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Potassium-rich antiperovskites K_3HTe and K_3FTe and their structural relation to lithium and sodium counterparts†

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Unlike perovskite oxides, antiperovskites M_3HCh and M_3FCh (M=Li, Na; Ch=S, Se, Te) mostly retain their ideal cubic structure over a wide range of compositions owing to anionic size flexibility and low-energy phonon modes that promote their ionic conductivity. In this study, we show the synthesis of potassium-based antiperovskites K_3HTe and K_3FTe and discuss the structural features in comparison with lithium and sodium analogues. It is shown experimentally and theoretically that both compounds maintain a cubic symmetry and can be prepared at ambient pressure, in contrast to most of the reported M_3HCh and M_3FCh which require high pressure synthesis. A systematic comparison of a series of cubic M_3HTe and M_3FTe (M=Li, Na, K) revealed that telluride anions contract in the order of K, Na, Li, with a pronounced contraction in the Li system. This result can be understood in terms of the difference in charge density of alkali metal ions as well as the size flexibility of Ch anions, contributing to the stability of the cubic symmetry.

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Introduction

Antiperovskites, represented by the chemical formula X_3BA (A, B: anions, X: cations), exhibit diverse physical and chemical properties owing to a versatile array of chemical compositions (Fig. 1). For example, lithium- and sodium-rich antiperovskites such as $M_3O(Cl_{1-x}Br_x)$ (M = Li, Na) exhibit excellent lithium/sodium ion conductivity that could be used in solid electrolytes for high energy density batteries. Recently, we have synthesized antiperovskites M_3HCh (M = Li, Na; Ch = S, Se, Te) with H anions at the B-site using high pressure reactions, and observed that the size flexibility of the H⁻ anion

stabilizes the ideal cubic structure $(Pm-\bar{3}m)$ over a wide compositional space, with the exception of Na₃HS.¹⁰ These compounds show a high ionic conductivity, which can be attributed to the low energy phonon modes associated with the rotation of HM₆ octahedra, lowering the migration barrier for lithium and sodium cations.¹⁰

Subsequently, antiperovskites M_3FCh (M = Li, Na; Ch = S, Se, Te) were synthesized mainly using the high pressure technique. Despite the use of the hard F anion instead of the soft H anion, M_3FCh also adopts the cubic structure with the

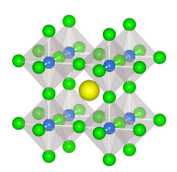


Fig. 1 Crystal structure of alkali metal rich antiperovskite M_3BTe (M=Li, Na, K; B=H, F). The newly synthesized K_3HTe and K_3FTe adopt the cubic symmetry ($Pm-\bar{3}m$). The B-site anion (H or F) (blue) occupies the octahedral center formed by K cations (green) and the A-site telluride anion (yellow) occupies the cuboctahedral site.

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[†] Electronic supplementary information (ESI) available: Laboratory X-ray diffraction patterns of K_3HTe and K_3FTe , size-flexibility of soft anions in other compounds, migration barriers in K_3HTe and K_3FTe , and measured ionic conduction of pristine and iodine substituted K_3HTe (PDF). See DOI: https://doi.org/10.1039/d3dt01039b

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exception of Na₃FS, and thus comparable ionic conduction is obtained due to the associated low energy FM6 octahedral rotation modes. These results, together with first-principles calculations, imply that highly polarizable Ch anions play a major role in stabilizing the cubic symmetry. Furthermore, we found a correlation between the soft phonon modes, the tolerance factor, 12 and the activation energy for the mobile cations.

This work aims to expand the available compositional space of M₃HCh and M₃FCh by including potassium instead of Li and Na. We have successfully synthesized K3HTe and K₃FTe under both high pressure and ambient pressure conditions. The preparation of a series of alkali metal-rich antiperovskites allows us to compare them systematically together with first-principles calculations. It is found that in M₃BCh (M = Li, Na, K; B = H, F; Ch = S, Se, Te), not only the size-flexibility of the Ch-site anions but also the charge density of the M-site cations has an impact on their structures.

Experimental

Polycrystalline samples of K₃HTe and K₃FTe were initially synthesized by a solid-state reaction under high pressure using washed KH (Sigma-Aldrich, 30 wt% dispersion mineral oil) with hexane and K₂Te (Kojundo, 99%), and KF (Wako, 99.0%) and K2Te as precursors, respectively. K3FTe was obtained with a stoichiometric ratio (i.e., KF: K_2 Te = 1:1), while the synthesis of K_3 HTe requires a hydrogen-rich ratio of $KH: K_2$ Te = 2:1, as is the case with other oxyhydrides synthesized under pressure including SrCrO₂H, ¹³ BaScO₂H, ¹⁴ BaVO_{3-x}H_x $(0.3 \le x \le 0.8)^{15}$ and $LaSrMnO_{3.3}H_{0.7}$. ¹⁶ The starting materials were thoroughly ground and pressed into a pellet, which was inserted into a boron nitride (BN) sleeve and sealed at both ends with BN caps. The assembly was then loaded into a graphite tube heater and sealed in a pyrophyllite cube, serving as a pressure medium. All the above procedures were conducted in a glovebox filled with N₂ (mBRAUN, Unilab-2000) due to the instability of the precursors and the products towards air and humidity. The cubic cells were pressed under 5 GPa, and then heated at 400 °C for 10 min for K3HTe and at 750 °C for 10 min for K₃FTe. After maintaining the temperature for 1 hour, the samples were cooled to room temperature in 5 min before the pressure was released. The recovered cells were placed back in the glovebox and the obtained polycrystalline samples were collected. We also attempted to synthesize both compounds under ambient pressure; pellets of the mixture of starting materials were vacuum sealed into Pyrex tubes and heated at 330 °C for 14 hours for K₃HTe and at 450 °C for 14 hours for K₃FTe. Iodine-substituted K₃HTe was synthesized using a mixture of KH, K2Te and KI with a hydrogen-rich ratio of 2:0.9:0.1 under the same conditions as those for pristine K₃HTe.

High-resolution synchrotron X-ray diffraction (SXRD) measurements were carried out at room temperature with a wavelength of $\lambda = 0.42$ Å at the BL02B2 beamline in SPring-8 (Japan), equipped with MYTHEN solid-state detectors. The K₃HTe and K₃FTe powder samples were sealed in borosilicate capillaries. Le Bail and Rietveld analyses of SXRD data were carried out using JANA2006 software, 17 while the crystal structures were visualized using VESTA.18

Ionic conductivities of cold-pressed pellets of K₃HTe and I-substituted K₃HTe were measured by electrochemical impedance spectroscopy (EIS) with a constant voltage of 10 mV in the frequency range from 1 MHz to 0.1 Hz using an ECS ModuLab potentiostat/galvanostat. Around 0.2 g of the powder specimen was placed between two stainless-steel rods as an ion-blocking electrode in a custom-made Swagelok cell and pressed into a 10 mm diameter pellet using a hydraulic press at 360 MPa (thickness: 1-1.5 mm). The temperature was varied from 25 °C to 100 °C. Data at each temperature were collected after being held for 115 min to ensure temperature stabilization.

Ab initio calculations on K3HTe, K3FTe, KH, KF and K2Te were performed under 0 and 5 GPa using the plane-wave basis projector augmented wave (PAW) method19 implemented in the Vienna *Ab initio* Simulation Package (VASP). 20,21 The generalized gradient approximation in the form of PBEsol²² was employed. The valence configurations in the PAW potentials used in this study are as follows: [3s² 3p⁶ 4s¹] for K, [1s¹] for H, [2s² 2p⁵] for F, and [5s² 5p⁴] for Te. The remaining electrons were treated as frozen core electrons. The plane-wave energy cutoff was set to 650 eV. The first Brillouin zone was sampled using Γ -centered k-point grids with a spacing of 0.2 \mathring{A}^{-1} . The total energy convergence was set to 1.0×10^{-6} eV per atom, and atomic positions and lattice constants were relaxed until the residual forces of atoms were less than 1.0×10^{-3} eV Å⁻¹.

Phonon band structures and vibrational free energies under harmonic approximation were calculated using the finite displacement method²³ implemented in Phonopy.²⁴ The displacement of atoms from their stationary positions to obtain second-order force constants was set to 0.01 Å. A 2 \times 2 \times 2 supercell with respect to the conventional unit cell was used to avoid the self-interaction between the displaced atoms. Nonanalytical term correction²⁵ was adopted using the Born effective charge tensors and dielectric constants calculated using density functional perturbation theory implemented in VASP. 20,21

The nudged elastic band (NEB) method²⁶ was used to calculate the energy barriers of K⁺ migration via vacancies and interstitial defects (dumbbell) in K₃HTe and K₃FTe. Charged supercells with the dimensions of $3 \times 3 \times 3$ were used. The first Brillouin zone was sampled with one k-point (0.25, 0.25, 0.25) due to the large size of supercells. For NEB calculations, the total energy and force convergence were set to 1.0×10^{-6} eV per cell and 2.0×10^{-2} eV Å⁻¹, respectively. Note that all the computational conditions employed here are consistent with our previous studies. 10,11

Results and discussion

As shown in Fig. S1a,† the XRD pattern of K₃HTe synthesized at 400 °C under 5 GPa is similar to that of the ideal antiperovs**Paper Dalton Transactions**

kite Na₃HTe $(Pm-\bar{3}m)$. Several additional reflections were assigned to unreacted KH, while the remaining small peaks could not be assigned to any known phases, nor to a superstructure as found in Na₃HS and Na₃FS with coherent octahedral rotations.^{10,11} The lattice parameter of K_3HTe , a =5.35927(2) Å, is longer than that of Na₃HTe (a = 4.76349(2) $m \AA),^{10}$ reflecting the difference in ionic radius between Na and K (Na⁺: 1.02 Å vs. K⁺: 1.38 Å).²⁷ Similarly, the XRD pattern of K₃FTe was indexed in the ideal cubic structure, along with KF and unknown impurities (Fig. S1b†). The lattice constant a =5.25451(1) Å is longer than that of the Na counterpart (a = 4.70315(1) Å).11

Rietveld refinement of the SXRD data for K3HTe assuming the cubic antiperovskite structure $(Pm-\bar{3}m)$ readily converged to GOF = 3.94, $R_p = 5.44\%$ and $R_{\rm wp} = 7.56\%$ (Fig. 2a), with reasonable isotropic atomic displacement parameter, U_{iso} values of 0.0385(2) Å² and 0.0239(1) Å² for K and Te, respectively. Refining occupancy factors, g, at K resulted in 0.988(1). Note that the U_{iso} of H and g of H and Te are not refined. Likewise, the Rietveld analysis of K₃FTe converged successfully, yielding GOF = 5.52, R_p = 5.33% and R_{wp} = 7.31%, U_{iso} of

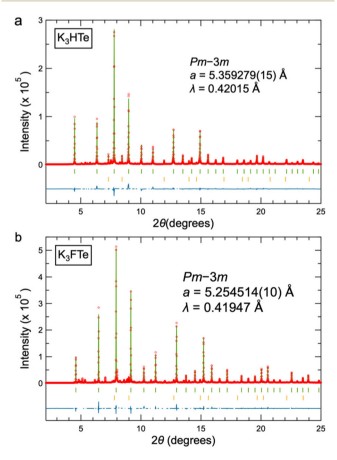


Fig. 2 Rietveld refinement of SXRD data of (a) K₃HTe and (b) K₃FTe. The red circles, the green solid line, and the blue solid line represent the observed, calculated, and difference intensities, respectively. The upper ticks indicate the calculated Bragg reflections for (a) K₃HTe and (b) K₃FTe, while the lower ticks for (a) KH and (b) KF. The unknown peaks are removed from the refinement.

 $0.0235(2) \text{ Å}^2 \text{ for K, } 0.0155(9) \text{ Å}^2 \text{ for F and } 0.0165(1) \text{ Å}^2 \text{ for Te}$ (Fig. 2b), and the occupancy of 0.990(1) for K and 0.960(5) for F. Thus, unless otherwise stated, we hereafter use the stoichiometric compositions of K₃HTe and K₃FTe.

Most of the alkali metal rich antiperovskites M3HCh and M₃FCh (M = Li, Na) reported so far have been synthesized at high pressure, 10,11 with the exception of Na₃HTe and Na₃FTe (unpublished data). First-principles calculations showed that the ideal cubic structure is dynamically stable at high pressure for most Na- and Li-based compounds, but at ambient pressure, many of them showed imaginary phonon modes appearing at the M and R points relevant to octahedral rotation. Here, we calculated the phonon band structures of K₃HTe and K₃FTe using the cubic structure at 0 and 5 GPa and found no imaginary phonon modes even under ambient pressure (Fig. 3a and b). The atom projected phonon density of states (DOS) for K3HTe indicates that the light H ion is responsible for high frequencies, whereas the heavy K and Te ions are for low frequencies. In K3FTe, this tendency is less pronounced. In addition, both compounds are lower in energy at ambient pressure than the starting materials K2Te and KH/ KF, suggesting that they are thermodynamically stable phases (Fig. 3c and d). These results led us to synthesize K₃HTe and K₃FTe under ambient pressure, and we indeed obtained the XRD patterns with identical lattice constants for both cases (Fig. S2 and S3†).

The discovery of K₃HTe and K₃FTe offers an opportunity to systematically compare the impact of the cations on stabiliz-

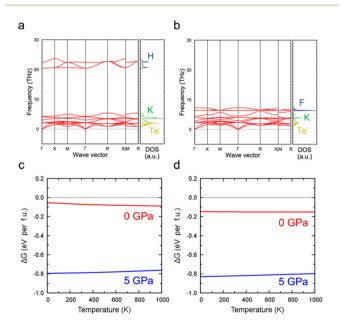


Fig. 3 Phonon band structures and partial phonon DOS of each ion of the cubic antiperovskites (a) K₃HTe and (b) K₃FTe at 0 GPa. The absence of imaginary phonon frequencies at the M and R points related to octahedral tilting, as found in, e.g., $\mathrm{Na_3HS}$ and $\mathrm{Na_3FS}$, $^{\mathrm{10,11}}$ means that both compounds are dynamically stable. The Gibbs free energy ΔG (ΔG = $\Delta U_{\text{internal}} + P\Delta V - \Delta F_{\text{phonon}}$, $\Delta F_{\text{phonon}} = \Delta E_{\text{ZPE}} - T\Delta S_{\text{phonon}}$) of (c) K₃HTe relative to that of KH + K₂Te and (d) K₃FTe relative to that of KF + K₂Te at 0 GPa, all calculated at ambient pressure.

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ation in the two sets of cubic antiperovskites, M3HTe and M_3 FTe (M = Li, Na, K). Fig. 4a shows the lattice parameters of cubic M3HTe and M3FTe as a function of Shannon ionic radii of the alkali metal cations, $r_{\rm M}$. In both systems, the lattice expands linearly with increasing $r_{\rm M}$, but the slope is obviously different, with a larger value for M2HTe, which suggests that this phenomenon is related to the size flexibility of hydride anions (vs. F⁻). 10,28 To obtain further insights into the structural features, the obtained lattice constants were used to estimate the ionic radius of H⁻ anions, $r_{\rm H}$, where $r_{\rm H} = a/2 - r_{\rm M}$ ($r_{\rm Li}$ = 0.76 Å, r_{Na} = 1.02 Å, r_{K} = 1.38 Å), as employed previously (Fig. 4b). 10 The expanded H anion in Na₃HTe compared to that in K₃HTe looks reasonable because the smaller ionic radius of Na⁺ (vs. K⁺) is compensated by the larger H⁻ anion. Following this trend, the H⁻ radius of Li₃HTe is expected to be larger than that of Na₃HTe. However, the opposite result, in which H contracts, was obtained, which implies that the hydride anion is not solely responsible for stabilizing and maintaining the cubic structure.

We hence took a theoretical approach to address this issue. We estimated the Bader volume²⁹ for each ion using the relaxed structures at ambient pressure by first-principles calculations. In terms of the H volume, its M dependence with a contraction at Li₃HTe appears to be reproduced, as shown in Fig. 5. However, similar dependence is also found in the F volume both qualitatively and quantitatively, with a smaller F volume in Li₃FTe than in Na₃FTe. Instead, a more pronounced M-dependence is seen in the tellurium anion, where the Te anion decreases its size in the order of K, Na, and Li.

The large variation in the size of the Te anion as a function of the alkali metal cations or unexpected contractions of the B-site anions in Li systems is likely related to the surface charge density of the alkali metal which, when increasing from K to Na and Li, leads to stiffer anions. We have noticed similar behaviour in lattice constants or anionic sizes in binary compounds, such as rock-salt MH30-35 and antifluorite M₂Te³⁶⁻³⁹ (Fig. S4†). Thus, the charge density of the M-site cation seems to predominantly modify the size of the Te anion to compensate for the size mismatch and preserve the cubic

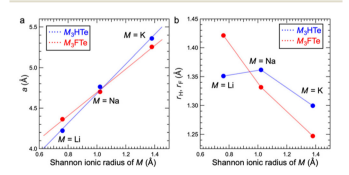


Fig. 4 (a) The lattice parameters of (blue) M_3HTe and (red) M_3FTe (M = Li, Na, K). 10,11 (b) The ionic radius of H $^-$ and F $^-$, $r_{\rm H}$ and $r_{\rm F}$, determined from the lattice constant, where $r_{\rm H} = (a - 2r_{\rm M})/2$ and $r_{\rm F} = (a - 2r_{\rm M})/2$. The fixed ionic radii of 0.76 Å, 1.02 Å and 1.38 Å were used for Li, Na and K, respectively.²⁷ Error bars are within the circles in both figures.

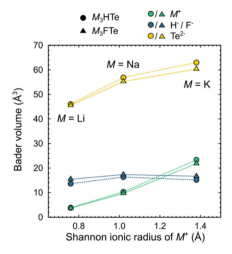


Fig. 5 The Bader volume of the alkali metal cation (green), H⁻/F⁻ anion (blue), and Te^{2-} anion (yellow) in M_3HTe (circles) and M_3FTe (triangles).

symmetry. Bearing in mind that Te has a high compressibility, the exceptionally small Te anion in Li₃HTe is consistent with the fact that it can only be synthesized under high pressure.

Since M_3HCh and M_3FCh (M = Li, Na; Ch = S, Se, Te) exhibit excellent lithium or sodium ion conductivity, 10,11 we performed nudged elastic band (NEB) calculations to estimate the migration energy barriers of K+ transport in K3HTe and K₂FTe. As seen in Fig. 6, their energy barriers are comparable to those of Li-rich and Na-rich antiperovskites, as low as 265 meV and 352 meV, respectively, for the vacancy mechanism, and 76 meV and 102 meV for the interstitial dumbbell mechanism. The low migration energy barriers in K-rich antiperovskites indicate their potential as ionic conductors, once vacancies or interstitial defects are introduced by the aliovalent substitution approach, for example. K3HTe has a lower energy barrier than K₃FTe, which is partly due to the softer phonon

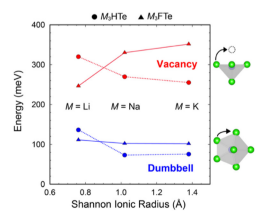


Fig. 6 Migration energy barriers of (circles) M₃HTe and (triangles) M_3FTe (where M = Li, Na, K). The results of vacancy and interstitial dumbbell mechanisms are shown in red and blue, respectively. The schematics on the right side show the migration pathways of M⁺ ions around the HM₆/FM₆ octahedron. The migration energy barriers of Liand Na-rich antiperovskites are taken from ref. 10 and 11.

mode relevant to HK₆ octahedral rotation (Fig. S5†). Attempts

to measure the K⁺ ion conductivity of pristine and I-doped K_3 HTe resulted in high resistances (at least >10⁶ Ω), which precluded proper estimation of bulk conductivities and activation energies (Fig. S6†). The large resistances may be attributed to the extremely high sensitivity of the sample surface to trace amounts of oxygen and water; thus, there is considerable room to adjust synthetic conditions in the future.

Conclusions

Paper

We prepared two potassium-rich cubic K3HTe and K3FTe under ambient conditions, unlike the reported sodium- and lithium-rich antiperovskites M₃HCh and M₃FCh (M = Li, Na; Ch = S, Se, Te) stabilized under high pressure conditions. The systematic comparison of the two sets of M3HTe and M3FTe compounds, using first-principles calculations, revealed the unexpected contraction of B-site anions in Li systems and alkali metal-dependent size variation in Te anions. In perovskite oxides, the crystal structure is mainly controlled by octahedral rotations in order to compensate for the size mismatch between A, B, and O ions; thus, the cubic phase is stable only in a narrow compositional space, which could be a problem for some applications such as ionic conductivity. In contrast, the cubic structure is stable over a wide compositional space in M₃HCh and M₃FCh. Depending on the surface charge density of the M-site cations, the Ch-site anions flexibly change their size to compensate for the size mismatch between the BM₆ octahedra and the Ch-site anion to maintain the cubic symmetry. The alkali metal-dependent size effects of other ions obtained in this study provide a new dimension in structural control, and thus control of physical and chemical properties, not limited to antiperovskites.

Author contributions

K. O. and S. F. contributed equally to this work. K. O. carried out all the synthetic work and was helped by S. G. K. O., H. U. and C. T. contributed to the refinement of XRD data. S. F. and A. K. performed theoretical calculations. P. W., K. Y. and Y. U. measured the ionic conductivities. H. K. designed and coordinated this study, and contributed to all measurements and analyses.

Conflicts of interest

There are no conflicts to declare.

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