UC Berkeley

UC Berkeley Previously Published Works

Title

Design principles for NASICON super-ionic conductors

Permalink

https://escholarship.org/uc/item/77r6n6n9

Journal

Nature Communications, 14(1)

ISSN

2041-1723

Authors

Wang, Jingyang

He, Tanjin

Yang, Xiaochen

et al.

Publication Date

2023

DOI

10.1038/s41467-023-40669-0

Peer reviewed

nature communications



Article

https://doi.org/10.1038/s41467-023-40669-0

Design principles for NASICON super-ionic conductors

Received: 2 November 2022

Accepted: 3 August 2023

Published online: 25 August 2023



Jingyang Wang^{1,2,3}, Tanjin He D^{1,2}, Xiaochen Yang^{1,2}, Zijian Cai D^{1,2}, Yan Wang D⁴, Valentina Lacivita⁴, Haegyeom Kim¹, Bin Ouyang⁵ ⋈ & Gerbrand Ceder ® ^{1,2} ⋈

Na Super Ionic Conductor (NASICON) materials are an important class of solidstate electrolytes owing to their high ionic conductivity and superior chemical and electrochemical stability. In this paper, we combine first-principles calculations, experimental synthesis and testing, and natural language-driven text-mined historical data on NASICON ionic conductivity to achieve clear insights into how chemical composition influences the Na-ion conductivity. These insights, together with a high-throughput first-principles analysis of the compositional space over which NASICONs are expected to be stable, lead to the successful synthesis and electrochemical investigation of several new NASICONs solid-state conductors. Among these, a high ionic conductivity of 1.2 mS cm⁻¹ could be achieved at 25 °C. We find that the ionic conductivity increases with average metal size up to a certain value and that the substitution of PO₄ polyanions by SiO₄ also enhances the ionic conductivity. While optimal ionic conductivity is found near a Na content of 3 per formula unit, the exact optimum depends on other compositional variables. Surprisingly, the Na content enhances the ionic conductivity mostly through its effect on the activation barrier, rather than through the carrier concentration. These deconvoluted design criteria may provide guidelines for the design of optimized NASICON conductors.

Solid-state batteries are a promising next-generation battery technology because of their potential for improvements in safety and energy density, stemming from the replacement of the conventional flammable organic electrolyte by a dense layer of a solid-state conductor that serves as both the alkali-ion-conducting electrolyte and separator^{1,2}. Numerous efforts have thus been devoted to the discovery of solid-state conductors with high ionic conductivity as well as good chemical and electrochemical stability³⁻⁹. Among these, sodium superionic conductors, or NASICONs for short, are of considerable interest. NASICON materials are a class of polyanionic materials with the general formula $Na_xM_2(AO_4)_3$, where M represents a transition or main group metal and AO₄ represents a polyanion^{10,11}. A robust crystalline framework with the rhombohedral (R-3c) symmetry is constructed by corner-sharing metal octahedra and polyanion tetrahedra, as illustrated in Fig. 1a, which creates two distinct Na sites. Therefore, Na ions can migrate from one site to another through two triangular bottlenecks and form a three-dimensional percolating channel for fast ion conduction¹². NASICON compounds can also form in a monoclinic symmetry (C2/ c) depending on the composition and temperature, which splits the Na2 site into two different sites 12. However, the framework skeleton remains the same, and thus, the ion-transport channel is similar. The high ionic conductivity and structural stability of NASICON-type materials are of great interest for building solid-state batteries and

¹Materials Sciences Division, Lawrence Berkeley National Laboratory, Berkeley, CA 94720, USA. ²Department of Materials Science and Engineering, University of California, Berkeley, CA 94720, USA. 3School of Sustainable Energy and Resources, School of Materials Science and Intelligent Engineering, Nanjing University, Suzhou, China. ⁴Advanced Materials Lab, Samsung Advanced Institute of Technology and Samsung Semiconductor, Inc, Cambridge, MA 02138, USA. 5Department of Chemistry and Biochemistry, Florida State University, Tallahassee, FL 32306, USA. 🖂 e-mail: bouyang@fsu.edu; gceder@berkeley.edu

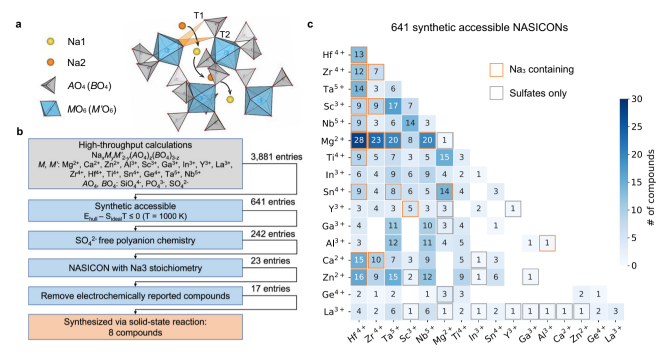


Fig. 1 | **Heat map of synthetically accessible NASICONs predicted by DFT. a** Crystal structure of a rhombohedral NASICON showing the Na-ion conduction pathway. Two bottleneck triangles are denoted as T1 and T2. **b** High-throughput computation workflow. **c** Distribution of the 641 DFT calculated synthetically

accessible NASICONs. The color map indicates the number of stable compounds containing a specific metal pair. The orange squares indicate blocks including NASICONs with Na₃ stoichiometry, and the grey squares mark blocks containing only sulfate-type NASICONs.

other applications such as ion-selective membranes and gas-sensing devices¹³⁻¹⁸.

The history of NASICON-type materials can be traced back to the 1960s. Hong and Goodenough et al. reported the prototype of NASI-CON, Na₃Zr₂(SiO₄)₂(PO₄). They discovered that the polyanion composition is quite flexible as the ratio between SiO₄ and PO₄ can be finely tuned. forming a complete solid-solution $Na_{1+x}Zr_2(SiO_4)_x(PO_4)_{3-x}$ $(0 \le x \le 3)$, with the Na content varying from 0 to 4 as required by charge neutrality^{10,11}. The ionic conductivity of this solid solution at 300 °C achieved an optimal value of $0.2 \,\mathrm{S\,cm^{-1}}$ when $x = 2^{11}$. However, the specific roles of the polyanion composition and Na content in determining the ionic conductivity have not been strictly decoupled¹⁹. Different polyanion groups, such as AsO₄³⁻, SO₄²⁻, and SeO₄²⁻, can be incorporated in the NASICON structure^{20,21}, though compounds with these polyanions have not been studied as extensively as the Si-P system. The ability of the NASICON structure to accommodate different ions on the M site has been extensively demonstrated. Divalent (Mg²⁺, Ca²⁺, etc.), trivalent (Al³⁺, Ga³⁺, Sc³⁺, etc.), tetravalent (Zr⁴⁺, Hf⁴⁺, Ti⁴⁺, etc.), and pentavalent (Nb⁵⁺, Sb⁵⁺, etc.) metal cations can all be stabilized in the M site with relatively large solubilities^{22,23}, and the ionic conductivity can be increased by one order of magnitude using aliovalent doping strategies^{14,23}. However, because these compositional variables (e.g., Na content, metal and polyanion species) are highly convoluted, their distinct effect on the ionic conductivity is challenging to differentiate and rationalize. For example, replacing Zr⁴⁺ with lower-valent Mg²⁺ also increases the Na content²⁴, Moreover, because the NASICON framework can accommodate a wide range of chemical species in the M and A sites^{11,21,23,25-35}, the possible compositional space is immense³⁶. Although we systematically studied the compositional bounds on phase stability of NASICONs in recent work³⁶, a general picture of the relationship between the compositional variables of NASICON materials and their conductivities, is essential to provide guiding principles for the design of high-performance NASICON conductors in such a large, yet partially uncharted chemical space.

In this study, we combine ab-initio calculations with text-mined data on the ionic conductivity of NASICONs from a corpus of several

million papers and our experimental work to disentangle the factors that control the Na-ion conductivity in NASICONs. High-throughput density-functional-theory (DFT) calculations are first used to investigate the stability of NASICONs. A subset of the predicted compositions is experimentally explored, leading to the successful synthesis of eight NASICONs. Five of these were introduced in our recent work³⁵ and three are new. In this paper we study the electrochemical performance of these eight recently discovered compounds as solid-state conductors, and present detailed crystal structure and electrochemical impedance. The ionic conductivity and activation energy of the as-synthesized NASICONs were investigated, and high ionic conductivity (~10⁻⁴ S cm⁻¹) at room temperature (~25 °C) was found in Na₃HfZr(SiO₄)₂(PO₄) and Na₃HfSc(SiO₄)(PO₄)₂. Natural language processing (NLP) tools were applied to text-mine NASICON literature and identify the trends of the ionic conductivity evolution with respect to different compositional variables. Guided by this text-mined information, we improved the NASICON room-temperature (~25 °C) ionic conductivity in this work up to 1.2 mS cm⁻¹, which is one of the highest conductivities among reported NASICONs. Ab-initio molecular dynamic (AIMD) simulations with well-controlled compositional variables revealed the following comprehensive compositional optimization strategies for NASICON conductors: (1) the optimal Na content is -x = 3 but depends on the polyanion chemistry, (2) high ionic conductivity is found in compositions with a large cation size until the size reaches an optimal value, and (3) high silicate content enhances the ionic conductivity. Our work provides a comprehensive guiding map to facilitate the discovery of new NASICONs and insights into the optimization of NASICON ionic conductivity toward the application of solid-state Na batteries.

Results

Map of synthetically accessible NASICONs predicted by highthroughput DFT calculations

High-throughput ground-state calculations were conducted to evaluate the phase stability of the NASICON structure across a wide range of compositions. The screening criteria are summarized in Fig. 1b. We considered the general formula $Na_xM_vM'_{2-v}$ $(AO_4)_z(BO_4)_{3-z}$, i.e., both the metal site and polyanion site can be occupied by up to two species. In particular, we considered SiO₄⁴, PO₄³ and SO₄² as possible polyanions and 16 possible electrochemical inactive metals for M and M'. (Fig. 1b). As the Na content x can take values from 0 to 4.0, y can take values from 0 to 2.0, and z can take values from 0 to 3.0. Using intervals of 0.5 for sampling these composition variables, 3881 charge-balanced compositions were enumerated, and their energy above the convex hull (E_{hull}) in the relevant chemical space was calculated using DFT. To guide the experimental exploration at high temperature³⁷, we further computed the ideal configurational entropy (S_{ideal}, see Methods) of each composition. We obtained 641 NASICONs with $E_{hull} - S_{ideal}T \le 0$ (T=1000 K), making them likely to be synthetically accessible. To narrow the target to possible solid-state conductors, SO_4^{2-} -containing NASICONs were excluded at this point because they are generally less stable at high temperature and therefore challenging to densify²⁸. Of the 242 remaining SO₄²⁻-free NASICONs, only 23 compounds with Na₃ stoichiometry were further considered, within which the electrochemical performance of 17 compounds has not yet been reported (Supplementary Table 1). Finally, 8 of these NASICON compounds were successfully synthesized through a classic solid-state synthesis. The distribution of the 641 NASICONs predicted to be synthetically accessible is presented in Fig. 1c, sorted by their metal chemistry. The color intensity and number in each box indicate the number of stable compounds found that contain both metal elements. The heat map is further annotated as follows: (1) blocks that contain at least one Na₃ compound are highlighted with orange rectangles; (2) blocks that only have sulfate NASICONs, and therefore not included in the experimental study, are marked with grey rectangles.

Experimental exploration of the predicted NASICONs as potential solid conductors

Experimental exploration via solid-state methods of the 17 NASICONs mentioned above led to the synthesis of 8 NASICON compounds. Supplementary Table 1 summarizes all the synthesis attempts, including failures. Although many silicate NASICONs containing metals with 5+ oxidation state were predicted to be synthesizable, they tended to form NaM $^{\rm 5+}O_3$ competing phases. Some failed synthesis attempts also involved the melting of precursors, even at relatively low temperatures (-600 °C). Note that for all the synthesis attempts, only the calcination temperatures were varied (600–1100 °C) to optimize the synthesis outcome; therefore, other variables such as the precursor selection, mixing, calcination atmosphere, or even alternative synthesis routes might be considered to access those phases $^{\rm 18-40}$.

X-ray diffraction (XRD) patterns of the successfully synthesized NASICONs are presented in Fig. 2. Among the 8 compositions, only $Na_3HfZrSi_2PO_{12}$ has a monoclinic structure (C2/c), whereas all the others crystallize in a rhombohedral (R-3c) structure. Representative Rietveld refinement profiles for the monoclinic ($Na_3HfZrSi_2PO_{12}$) and rhombohedral ($Na_3HfScSiP_2O_{12}$) structures are presented in Figs. 2a and 2b, respectively, both of which show a small amount of impurity phase, consistent with previous findings in the literature⁴¹. Supplementary Fig. 1 presents the refined XRD patterns for other NASICONs, with the extracted lattice parameters summarized in Supplementary Table 2–9. In general, the c lattice parameter is mainly affected by the polyanion species, which determine the unit-cell volume, with a larger volume for increasing silicate content. The a lattice parameter, instead, is primarily controlled by the metal species.

Dense pellets of the as-synthesized materials were prepared (Methods and Supplementary Table 10) for the electrochemical impedance spectroscopy (EIS) tests. The bulk and grain-boundary

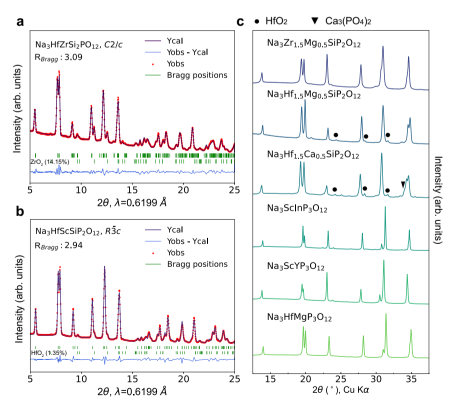


Fig. 2 | **XRD patterns of the as-synthesized NASICONs.** Rietveld refinement results of synchrotron XRD data of Na₃HfZrSi₂PO₁₂ with monoclinic *C2/c* symmetry (a), and of Na₃HfScSiP₂O₁₂ with rhombohedral *R-3c* symmetry (b). c Lab XRD

patterns of other as-synthesized NASICONs. The corresponding refinement results are presented in Supplementary Fig. 1, and structural parameters in Supplementary Table 2-9.

conductivities at room temperature (~25 °C) are summarized in the top panel of Fig. 3, and the corresponding Nyquist plots, equivalent circuits and fitting parameters are presented in Supplementary Fig. 3-4, Supplementary Table 11, respectively. In Fig. 3, the compounds are grouped by their polyanion chemistry, i.e., phosphates (PO₄)₃ and mixed silicate-phosphates (SiO₄)(PO₄)₂ and (SiO₄)₂(PO₄). In each polyanion group, compounds are sorted by ascending average metal radius (\bar{r}_M) . In general, the ionic conductivity increases as the silicate content increases¹⁹, irrespective of the cations. Specifically, the total conductivities of the pure phosphate group are in the range of 10⁻⁶ to 10^{-5} S cm⁻¹, whereas those of the (SiO₄)(PO₄)₂ group are in the range of 10^{-5} to 10^{-4} S cm⁻¹. The highest ionic conductivity of 4.4×10^{-4} S cm⁻¹ is achieved in Na₃HfZr(SiO₄)₂(PO₄). Within each polyanion group, the ionic conductivity shows a non-monotonic trend with \bar{r}_M . Indeed, the ionic conductivity increases with \bar{r}_M only up to an optimal value. For example, the ionic conductivity increases from 2.40×10^{-6} to 6.48×10^{-5} S cm⁻¹ in going from Na₃HfMg(PO₄)₃ ($\bar{r}_M = 0.715 \text{ Å}$) to Na₃ScIn(PO₄)₃ $(\bar{r}_M = 0.773 \text{ Å})$ and from $6.17 \times 10^{-5} \text{ S cm}^{-1}$ for $Na_3Hf_{1.5}Mg_{0.5}(SiO_4)(PO_4)_2$ $(\bar{r}_M = 0.713 \text{ Å})$ to $1.87 \times 10^{-4} \text{ S cm}^{-1}$ for $Na_3HfSc(SiO_4)(PO_4)_2$ $(\bar{r}_M = 0.728 \text{ Å})$. Further increasing \bar{r}_M is unfavorable with both polyanion compositions, as Na₃ScY(PO₄)₃ ($\bar{r}_M = 0.823 \,\text{Å}$) and $Na_3Hf_{1.5}Ca_{0.5}(SiO_4)(PO_4)_2$ ($\bar{r}_M = 0.783 \text{ Å}$) appear to have lower conductivities of 4.10×10⁻⁶ and 2.93×10⁻⁵ S cm⁻¹, respectively. The refined compositions are listed in Supplementary Table 2 - 9, and agree reasonably with the nominal composition, so that the trend of the average cation radius presented in Fig. 3 remains valid. Therefore, nominal compositions are used in the analysis for simplicity. However, Na content <3 are observed for the refined compositions, probably due to the Na loss during the high temperature annealing.

The ionic conductivity of each NASICON was measured at various temperatures (Supplementary Fig. 6), revealing typical Arrhenius behavior, and the extracted activation energies are plotted in the bottom panel of Fig. 3. Consistent with the trend in the conductivities, as the silicate content increases, the activation energy generally decreases. Similarly, the activation energy is reduced when large M

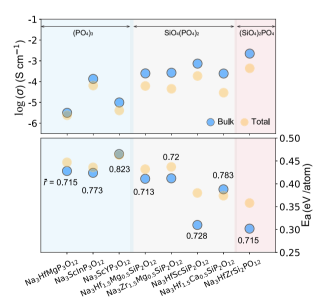


Fig. 3 | **Measured ionic conductivity of as-synthesized NASICONs.** Scatter plots of room-temperature (-25 °C) conductivities (log scale, upper panel) fitted from Nyquist plots (Supplementary Fig. 3) and activation energies (lower panel) extracted from the linear fitting of the Arrhenius plots (Supplementary Fig. 6). The bulk properties are indicated by blue circles, whereas the total properties are indicated by orange ones. On the x-axis, the compounds are grouped by polyanion chemistry, and in each group, compounds are ordered by increasing average metal radius (\bar{r}_M , in Å). \bar{r}_M of each compound is also labeled in the bottom panel.

cations are present, though there is a limit to the benefits of this size effect. Na₃HfZr(SiO₄)₂(PO₄), which has the highest ionic conductivity, also exhibits the lowest bulk activation energy of 0.302 eV. These experimental observations indicate that for our as-synthesized NASI-CONs, the ionic conductivity is primarily determined by the activation energy. It is worth noting that though the nominal Na content of the assynthesized compounds are all equal to 3, the ratio between Na1, Na2 site occupancies can be different (Supplementary Table 2–9, Supplementary Fig. 7). The Na1 site occupancy decreases as the silicate content increases¹⁹: the three NASICONs with pure phosphates have Na1 occupancy of -0.8, whereas the monoclinic Na₃HfZr(SiO₄)₂(PO₄) phase with the highest ionic conductivity exhibits the lowest Na1 occupancy of -0.63.

Further compositional optimization based on text-mined ionic conductivity data

To generalize our findings, we investigated a text-mined dataset containing the experimentally measured ionic conductivity of 475 reported NASICONs. The corresponding papers were identified by screening over two million materials science articles with criteria for specific materials (i.e., NASICONs) and properties (i.e., ionic conductivity) via chemical named entity recognition⁴²⁻⁴⁴. Three compositional variables were used to visualize the distribution of the ionic conductivity data obtained from the literature: the Na content (x), average metal radius (\bar{r}_M) , and average A site radius (\bar{r}_A) . As the ionic radius of Si⁴⁺ (0.26 Å) is larger than that of P^{5+} (0.17 Å), the variation of the silicate content in Fig. 3 can be appropriately reflected by \bar{r}_A . Therefore, a threedimensional compositional space of the NASICONs was constructed, as shown in Fig. 4a, with the color of each point indicating its ionic conductivity on a log scale. Overall, the ionic conductivity of NASI-CONs varies greatly with composition, ranging from 10⁻¹⁴ to 10⁻³ S cm⁻¹. However, this finding should be taken with caution because some of this difference may also result from inconsistent sample preparation or testing methods between the various literature studies. In particular, pellet density and varying interfacial resistance may affect the measured ionic conductivity. These experimental details are often not explicitly reported in the literature¹⁹. In Fig. 4a, most high ionic conductivity values can be found in certain narrow subspaces. For instance, two thin sections with $0.70 \le \bar{r}_M \le 0.74$ Å and $2.5 \le x \le 3.5$ are highlighted, respectively, and the projections of data points within those two sections are plotted in Fig. 4b and Fig. 4c.

Figure 4b indicates that, within the group of pure phosphates $(\bar{r}_A = 0.17 \text{ Å})$, the high-ionic-conductivity compositions have Na content around x = 2.5 (though the limited number of Na-rich phosphates that have been reported limit the confidence level of this finding). As the silicate content increases (\bar{r}_A increasing), the Na content, at which point the ionic conductivity is maximized shifts toward higher values. For NASICONs with silicate contents equal to or larger than two (per formula unit of three polyanion groups), the optimal Na content is between 3.0 and 3.4. Note that the optimal Na content has already been investigated in previous studies 19,23,36,45. However, we argue that the optimal value might be different for different polyanion compositions, and in the following section we investigate the Na content effect in more depth by setting other compositional variables constant in AIMD simulations. Figure 4c shows that the maximum ionic conductivity occurs when \bar{r}_M is slightly above 0.72 Å (though the exact optimal value might be dependent on polyanion species), whereas NASICONs with a \bar{r}_M that is too small or too large are less likely to exhibit high ionic conductivity. Some exceptions are found among phosphate NASICONs with large lanthanide elements such as Er or Dy as doping cations that exhibit high ionic conductivity46, however, the temperature at which EIS measurements were taken was not explicitly mentioned.

Among all the NASICONs synthesized in our limited set of experiments, Na₃HfZrSi₂PO₁₂ exhibits the highest ionic conductivity.

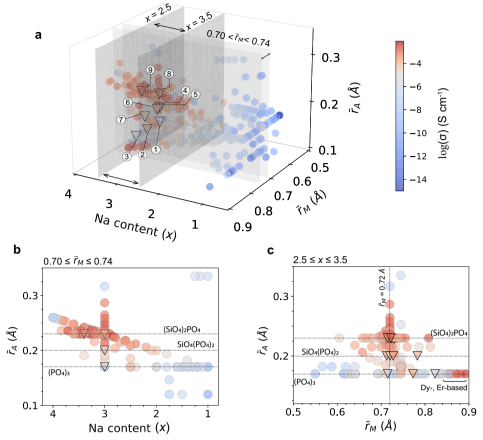


Fig. 4 | **Reported room-temperature (~25 °C) NASICON conductivities as a function of compositional variables. a** Three-dimensional scatter plot of the reported NASICON conductivities as a function of Na content (x), average M site radius (\bar{r}_M) , and average A site radius (\bar{r}_A) . The ionic conductivity (on a log scale) of each NASICON is represented by the color bar. **b** Projection of data points between $\bar{r}_M = 0.70$ Å and $\bar{r}_M = 0.74$ Å in **a** on the Na content $\times \bar{r}_A$ plane. **c** Projection of the

data points between x = 2.5 and x = 3.5 in **a** on the $\overline{r}_M \times \overline{r}_A$ plane. The as-synthesized NASICONs in this work are represented by triangles: 1: Na₃HfMg(PO₄)₃, 2: Na₃ScIn(PO₄)₃, 3: Na₃ScY(PO₄)₃, 4: Na₃Hf_{1.5}Mg_{0.5}(SiO₄)(PO₄)₂, 5: Na₃Zr_{1.5}Mg_{0.5}(SiO₄)(PO₄)₂, 6: Na₃HfSc(SiO₄)(PO₄)₂, 7: Na₃Hf_{1.5}Ca_{0.5}(SiO₄)(PO₄)₂, 8: Na₃HfZr(SiO₄)₂(PO₄), 9: Na_{3.4}Hf_{0.6}Sc_{0.4}ZrSi₂PO₁₂.

However, the data in Fig. 4 indicate that the ionic conductivity might still be further improved by modifying the average cation radius and stoichiometry of Na₃HfZrSi₂PO₁₂. Therefore, we attempted to use Sc³⁺ as a large-size dopant with a lower valence to bring both the average cation radius and Na content toward the optimized values. By replacing 0.2, 0.4 Hf⁴⁺ per f.u. with Sc³⁺, the average cation radius can be increased from 0.715 Å to 0.719 and 0.722 Å, and the Na content to 3.2 and 3.4, respectively. Figure 5a presents the XRD patterns of Scsubstituted Na_{3,2}Hf_{0.8}Sc_{0.2}ZrSi₂PO₁₂ and Na_{3,4}Hf_{0.6}Sc_{0.4}ZrSi₂PO₁₂. Both compounds show a majority of NASICON phase with a small amount of ZrO₂ impurity. As the Sc content increases, Na_{3.4}Hf_{0.6}Sc_{0.4}ZrSi₂PO₁₂ no longer retains the monoclinic (C2/c) symmetry and can be properly refined with a rhombohedral model (Supplementary Fig. 5). EIS spectra of both compounds using Na metal as electrodes are presented in Compared with pristine Na₃HfZrSi₂PO₁₂, $Na_{3.2}Hf_{0.8}Sc_{0.2}ZrSi_2PO_{12}$ and $Na_{3.4}Hf_{0.6}Sc_{0.4}ZrSi_2PO_{12}$ exhibit higher total ionic conductivity of 0.48 and 1.2 mS cm⁻¹, respectively. In particular, 1.2 mS cm⁻¹ is one of the highest conductivities among reported NASICON conductors at room temperature (~25 °C)^{47,48}. It is worth noting that the ionic conductivity of this material may still be improved by increasing the silicate content, as inferred from the experimental and text-mined data, or by optimizing the sintering condition via hot pressing or spark plasma sintering techniques, instead of the coldpressing method used in this work. However, the former essentially requires the incorporation of high-valence charge compensators, which may be inhibited by the challenges related to the synthesis of

materials with 5+ metals and high silicate content (Supplementary Table 1).

The stability of Na $_{3.4}$ Hf $_{0.6}$ Sc $_{0.4}$ ZrSi $_2$ PO $_{12}$ against a Na-metal anode was evaluated by constructing a Na | Na $_{3.4}$ Hf $_{0.6}$ Sc $_{0.4}$ ZrSi $_2$ PO $_{12}$ | Na symmetric cell. As shown in Fig. 5c, stable Na stripping and plating are achieved, and the small overpotentials are stable throughout the 200-h test. EIS spectra of the Na-metal symmetric cell were collected after 100-h and 200-h cycling, showing no obvious change in the overall impedance (Supplementary Fig. 9), indicating that the NASICON–Na metal interface is stable.

Discussion

Our experimental analysis, combined with the large-scale extraction of ionic conductivity data from the literature, establishes optimal ranges for Na content, polyanion chemistry, and average metal radius. To better understand how these various compositional variables affect the ionic conductivity, ab initio molecular dynamic (AIMD) simulations were performed, and the results are presented in Fig. 6. Three different NASICON groups with different controlled variations in composition were designed so as to separate the effect of each variable as independently as possible:

1. Na-content variation: The polyanion chemistry per formula unit was fixed to be either $(PO_4)_3$ or $(PO_4)_2(SiO_4)$ with the average cation radius kept near 0.715 Å while varying the Na content by modifying the ratio between Hf^{4+} (0.71 Å) and Mg^{2+} (0.72 Å) with similar size.

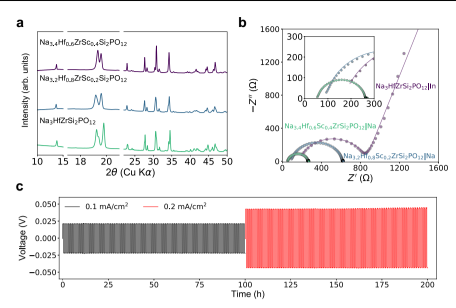


Fig. 5 | **Optimization of ionic conductivity via Sc doping. a** XRD patterns. The peaks between 18° and 20° are highlighted to show the change in crystal symmetry. **b** Nyquist plots obtained from EIS at room temperature (-25 °C). Indium metal was used as blocking electrodes for Na₃HfZrSi₂PO₁₂, whereas Na metal was used as the

electrodes for Na $_{3.2}$ Hf $_{0.8}$ Sc $_{0.2}$ ZrSi $_2$ PO $_{12}$ and Na $_{3.4}$ Hf $_{0.6}$ Sc $_{0.4}$ ZrSi $_2$ PO $_{12}$. c Na stripping and plating tests in a temperature chamber at 25 °C at various current rates of a Na| Na $_{3.4}$ Hf $_{0.6}$ Sc $_{0.4}$ ZrSi $_2$ PO $_{12}$ | Na symmetric cell.

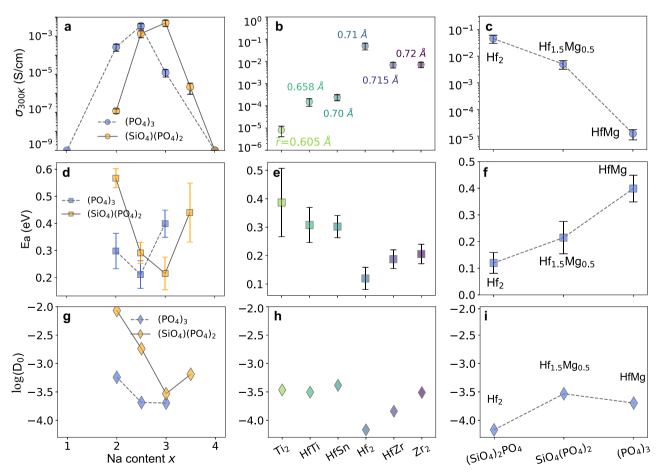


Fig. 6 | Bulk ionic conductivities, activation energies, and prefactors calculated by AIMD. Calculated room-temperature (300 K) bulk ionic conductivities (a) and activation energies (d) with error bars⁷⁷, and prefactors (g) of Na_{1+2y}Hf_{2-y}Mg_yP₃O₁₂ (y = 0, 0.5, 0.75, 1, 1.5, labeled blue) and Na_{2+2z}Hf_{2-z}Mg_zSiP₂O_{1z} (z = 0, 0.25, 0.5, 0.75, 1, labeled yellow). In these two groups, both the polyanion composition and cation radius are kept constant. Note that for

compounds with y=0, 1.5 and z=1, the conductivities were not converged and therefore set as « 10^{-9} mS cm⁻¹. **b**, **e**, **h** Group of Na₃MMSi₂ PO_{12} (M, M'=Ti, Hf, Sn, Zr), with constant Na content x and polyanion composition. **c**, **f**, **i** Group of Na₃ $Hf_{2-z}Mg_zSi_{2-2z}P_{1+2z}O_{12}$ (z=0, 0.5, 1) where \bar{r}_M and the Na content x are kept constant. The error bars quantifying the statistical confidence levels of diffusion results from AIMD are calculated according to He et al. 77 .

- Variation of the cation size: The Na content was fixed to be Na₃ per formula unit with (PO₄)(SiO₄)₂ as the polyanion while varying the average cation radius by modifying the combination of different M⁴⁺ metal cations.
- 3. Polyanion composition: The Na content was fixed to be Na₃ per formula unit with the cation radius kept at -0.715 Å while modifying the polyanion and charge compensating through the ratio between Hf⁴⁺ (0.71 Å) and Mg²⁺ (0.72 Å).

Figure 6 shows the AIMD calculated ionic conductivity at 300 K (Fig. 6a-c), the activation energy (Fig. 6d-f), and the prefactor for Na diffusion (Fig. 6g-i) for the three compound groups. The correlations between the ionic conductivity and compositional variables in Fig. 6a-c are generally consistent with our experimental and textmined ionic conductivity trends. Regarding the Na content effect, in Figs. 6a and 6d, conductivities and activation energies vary simultaneously with Na content, indicating that Na content not only determines the charge-carrier concentration, as expected, but also affects the ionic conductivity through the migration barrier. The correlation between Na content and migration barrier can be rationalized as follow: the migration barrier is determined by the size of the bottleneck for ion hopping, which is then controlled by the lattice parameter. For NASICONs with 1Na per formula unit, Na ion sits in the Na1 (6b) site that is face-sharing with two MO_{6} , as shown in Fig. 1a. Increasing Na content will displace the Na⁺ into the Na2 (18e) site, which results in reduced occupancy of the Na1 site and increased electrostatic repulsion between MO₆ units, leading to larger unit cell volume and therefore bottleneck size³⁷. This trend is supported by the correlation between Na content and lattice parameter, as shown in Supplementary Fig. 10. Furthermore, consistent with the text-mined data in Fig. 4b, Fig. 6a also shows that the optimal Na content for ionic conductivity is polyanion dependent: for phosphates the optimal value is 2.5 while for mixed polyanions the optimal value is near 3. Correspondingly in Fig. 6d, the Na content that is optimal for lowering the migration barrier is also lower for phosphates than for mixed polyanions. This result indicates that the optimal Na content for different polyanion composition appears to be mostly driven by Na's effect on the activation energy rather than by its direct relation to the carrier concentration.

The correlation between the cation radius and activation energy. displayed in Fig. 6e, is consistent with our experimental observation, i.e., the activation energy generally decreases with increasing cation radius but becomes mostly insensitive once surpassing the optimal value. Our finding is also in line with previous studies indicating that an optimal cation radius exists^{23,47}, and here we demonstrate this effect more rigorously by keeping other compositional variables constant. The results in Fig. 6f show that by keeping the Na content and cation radius constant, the activation energy can be reduced up to ~200 meV solely by increasing the silicate content from 0 to 2/3, implying that the highest ionic conductivity might be obtained in pure silicate NASI-CONs or NASICONs with SiO₄⁴⁻ content above 2/3. However, the experimental validation of this optimum is challenging as synthesis attempts for this type of NASICON were unsuccessful via the solidstate method (Supplementary Table 1). Increasing the Na content, metal radius, and/or Si content all lead to the increase of the lattice constant of NASICON and thereby enhance the ionic conductivity. This observation would indicate that Na mobility is mostly controlled by the size of some migration bottleneck, and not by other effect such as cation-cation interaction⁴⁹⁻⁵², topology^{5,51,53,54}, or polyanion-assisted motion55-60.

While the dominant effect of the compositional variables is likely the control of the size of the ion migration bottleneck and, therefore, the activation energy, the composition also has a more subtle effect on the ordering tendency of Na ions and lattice softness of the NASICON framework, which is reflected in the variation of the prefactor for Na diffusion. The evolution of the prefactor differs from that of the activation energy and ionic conductivity. Generally, it becomes smaller as

the ionic conductivity becomes larger, consistent with prior findings in the literature⁶¹⁻⁶³. When tuning Na content and metal radius (Figs. 6g. 6h), the prefactor is minimal when the ionic conductivity is maximal. consistent with a reduction in attempt frequency for ion migration when the compound softens due to the increase in lattice parameter. The polyanion chemistry shows less (inverse) correlation with the activation energy as shown in Fig. 6i. Such a divergent evolution suggests that the prefactor can be used to fine-tune the ionic conductivity. Even though the effect of the prefactor can be neglected in the composition region where the magnitude of the ionic conductivity is primarily determined by the activation energy, when the activation energy is reduced to a sufficiently low value (e.g., ~300 meV), the prefactor should be taken into consideration to further enhance the ionic conductivity. While it is generally assumed that changes in the prefactor are related to a softening of the relevant phonon modes that determine the attempt frequency for hopping, they may also reflect variations in effective carrier concentration caused by more or less local Na ordering.

In summary, through experiments, text-mined literature data, and ab-initio modeling, we have quantified the relationship between the various composition variables of NASICON compounds and the Na-ion conductivity. We find that Na-ion conductivity can be optimized by: (1) an average cation radius of slightly above 0.72 Å for NASICONs with a $(\mathrm{SiO_4})_2\mathrm{PO_4}$ polyanion composition; (2) high $\mathrm{SiO_4}^4$ polyanion content, and (3) a carefully tuned Na content around Na₃ per formula unit but depending on the polyanion composition. In addition, by constructing a first-principles synthesis map for the possible chemical space of NASICON compounds, eight NASICONs were successfully synthesized and investigated as solid-state conductors, and room-temperature (-25 °C) ionic conductivity of 1.2 mS cm $^{-1}$ was achieved in Na_{3.4}Hf_{0.6}Sc_{0.4}ZrSi₂PO₁₂.

Methods

Experiments

For the solid-state synthesis of the NASICON compounds, typical metal oxides or hydroxides (HfO₂ (Aldrich, 99.8%), MgO (Aldrich, ≥99.99%), Sc₂O₃ (Sigma, 99.9%), Zr(OH)₄ (Aldrich, 97%), SnO₂ (Alfa Aesar, 99.9%), CaO (Sigma-Aldrich, 99.9%), In₂O₃ (Sigma, 99.998%), Y₂O₃ (Sigma, nanopowder, <50 nm particle size)) were used as precursors to introduce metal cations. SiO₂ (Sigma-Aldrich, nanopowder) and NaH₂PO₄ (Sigma, >99%) were used as silicate and phosphate sources. Na₂CO₃ (Sigma-Aldrich, >99%) was used as an extra sodium source. In addition, 10% excess NaH₂PO₄ was introduced to compensate for the possible sodium and phosphate loss during the high-temperature treatment. The powder mixtures were wet ball-milled (3 mL ethanol per 50 mL jar) with stainless steel balls and jar for 12 h using a planetary ball mill (PM200, Retsch) at 250 rpm for thorough mixing before pressing into pellets. The pelletized samples were first annealed to form the target phase, then grounded with a mortar and pestle, and wet ball-milled (3 mL ethanol per 50 mL jar) again using a high-energy ball mill (SPEX 8000 M Mixer/Mill) with zirconia balls and jar to reduce the particle size. The powder was then dried and pressed into pellets with a uniaxial press. The pellets were again pressed using a cold isostatic press (YLJ-CIP-20B, MTI), then wrapped with Pt foil for the second hightemperature annealing in an alumina crucible. The detailed annealing conditions for the first and second steps are provided in Supplemen-Table 10. Synchrotron powder diffraction data $Na_3HfZr(SiO_4)_2(PO_4)$, $Na_3HfSc(SiO_4)(PO_4)_2$, $Na_{3.4}Hf_{0.6}Sc_{0.4}ZrSi_2PO_{12}$ were collected at beamline 7-BM, National Synchrotron Light Source II, Brookhaven Nation Lab using an average wavelength of 0.6199 Å. Other obtained materials were analyzed using Rigaku Miniflex 600, Bruker D8 Diffractometer with Cu Kα radiation, scanned over 10–70° two theta range, with data points collected every 0.02° 2θ and scan speed of ~0.005° s⁻¹. Data analysis was carried out via Rietveld refinement⁶⁴ using Fullprof program⁶⁵. Certain steps were carried out during the refinements with regard to the site occupancies: (1) First the atomic occupancies were set to their nominal compositions. (2) The occupancy of M1. M2 were refined with the constraint that their occupancies sum to 2: (3) The occupancy of Na1, Na2 (and Na3 for monoclinic NASICON) were then refined freely; (4) To ensure the charge neutrality, with the refined occupancy of Na1, Na2, M1 and M2. the Si, P occupancies were manually calculated with the constraint of their occupancies summing to 3; (5) We applied the manually calculated Si, P occupancies to the refined model, and performed step (2) and step (3) again. During this step, the change of Na1, Na2, M1 and M2 occupancies were <1%, therefore charge neutrality was maintained. For pure phosphate compounds, the P occupancy was fixed at 3, and the occupancy of Na1, Na2, M1, M2 were refined with constraints that charge neutrality was ensured, i.e., for Na₃ScYP₃O₁₂, Sc_{Occ} + Y_{Occ} = 2, $Na1_{Occ} + Na2_{Occ} = 3$; $Na_3ScInP_3O_{12}$, $Sc_{Occ} + In_{Occ} = 2$, $Na1_{Occ} + Na2_{Occ} = 3$; for Na₃HfMgP₃O₁₂, Hf_{Occ} + Mg_{Occ} = 2, Δ Hf_{Occ} + Δ Na1_{Occ} + Δ Na2_{Occ} = 0.

The ionic conductivity was evaluated using EIS at temperatures ranging from ~0 to ~100 °C. EIS analysis was performed using a Bio-Logic VMP-300 system at the initial open-circuit voltage in the frequency range from 7 MHz to 100 mHz with the application of a 10 mV signal amplitude. The measurements were performed using a BioLogic controlled environment sample holder assembled and sealed in an Arfilled glovebox. Indium foils were directly pressed onto both sides of the sample pellet as blocking electrodes. Sample pellets are ~ 6 mm in diameter and ~ 1 mm in thickness. For the Sc-doped samples, Na foils were pressed onto both sides of the pellet in a Ar-filled glove box. The Na|NASICON|Na symmetric cell was placed in a customized solid-state pressure cell with an internal pressure of 3 MPa. EIS and cell cycling tests were performed in sequence using the same cell.

The bulk and total ionic conductivities were obtained by fitting the Nyquist plot with two theoretical circuit models: (1) 1 R (bulk) + 1RC (grain boundary) + C for Na₃HfZrSi₂PO₁₂ and Na₃HfScSiP₂O₁₂ and (2) 2RC (bulk, grain boundary) + C for the others (R: resistance, C: constant phase element, RC: resistance and constant phase element in parallel.) For the Sc-doped samples, only the total resistance (ionic conductivity) was estimated by taking the low-frequency intercept of the semi-circle on the x-axis.

First-principles calculations

First-principles total energies calculations were performed using the Vienna ab initio simulation package (VASP) with a plane-wave basis set66. Projector augmented-wave potentials67 with a kinetic energy cutoff of 520 eV and the exchange-correlation form in the Perdew-Burke-Ernzerhof generalized gradient approximation (GGA-PBE)⁶⁷ were employed for all the structural optimizations and total energy calculations. For all the calculations, a reciprocal space discretization of 25 k-points per Å⁻¹ was applied, and the convergence criteria were set to 10⁻⁶ eV for electronic iterations and 0.02 eV/Å for ionic iterations. A rhombohedral conventional cell was used for each of the NASICON structures. The Na-vacancy ordering, cation ordering, and anion ordering were set as the one with the lowest electrostatic energy⁶⁸⁻⁷². It should be noted that NASICONs can also exist with other space groups, such as the monoclinic form (C2/c). The monoclinic NASICONs can be regarded as an ordered version of rhombohedral NASICONs⁷³. Therefore, the energy difference between rhombohedral and monoclinic NASICONs is expected to be low as it relates to the Na disordering energy. To provide an estimate, the disordering energy for Na is typically 30–40 meV, while the contribution to E_{hull} must be normalized by the number of atoms per formula unit, i.e., ~30–40 meV/(17–21 atoms per formula unit). This energy variation is much smaller than that caused by chemistry variation. With such considerations, all our calculations and analysis were only performed for rhombohedral NASICONs. In addition to the 3881 NASICON structures, all the competing phases in the relevant chemical spaces that are given in The Materials Project⁷⁴ were also calculated to

construct the phase diagrams. To estimate the synthetic accessibility of the NASICON at finite temperature, we included the ideal configurational entropy and assumed that a NASICON is synthetically accessible when $E_{hull} - TS_{Ideal} \le 0$. The ideal entropy was calculated by assuming a fully disordered distribution of Na, cation, and anion sites. For a NASICON with the chemical formula $Na_x M_y M'_{2-y} (AO_4)_z (BO_4)_{3-z}$, the ideal entropy is calculated as follows with the units of eV/atom:

$$S_{ideal} = -k_b \frac{1}{x+17} \left(\frac{1}{4} \left(\frac{x}{4} \ln \left(\frac{x}{4} \right) + \left(1 - \frac{x}{4} \right) \ln \left(1 - \frac{x}{4} \right) \right) + \frac{1}{2} \left(\frac{y}{2} \ln \left(\frac{y}{2} \right) + \left(1 - \frac{y}{2} \right) \ln \left(1 - \frac{y}{2} \right) \right) + \frac{1}{3} \left(\frac{z}{3} \ln \left(\frac{z}{3} \right) + \left(1 - \frac{z}{3} \right) \ln \left(1 - \frac{z}{3} \right) \right) \right)$$
(1)

While this is likely an overestimation of the entropic stabilization that can be achieved, this more generous filter allows us to capture NASICONs which would be synthesizable as metastable phases.

Ab initio molecular dynamic (AIMD), initialized from the DFT relaxed electrostatic ground states of each NASICON composition, was used to investigate the ionic conductivity. All the AIMD calculations were performed in an NVT ensemble with a time step of 2 fs using a Nosé–Hoover thermostat⁷⁵ for a period of 160 ps. A minimal Γ -point-only k-point grid was used for all the calculations. The AIMD simulations were run at 600 K, 800 K, 1000 K, 1200 K, and 1500 K. All the data were fitted assuming Arrhenius behavior to obtain the activation energy, diffusion prefactor, and room-temperature (-25 °C) diffusivity.

Text mining for NASICON conductivity data

In order to gather NASICON conductivity data spanning a variety of chemical systems from scientific literature, we employed a hybrid strategy combining named entity recognition methods along with human assistance. Initially, NASICON-related papers were identified by applying named entity recognition algorithms 43,44,76 and specific selection criteria to over two million material science papers, including (1) any mention of "NASICON" or known NASICON compositions within abstracts and experimental sections and (2) any mention of "conductivity" or properties associated with conductivity in the abstract. In each NASICON-related paper, the NASICON conductivity data were manually curated from tables and figures when available, accounting for the ionic conductivity σ and activation energy E_a corresponding to different temperatures. When high-temperature conductivity and activation energy rather than room-temperature conductivity were reported, the Arrhenius relationship

$$\sigma T = A \exp\left(-E_a/kT\right) \tag{2}$$

was used to project the high temperature conductivity to 300 K, where A is the pre-exponential factor and k is the Boltzmann constant. The compiled NASICON conductivity data have been included in the Source Data file.

Data availability

Source data are provided with this paper. Other relevant data are available from the corresponding authors upon reasonable request. Source data are provided with this paper.

References

- Tian, Y. et al. Promises and challenges of next-generation "Beyond Li-ion" batteries for electric vehicles and grid decarbonization. Chem. Rev. 121, 1623–1669 (2021).
- Xiao, Y. et al. Understanding interface stability in solid-state batteries. Nat. Rev. Mater. 5, 105–126 (2020).
- Manthiram, A., Yu, X. & Wang, S. Lithium battery chemistries enabled by solid-state electrolytes. Nat. Rev. Mater. 2, 1–16 (2017).
- Famprikis, T., Canepa, P., Dawson, J. A., Islam, M. S. & Masquelier, C. Fundamentals of inorganic solid-state electrolytes for batteries. Nat. Mater. 18, 1278–1291 (2019).

- Wang, Y. et al. Design principles for solid-state lithium superionic conductors. Nat. Mater. 14, 1026–1031 (2015).
- Zhou, L. et al. Solvent-engineered design of argyrodite Li6PS5X (X = Cl, Br, I) solid electrolytes with high ionic conductivity. ACS Energy Lett. 4, 265–270 (2018).
- Park, K.-H. et al. High-voltage superionic halide solid electrolytes for all-solid-state Li-ion batteries. ACS Energy Lett. 5, 533–539 (2020).
- 8. Zhang, Z. et al. Na 11 Sn 2 PS 12: a new solid state sodium superionic conductor. *Energy Environ. Sci.* **11**, 87–93 (2018).
- Adeli, P. et al. Boosting solid-state diffusivity and conductivity in lithium superionic argyrodites by halide substitution. *Angew. Chem.* 131, 8773–8778 (2019).
- Hong, H.-P. Crystal structures and crystal chemistry in the system Na1+ xZr2SixP3- xO12. Mater. Res. Bull. 11, 173-xO182 (1976).
- Goodenough, J. B., Hong, H.-P. & Kafalas, J. Fast Na+-ion transport in skeleton structures. *Mater. Res. Bull.* 11, 203–220 (1976).
- Park, H., Jung, K., Nezafati, M., Kim, C.-S. & Kang, B. Sodium ion diffusion in Nasicon (Na3Zr2Si2PO12) solid electrolytes: effects of excess sodium. ACS Appl. Mater. interfaces 8, 27814–27824 (2016).
- Shimizu, Y. & Yamashita, N. Solid electrolyte CO2 sensor using NASICON and perovskite-type oxide electrode. Sens. Actuators B: Chem. 64, 102–106 (2000).
- Anantharamulu, N. et al. A wide-ranging review on Nasicon type materials. J. Mater. Sci. 46, 2821–2837 (2011).
- 15. Jian, Z., Hu, Y. S., Ji, X. & Chen, W. Nasicon-structured materials for energy storage. *Adv. Mater.* **29**, 1601925 (2017).
- Fabry, P., Gros, J., Million-Brodaz, J. & Kleitz, M. NASICON, an ionic conductor for solid-state Na+-selective electrode. Sens. Actuators 15, 33–49 (1988).
- Tang, B., Jaschin, P. W., Li, X., Bo, S.-H. & Zhou, Z. Critical interface between inorganic solid-state electrolyte and sodium metal. *Materials Today* https://doi.org/10.1016/j.mattod.2020.08.016 (2020).
- Yang, Z., Tang, B., Xie, Z. & Zhou, Z. NASICON-Type Na3Zr2Si2PO12 solid-state electrolytes for sodium batteries. ChemElectroChem 8, 1035–1047 (2021).
- Deng, Z. et al. Fundamental investigations on the sodium-ion transport properties of mixed polyanion solid-state battery electrolytes. Nat. Commun. 13, 4470 (2022).
- Slater, P. R. & Greaves, C. Synthesis and conductivities of Sulfate/ Selenate phases related to Nasicon: NaxM'(II)xM"(III)2-x(SO4)3y(SeO4)y. J. Solid State Chem. 107, 12–18 (1993).
- 21. Cava, R., Vogel, E. M. & Johnson, D. W. Jr Effect of homovalent framework cation substitutions on the sodium ion conductivity in Na3Zr2Si2PO12. *J. Am. Ceram.* Soc. **65**, c157–c159 (1982).
- Singh, B. et al. A chemical map of NaSICON electrode materials for sodium-ion batteries. J. Mater. Chem. A 9, 281–292 (2021).
- 23. Guin, M. & Tietz, F. Survey of the transport properties of sodium superionic conductor materials for use in sodium batteries. *J. power sources* **273**, 1056–1064 (2015).
- Song, S., Duong, H. M., Korsunsky, A. M., Hu, N. & Lu, L. A Na+ superionic conductor for room-temperature sodium batteries. Sci. Rep. 6, 32330 (2016).
- 25. Lu, Y. et al. A high-performance monolithic solid-state sodium battery with Ca2+ doped Na3Zr2Si2PO12 electrolyte. *Adv. Energy Mater.* **9**, 1901205 (2019).
- Ma, Q. et al. Scandium-substituted Na3Zr2 (SiO4) 2 (PO4) prepared by a solution-assisted solid-state reaction method as sodium-ion conductors. Chem. Mater. 28, 4821–4828 (2016).
- Yue, Y. & Pang, W. Hydrothermal synthesis and characterization of NaSn2 (PO4) 3. J. Mater. Sci. Lett. 11, 148–149 (1992).
- 28. Slater, P. & Greaves, C. Synthesis and conductivities of sulfate/selenate phases related to nasicon: NaxM'(II) xM "(III) 2-x (SO4) 3-y (SeO4) y. J. Solid State Chem. 107, 12–18 (1993).
- Winand, J.-M., Rulmont, A. & Tarte, P. Nouvelles solutions solides LI (MIV) 2- x (NIV) x (PO4) 3 (L= Li, Na M, N= Ge, Sn, Ti, Zr, Hf) synthèse

- et étude par diffraction x et conductivité ionique. *J. Solid State Chem.* **93**. 341–349 (1991).
- 30. Zahir, M., Olazcuaga, R. & Hagenmuller, P. Crystal chemistry and ionic conductivity in Nasicon-type phases Na1+ xZr2− xYbx (AsO4) 3 with 0≤ x≤ 0.45. *Mater. Lett.* **2**, 234–236 (1984).
- 31. Fujitsu, S., Nagai, M., Kanazawa, T. & Yasui, I. Conduction paths in sintered ionic conductive material Na1+ xYxZr2- x (PO4) 3. *Mater. Res. Bull.* **16**, 1299–1309 (1981).
- 32. Cherkaoui, F., Viala, J., Delmas, C. & Hagenmuller, P. Crystal chemistry and ionic conductivity of a new Nasicon-related solid solution Na1+ xZr2- x2Mgx2 (PO4) 3. *Solid State Ion.* **21**, 333–337 (1986).
- Saito, Y., Ado, K., Asai, T., Kageyama, H. & Nakamura, O. Ionic conductivity of NASICON-type conductors Na1. 5MO. 5Zr1. 5 (PO4) 3 (M: Al3+, Ga3+, Cr3+, Sc3+, Fe3+, In3+, Yb3+, Y3+). Solid State Ion. 58, 327–331 (1992).
- Takahashi, T., Kuwabara, K. & Shibata, M. Solid-state ionics-conductivities of Na+ ion conductors based on NASICON. Solid State Ion. 1, 163–175 (1980).
- Kapoor, V., Singh, B., Sai Gautam, G., Cheetham, A. K. & Canepa, P. Rational design of mixed polyanion electrodes Na x V2P3-i (Si/S) i O12 for sodium batteries. *Chem. Mater.* 34, 3373–3382 (2022).
- 36. Ouyang, B. et al. Synthetic accessibility and stability rules of NASICONs. *Nat. Commun.* **12**, 1–11 (2021).
- 37. Deng, Z. et al. Phase behavior in rhombohedral NaSiCON electrolytes and electrodes. *Chem. Mater.* **32**, 7908–7920 (2020).
- Bianchini, M. et al. The interplay between thermodynamics and kinetics in the solid-state synthesis of layered oxides. *Nat. Mater.* 19, 1088–1095 (2020).
- Bai, J. et al. Kinetic pathways templated by low-temperature intermediates during solid-state synthesis of layered oxides. *Chem. Mater.* 32, 9906–9913 (2020).
- Miura, A. et al. Selective metathesis synthesis of MgCr2S4 by control of thermodynamic driving forces. *Mater. Horiz.* 7, 1310–1316 (2020).
- 41. Ruan, Y. et al. Optimization of Na3Zr2Si2PO12 ceramic electrolyte and interface for high performance solid-state sodium battery. *Ceram. Int.* **45**, 1770–1776 (2019).
- 42. Kononova, O. et al. Opportunities and challenges of text mining in materials research. *Iscience* **24**, 102155 (2021).
- He, T. et al. Similarity of precursors in solid-state synthesis as textmined from scientific literature. *Chem. Mater.* 32, 7861–7873 (2020).
- Weston, L. et al. Named entity recognition and normalization applied to large-scale information extraction from the materials science literature. J. Chem. Inf. Model. 59, 3692–3702 (2019).
- Boilot, J. P., Collin, G. & Colomban, P. Relation structure-fast ion conduction in the NASICON solid solution. J. Solid State Chem. 73, 160–171 (1988).
- Miyajima, Y., Saito, Y., Matsuoka, M. & Yamamoto, Y. Ionic conductivity of NASICON-type Na1+ xMxZr2- xP3O12 (M: Yb, Er, Dy). Solid State Ion. 84, 61–64 (1996).
- 47. Jolley, A. G., Cohn, G., Hitz, G. T. & Wachsman, E. D. Improving the ionic conductivity of NASICON through aliovalent cation substitution of Na 3 Zr 2 Si 2 PO 12. *Ionics* **21**, 3031–3038 (2015).
- Zhang, Z. et al. Correlated migration invokes higher Na+-Ion conductivity in NaSICON-type solid electrolytes. Adv. Energy Mater. 9, 1902373 (2019).
- Xiao, Y. et al. Lithium oxide superionic conductors inspired by garnet and NASICON structures. Adv. Energy Mater. 11, 2101437 (2021).
- Xiong, S. et al. Computation-guided design of LiTaSiO5, a new lithium ionic conductor with sphene structure. Adv. Energy Mater.
 9, 1803821 (2019).

- He, X. et al. Crystal structural framework of lithium super-ionic conductors. Adv. Energy Mater. 9, 1902078 (2019).
- 52. Kozinsky, B. et al. Effects of sublattice symmetry and frustration on ionic transport in garnet solid electrolytes. *Phys. Rev. Lett.* **116**, 055901 (2016).
- 53. Zhang, Y. et al. Unsupervised discovery of solid-state lithium ion conductors. *Nat. Commun.* **10**, 1–7 (2019).
- 54. Jun, K. et al. Lithium superionic conductors with corner-sharing frameworks. *Nat. Mater.* **21**, 924–931 (2022).
- Fang, H. & Jena, P. Argyrodite-type advanced lithium conductors and transport mechanisms beyond peddle-wheel effect. *Nat. Commun.* 13, 1–11 (2022).
- Zhang, Z. & Nazar, L. F. Exploiting the paddle-wheel mechanism for the design of fast ion conductors. Nat. Rev. Mater. 7, 389–405 (2022).
- 57. Zhang, Z. et al. Targeting superionic conductivity by turning on anion rotation at room temperature in fast ion conductors. *Matter* **2**, 1667–1684 (2020).
- Smith, J. G. & Siegel, D. J. Low-temperature paddlewheel effect in glassy solid electrolytes. Nat. Commun. 11, 1–11 (2020).
- Zhang, Z., Roy, P.-N., Li, H., Avdeev, M. & Nazar, L. F. Coupled cation–anion dynamics enhances cation mobility in roomtemperature superionic solid-state electrolytes. *J. Am. Chem. Soc.* 141, 19360–19372 (2019).
- Sun, Y. et al. Enhanced ionic conductivity and lack of paddle-wheel effect in pseudohalogen-substituted Li argyrodites. *Matter* 7, 4379–4395 (2022).
- Muy, S. et al. Tuning mobility and stability of lithium ion conductors based on lattice dynamics. Energy Environ. Sci. 11, 850–859 (2018).
- Krauskopf, T. et al. Comparing the descriptors for investigating the influence of lattice dynamics on ionic transport using the superionic conductor Na3PS_{4-x}Se_x. J. Am. Chem. Soc. 140, 14464–14473 (2018).
- 63. Muy, S., Schlem, R., Shao-Horn, Y. & Zeier, W. G. Phonon-ion interactions: designing ion mobility based on lattice dynamics. *Adv. Energy Mater.* **11**. 2002787 (2021).
- Rietveld, H. M. A profile refinement method for nuclear and magnetic structures. J. Appl. Crystallogr. 2, 65–71 (1969).
- Rodríguez-Carvajal, J. Recent advances in magnetic structure determination by neutron powder diffraction. *Phys. B: Condens. Matter* 192, 55–69 (1993).
- Kresse, G. & Furthmüller, J. Efficient iterative schemes for ab initio total-energy calculations using a plane-wave basis set. *Phys. Rev. B* 54, 11169–11186 (1996).
- Perdew, J. P., Burke, K. & Ernzerhof, M. Generalized gradient approximation made simple. *Phys. Rev. Lett.* 77, 3865–3868 (1996).
- Ouyang, B., Wang, Y., Sun, Y. & Ceder, G. Computational investigation of Halogen-substituted Na argyrodites as solid-state superionic conductors. *Chem. Mater.* 32, 1896–1903 (2020).
- Wang, J. et al. A high-energy NASICON-type cathode material for Na-Ion batteries. Adv. Energy Mater. 10, 1903968 (2020).
- Kim, H., Ishado, Y., Tian, Y. & Ceder, G. Investigation of Alkali-Ion (Li, Na, and K) Intercalation in KxVPO4F (x~ 0) Cathode. Adv. Funct. Mater. 29, 1902392 (2019).
- Seo, D.-H. et al. The structural and chemical origin of the oxygen redox activity in layered and cation-disordered Li-excess cathode materials. Nat. Chem. 8, 692 (2016).
- Dacek, S. T., Richards, W. D., Kitchaev, D. A. & Ceder, G. Structure and dynamics of fluorophosphate Na-ion battery cathodes. *Chem. Mater.* 28, 5450–5460 (2016).
- Morgan, D. et al. Experimental and computational study of the structure and electrochemical properties of Li x M2 (PO4) 3

- compounds with the monoclinic and rhombohedral structure. *Chem. Mater.* **14**, 4684–4693 (2002).
- Jain, A. et al. Commentary: The materials project: a materials genome approach to accelerating materials innovation. *Apl. Mater.* 1, 011002 (2013).
- 75. Nosé, S. A unified formulation of the constant temperature molecular dynamics methods. *J. Chem. Phys.* **81**, 511–519 (1984).
- Kononova, O. et al. Opportunities and challenges of text mining in materials research. *Iscience* 24, 102155 (2021).
- He, X., Zhu, Y., Epstein, A. & Mo, Y. Statistical variances of diffusional properties from ab initio molecular dynamics simulations. *npj* Comput. Mater. 4, 18 (2018).

Acknowledgements

This work was supported by the Samsung Advanced Institute of Technology. The computational analysis was performed using computational resources sponsored by the Department of Energy's Office of Energy Efficiency and Renewable Energy at the National Renewable Energy Laboratory. Computational resources were also provided by the Extreme Science and Engineering Discovery Environment (XSEDE), which is supported by the National Science Foundation grant number ACI1053575 and the National Energy Research Scientific Computing Center (NERSC), a DOE Office of Science User Facility supported by the Office of Science and the U.S. Department of Energy under contract no. DE-ACO2- 05CH11231. This research used beamline 7-BM of the National Synchrotron Light Source II, a U.S. Department of Energy (DOE) Office of Science User Facility operated for the DOE Office of Science by Brookhaven National Laboratory under Contract No. DE-SC0012704.

Author contributions

J.W., B.O., and G.C. initiated and designed the project. G.C. supervised all aspects of the research. B.O. performed the high-throughput DFT calculations and AIMD simulations. B.O. and J.W. analyzed the calculation data. J.W. synthesized the materials and performed characterizations and electrochemical tests with the help of Z.C. and X.Y. X. Y. acquired the synchrotron diffraction data. In addition, T. H. and J.W. performed the text mining and analyzed the data. Y.W., V.L., and H.K. contributed valuable discussion and insights. J.W., B.O., and G.C. wrote the manuscript, and the manuscript was revised by all the authors.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41467-023-40669-0.

Correspondence and requests for materials should be addressed to Bin Ouyang or Gerbrand Ceder.

Peer review information *Nature Communications* thanks Marco Amores and the other, anonymous, reviewers for their contribution to the peer review of this work. A peer review file is available.

Reprints and permissions information is available at http://www.nature.com/reprints

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit https://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2023