K_{crit}$ . The minimum unit of an example tree section illustrated in Fig. 4(a), together with the tree with more layers and branches depicted in Fig. S3 (ESI<sup>†</sup>), confirms the importance of electronegative anions for maintaining a low  $K_{crit}$ . For example, the left most branch at each layer implements a conditional statement on O or Cl. If the material has a low composition of those anions, the tree is with a higher predicted value of  $K_{crit}$ . Similarly, the right most branch states that if the material has a high composition of Li, F, Al, the tree is with a lower predicted value of  $K_{crit}$ . While each node corresponds to the evaluation of a single element, each element may be evaluated many times throughout the tree. Thus, each tree represents an approximate mapping of the  $K_{crit}$  manifold. The ensemble then performs a weighted average of the predicted manifolds for the final value, *i.e.*, to predict a value for a given composition and a weighted average then gives a final prediction for the input composition. As shown in Fig. 4(b), which compares the predicted and actual  $K_{crit}$  values for a test set of 10 000 SEIs, this approach can readily capture the complicated compositional dependence, especially in the lower range of  $K_{crit}$ . The root mean square error (RMSE) for this test set is 2.59 GPa.

Using the trained decision tree ensemble, we can modify the composition of known lithium-ion conductors to decrease the  $K_{crit}$  and improve the SEI stability. To help us understand the role of each element on interface  $K_{crit}$ , we applied a feature-attribution model SHAP, or Shapley additive explanations,<sup>17</sup> to our trained ensemble to extract the influence of the composition of each element on the  $K_{crit}$ . More technical details about the





**Fig. 4** Decision tree method and results for composition design. (a) Illustration of the structure and mechanism of decision trees. The input composition is parsed through a hierarchy of criteria on the composition and generates output (see Fig. S3 for larger illustration, ESI<sup>†</sup>). (b) Comparison of the predicted  $K_{\text{crit}}$  by the decision tree and the calculated  $K_{\text{crit}}$  at the interface to LSPS at 4 V from our constrained ensemble *ab initio* platform, with a root-mean-square error (RMSE) of 2.59 GPa. (c) Optimization from (top to bottom) parent glassy material to child A (5% compositional change) to child B (10% compositional change) to child C (unlimited compositional change). The individual influence of each element is calculated by the SHAP model. Three intermediate states during the optimization process illustrate how the O, F, and Cl elements are introduced into the optimized Child A material as suggested by the SHAP model.

optimization method is described in the Methods. We tested various materials with changing levels of restriction on the elemental composition to better understand how  $K_{\text{crit}}$  will change based on the doping amount. Table S1 (ESI<sup>†</sup>) shows compositional optimization of 10 well known sulfide glass SSE's and the corresponding child phases where the composition has been changed to lower the interface  $K_{\text{crit}}$  to LSPS. Child A, B, and C correspond to maximum allowed changes in the stoichiometry of each element as 0.05, 0.1, or no constraint, respectively. Note that in practice it is preferable to make the minimum possible compositional changes to lower the  $K_{\text{crit}}$ , as the ionic conductivity will likely fall quickly as composition

is deviated. In Table S1 (ESI<sup>†</sup>), the known sulfide glasses have  $K_{\text{crit}}$  values that are characteristically on the order of 15–17 GPa from our prediction. These values decrease to 10–13 GPa in child A phases, 6–9 GPa in child B phases, and 1.5–2 GPa in child C phases. This  $K_{\text{crit}}$  decrease is expected to remarkably improve interface stability in actual solid-state battery cells, as the local mechanical constriction  $K_{\text{eff}}$  is not necessarily at a sufficiently high level, depending on materials and battery assembly approaches.

We now use the first parent compound and its child compositions in Table S1 (ESI<sup>†</sup>) as an example to illustrate how our platform finds new compositions with lowered  $K_{\text{crit}}$ .

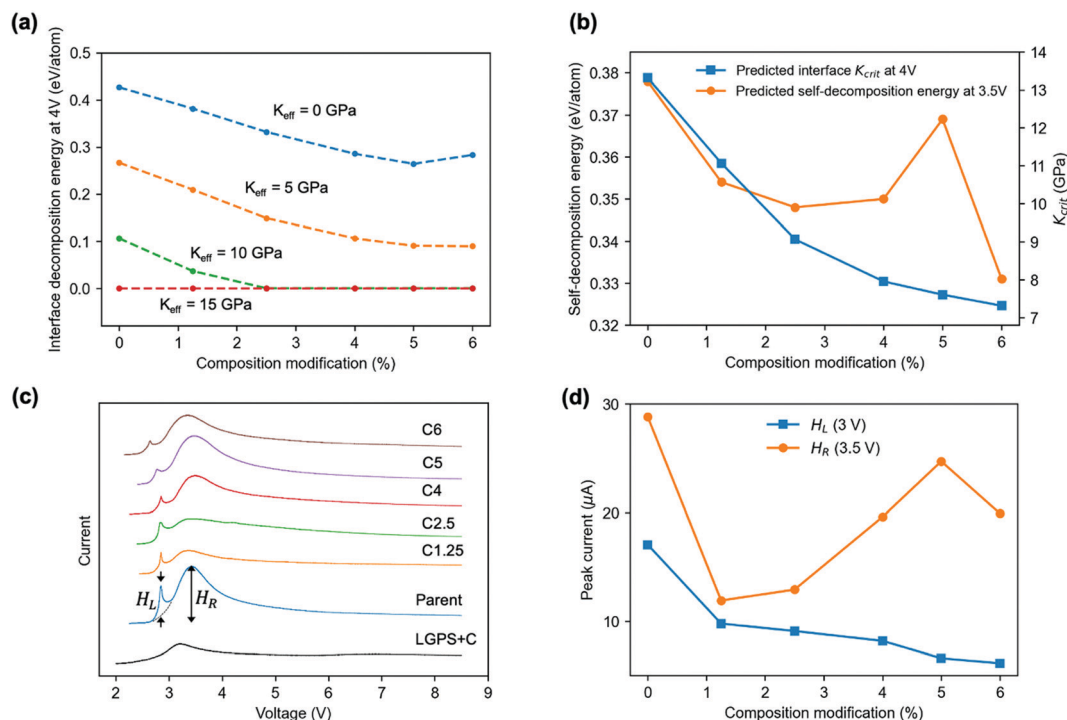


In Fig. 4c, the length of each bar (in the unit of GPa) approximates how much the corresponding element increases (red) or decreases (blue) the  $K_{\text{crit}}$ . The SHAP model analyzed that for the parent glass of  $\text{Li}_{0.364}\text{P}_{0.110}\text{S}_{0.388}\text{I}_{0.138}$ , the absence of O, F and Cl are dominantly responsible for the relatively high  $K_{\text{crit}}$ , as the first three long red bars are with O = 0, F = 0 and Cl = 0 in Fig. 4c, indicating that the presence of O, F and Cl in the compound will reduce  $K_{\text{crit}}$ . Meanwhile, the blue bars for existing elements of Li, I, P in the parent glass indicate that having them as existing nonzero compositions is not in conflict with the optimization goal of reducing  $K_{\text{crit}}$ .

It's natural to understand that the optimization effort should be given to the features with red bars, since the length of red bar also roughly indicates how much we can further decrease the  $K_{\text{crit}}$  if we fully optimize the composition of the corresponding elements starting from a given composition. We take an approach to efficiently optimize  $K_{\text{crit}}$  by focusing on the element corresponding to either the longest red bar at each step, or the 2nd longest red bar if the longest bar element has just been updated in the previous step (*i.e.*, a pseudo-longest red bar). The actual optimization process of  $K_{\text{crit}}$  is also influenced by the concurrent composition of other elements brought by changing the composition of one element. For example, the child A, B, C compositions bring down the  $K_{\text{crit}}$  from the 16.07 GPa (parent glass) to 12.07 GPa (Child A),

6.89 (Child B), and 2.05 GPa (Child C) with an increasing modification range of compositions. F, O and Cl are first introduced to Child A composition of  $\text{Li}_{0.324}\text{P}_{0.098}\text{S}_{0.346}\text{I}_{0.138}\text{Cl}_{0.026}\text{F}_{0.042}\text{O}_{0.040}$ , because these three elements all appear as either the longest or the pseudo-longest red bar in the initial intermediate optimization steps from the Parent toward the Child A composition (Fig. 4c), with Li, P, S, I compositions being forced to make slight concurrent decreases. The algorithm is very efficient in finding the right composition, as the intermediate compositions after the first 3 intermediate steps are very close to the Child A composition, with the remaining 17 intermediate steps just as a fine-tune of the composition. Through the composition optimization from Parent to Child A, B, C compounds, the total length of red bars shrinks and the  $K_{\text{crit}}$  decreases generally. At Child C with unconstrained compositional change, the red bars are very short, suggesting a very limited remaining room to further decrease  $K_{\text{crit}}$  through compositional modification. The general trend of adding more F, O, Cl, Br for reduced  $K_{\text{crit}}$  as predicted by the SHAP model is also in good agreement with the elemental correlation map (Fig. 3(b)).

Using the predicted  $K_{\text{crit}}$  of SEI at 4 V, we further predict the 4 V interface (to LGPS) decomposition energy for the SEI under various levels of composition modification (Fig. 5(a)). Similar to  $K_{\text{crit}}$ , the interface decomposition energy also tends to decrease with the level of modification. A  $K_{\text{eff}}$  of 5 GPa can further lower



**Fig. 5** Material modification and experimental validation based on parent glass  $\text{Li}_{0.364}\text{P}_{0.110}\text{S}_{0.388}\text{I}_{0.138}$  (Table S2, ESI†). (a) Predicted interface decomposition energy at 4 V for the parent (0%) and modified glass coatings (1.25, 2.5, 4, 5, 6%) to LGPS electrolyte at different  $K_{\text{eff}}$ . The x% composition modification suggested by our SHAP model means the stoichiometry of each element is allowed to change up to a number of x/100, regardless of the initial amount of this element. (b) Predicted  $K_{\text{crit}}$  of coating materials at the interface to LGPS electrolyte (blue) and the self-decomposition energy of coating materials at  $K_{\text{eff}} = 0$  GPa (orange) vs. composition modification level. (c) CV test of the synthesized parent glass and five modified child glasses, labelled as Cx, where x corresponds to the modification level in (a).  $H_L$  and  $H_R$  mark the peak height for the decomposition current at around 3 V (pre-peak) and 3.5 V (broad peak), respectively. (d) Pre-peak ( $H_L$ ) and broad peak ( $H_R$ ) decomposition currents vs. composition modification level.



the interface decomposition energies from that at 0 GPa while keeping the similar decreasing trend with composition modification. A  $K_{\text{eff}}$  of 10 GPa can stabilize child materials at the 4 V interface to LGPS with a composition modification larger than 2.5%, for which  $K_{\text{eff}} > K_{\text{crit}}$  is satisfied due to the reduced  $K_{\text{crit}}$  by composition modifications (Fig. 5b). With a 15 GPa  $K_{\text{eff}}$ , all the materials, including the parent material will be stabilized. The trends described in Fig. 5a and b are qualitatively the same for interfaces calculated at 2 V and 3 V (Fig. S4, ESI<sup>†</sup>), except for some quantitative changes by voltage. Note that the  $K_{\text{crit}}$  at 2 V are all below 4.6 GPa and that at 3 V are all above 4.9 GPa at all modifications.

## Testing the model by experiment

To further test the predictability of our model, we synthesized the parent and predicted child glasses of the first row of Table S2 (ESI<sup>†</sup>) and performed cyclic voltammetry (CV) test based on batteries of normalized cathode weight, area and composition, as shown in Fig. 5(c) (Methods). Two decomposition current peaks labelled as  $H_L$  and  $H_R$  are observed at 3 V and 3.5 V, respectively. The pre-peak  $H_L$  is found to decrease monotonically with modification, while the broad peak  $H_R$  is oscillatory (Fig. 5(d)). We found that the predicted self-decomposition energy at 3.5 V at  $K_{\text{eff}} = 0$  GPa in Fig. 5(b) exhibits a very similar trend to the measured  $H_R$  curve, suggesting that the change of broad peak intensity around 3.5 V with composition modification should be mainly caused by the self-decomposition current of the glasses in the initial charge, where the decomposition happens at the glass region near the many inevitable pores in the cold pressed pellet,<sup>18</sup> where  $K_{\text{eff}} = 0$  GPa. Note that without glasses, the control cathode of LGPS mixed only with amorphous conductive carbon shows a broad peak of decomposition current below 9  $\mu\text{A}$  in the CV test, which defines the background contribution from the LGPS self-decomposition to the measured current that is independent from the modification to the glasses. Thus, the pre-peak  $H_L$  has to be contributed by the interface decomposition between LGPS and glass at a weak local mechanical constriction in the initial charge, consistent with the general decreasing trend for both curves of the measured  $H_L$  (Fig. 5d) and the predicted interface decomposition energy at a low  $K_{\text{eff}}$  (Fig. 5a and Fig. S4, ESI<sup>†</sup>) versus modifications.

The fact that all the  $H_L$  pre-peaks are around 3 V suggests that the local  $K_{\text{eff}}$  at the interface in the initial charge is around 4.6 to 4.9 GPa so that it is higher than all the  $K_{\text{crit}}$  at 2 V (Fig. S4a, ESI<sup>†</sup>) but lower than all the  $K_{\text{crit}}$  at 3 V (Fig. S4b, ESI<sup>†</sup>). Such an initial interface  $K_{\text{eff}}$  is decided by the mechanical property of the two materials that form the interface (glass sulfide and LGPS here) and the processing and testing procedures (hand mixing, cold initial and stack press here) that define the interface contact. Only when the voltage is charged approaching 3 V, the local  $K_{\text{eff}}$  starts to be smaller than  $K_{\text{crit}}$  at the interface, as  $K_{\text{crit}}$  increases with charge (comparing Fig. 5b and Fig. S4b, ESI<sup>†</sup>), so that the interface decomposition

happens. However, after the initial charge, the decomposition current almost vanishes in discharge and the following CV cycles below 4 V (Fig. S5, ESI<sup>†</sup>), suggesting that the local  $K_{\text{eff}}$  of the glass-LGPS mixture upon the initial self and interface decompositions is increased beyond the  $K_{\text{crit}}$  at all modifications both at the interface ( $K_{\text{crit}} < 14$  GPa, Fig. 5b) and inside the glass ( $K_{\text{crit}}^{\text{self}} < 6$  GPa, Fig. S6, ESI<sup>†</sup>). Therefore, it is through these local decompositions in the initial charge that the order of  $K_{\text{eff}}$  and  $K_{\text{crit}}$  is switched at some low  $K_{\text{eff}}$  regions, leading to the largely suppressed decomposition in further cycling, where the local  $K_{\text{eff}}$  is increased beyond 14 GPa throughout the charged pellet and  $K_{\text{eff}} > K_{\text{crit}}$  is generally satisfied below 4 V. Note that through the comparison between computation and experiment an upper limit of  $K_{\text{eff}}$  value around 15 to 20 GPa is also estimated for crystalline phase of LSPS<sup>12</sup> and LGPS.<sup>1,4,13</sup> Such a  $K_{\text{eff}}$  is largely decided by the mechanical modulus of the electrolyte materials, which is similar for crystalline<sup>19</sup> and amorphous<sup>20</sup> sulfides.

We have shown that within the constrained ensemble that is unique to solid-state batteries, the use of randomly initialized decision tree ensembles has significant potential for optimizing SEIs at multiple solid–solid interfaces. Fine-tuning the SEI composition based on the decision tree results has been able to reduce the  $K_{\text{crit}}$  significantly. Although the examples given here are tested at an experimental condition with strong mechanical constriction (see Methods) so that the decomposition current is largely inhibited after the initial charge regardless of the composition, in practice, the composition modification strategy for the electrolyte and coating materials could greatly reduce the burden of void elimination procedures during battery assembly and micro-crack prevention approaches during battery test, favoring the ideal manufacturing conditions at weak mechanical constriction. As a result, the method opens the door to realizable solid-state battery design and production based on mechanical constriction induced metastability.

## Methods

### Experiment

The amorphous solid electrolytes (P1-C6) are synthesized by high energy ball milling at 350 rpm. Stoichiometric amounts of  $\text{Li}_2\text{S}$  (>99.9% purity, Alfa Aesar),  $\text{P}_2\text{S}_5$  (>99% purity, Sigma Aldrich),  $\text{LiI}$  (>99% purity, Sigma Aldrich),  $\text{LiF}$  (>99.99% purity, Sigma Aldrich), and  $\text{Li}_2\text{O}$  (>97%, Sigma Aldrich) were milled for 20 hours. A total of 1 mg LGPS–amorphous SE–carbon composite cathode (weight ratio = 45:45:10) was used as the cathode. The composite was mixed and rolled into a thin film by a dry method with the addition of 5% PTFE. The diameter of the cathode film is 0.49  $\text{cm}^2$ . The area and weight of all batteries were kept constant for all CV measurements. 120 mg LGPS was applied as the electrolyte while Li/graphite (capacity ratio = 2.5:1) was applied as an anode. The battery of Li/graphite–LPSCl–LGPS–cathode was pressed in a homemade pressurized cell, where LPSCl represents  $\text{Li}_{5.5}\text{PS}_{4.5}\text{Cl}_{1.5}$  that is used to guarantee a stable anode interface following our



recent approach.<sup>21</sup> The batteries were initially pressed at 467 MPa while the operating pressure was kept at 250 MPa during battery test to ensure the mechanical constriction. In the CV test, a voltage ranges from the open circuit voltage to 8.5 V was applied with the scan rate of 0.1 mV s<sup>-1</sup>. The control sample labelled as LGPS + C in Fig. 5 is constructed with the similar procedure but using only 0.5 mg of LGPS + C (weight ratio = 90 : 10) as cathode, without mixing with the amorphous glass.

### High-throughput computation

Thermodynamical chemical decomposition reactions between materials (DFT data obtained from Materials Project) and an electrolyte were first calculated to generate an interphase for each material/electrolyte combination by constructing phase diagram using the Python Materials Genomics (pymatgen) library<sup>22</sup> with the binary-search algorithm.<sup>9</sup> The electrochemical stability of each interphase is then calculated from its grand potential phase diagram to obtain decomposition energy and reaction strain for calculating  $K_{\text{crit}}$  by eqn (2) and (3). All *ab initio* calculations of phase data were obtained following the Material Project calculation guidelines in the Vienna *ab initio* Software Package (VASP). The mechanical constriction induced metastability calculations were performed on the computational platform we developed following the perturbation or Lagrangian optimization methods.<sup>11–13</sup>

### Decision tree learning

A decision tree consists of hierarchical computation (decision) nodes. A typical decision tree is depicted in Fig. S3 (ESI†). The input data to the decision trees is in the form  $(X, y) = (\{x_1, x_2, \dots, x_n\}, y)$  where  $x_i$  are the features and  $y$  is a target value. The decision tree can perform both the regression and classification tasks, depending on whether the nature of target variable  $y$  being continuous or a finite number of classes. Starting with the input features, each node of the tree applies a conditional statement on the value of a feature, then moves to a subsequent node based on the truth of that statement. The optimization of the tree includes choosing both the feature and threshold for the criteria for each node that overall best splits the set of items. Instead of measuring the error, better metrics such as the cross entropy and the Gini index are generally used to measure the goodness of the choice of criteria and data split.<sup>23</sup>

We use an ensemble model of individual decision trees, the Extremely Randomized Tree model.<sup>24</sup> In such models, a number of  $N$  trees are initialized simultaneously ( $N = 30$  in our setting). Each tree in the ensemble is fed with training data sampled from the training set. A random subset of candidate features is used, from which thresholds are drawn at random for each candidate feature, and the best of these randomly generated thresholds is picked as the splitting rule.

### Feature attribution and composition optimization

SHAP, or Shapley additive explanations, is a game theoretic approach to explain the output of any machine learning model.<sup>17</sup> It connects optimal credit allocation with local explanations using

the classic Shapley values from game theory and their related extensions and obtain the influence of the individual features on the output. In our case, we apply it to extract how each elemental composition of the SEI influences the  $K_{\text{crit}}$ .

We compute the SHAP values and choose the element with the most positive SHAP value, *i.e.*, the presence of which has the most positive influence on the  $K_{\text{crit}}$ . We then optimize the composition of this chosen element under the given constraint. With the composition optimized, we compute the SHAP value again and optimize the next element with the most positive SHAP value. Due to the composition constraint, it is possible that the element that is just updated still has the highest SHAP value. In this case, we move to the element with the second-highest SHAP value (pseudo-longest red bar in Fig. 4c) in order to proceed the optimization. The optimization process ended until the  $K_{\text{crit}}$  cannot be further reduced.

## Conflicts of interest

The authors declare no competing financial or non-financial interests.

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