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Discovery of multi-anion antiperovskites X_6NFSn_2 (X = Ca, Sr) as promising thermoelectric materials by computational screening

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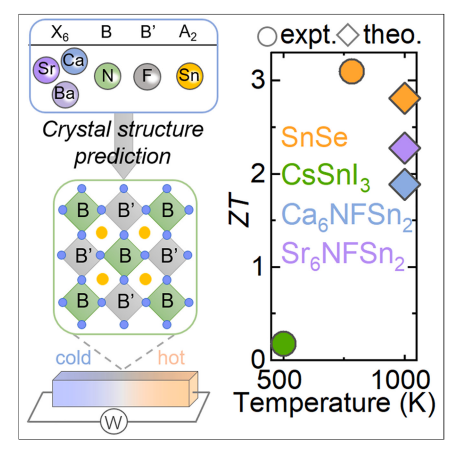
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Matter

Article

Discovery of multi-anion antiperovskites X_6 NFSn₂ (X = Ca, Sr) as promising thermoelectric materials by computational screening



We design new multinary antiperovskites X_6NFSn_2 (X = Ca, Sr, Ba). By exploring the phase stability, electrical, and thermal properties, Ca_6NFSn_2 and Sr_6NFSn_2 are identified as promising thermoelectric materials.



Discovery A new material or phenomena Han et al., Matter 7, 158–174 January 3, 2024 © 2024 The Authors. Published by Elsevier Inc. https://doi.org/10.1016/j.matt.2023.10.022



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Highlights

Designing phase-stable multianion antiperovskites X_6 NFSn₂ (X =Ca, Sr, and Ba)

Identifying the promising thermoelectric properties of multi-anion antiperovskites

Extracting lattice thermal conductivity assisted by machine learning

Unraveling the origin of low lattice thermal conductivity of X_6 NFSn₂ (X = Ca, Sr)



Article

Matter

Discovery of multi-anion antiperovskites X_6 NFSn₂ (X = Ca, Sr) as promising thermoelectric materials by computational screening

Dan Han,^{1,2,3,5,6,*} Bonan Zhu,^{2,5} Zenghua Cai,⁴ Kieran B. Spooner,^{2,3} Stefan S. Rudel,¹ Wolfgang Schnick,¹ Thomas Bein,¹ David O. Scanlon,^{2,3,*} and Hubert Ebert¹

SUMMARY

The thermoelectric performance of existing perovskites lags far behind that of state-of-the-art thermoelectric materials such as SnSe. Despite halide perovskites showing promising thermoelectric properties, namely, high Seebeck coefficients and ultralow thermal conductivities, their thermoelectric performance is significantly restricted by low electrical conductivities. Here, we explore new multi-anion antiperovskites X_6NFSn_2 (X = Ca, Sr, and Ba) via B-site anion mutation in antiperovskite and global structure searches and demonstrate their phase stability by first-principles calculations. Ca₆NFSn₂ and Sr₆NFSn₂ exhibit decent Seebeck coefficients and ultralow lattice thermal conductivities (<1 W $m^{-1} K^{-1}$). Notably, Ca₆NFSn₂ and Sr₆NFSn₂ show remarkably larger electrical conductivities compared to the halide perovskite CsSnI₃. The combined superior electrical and thermal properties of Ca₆NFSn₂ and Sr₆NFSn₂ lead to high thermoelectric figures of merit (ZTs) of \sim 1.9 and \sim 2.3 at high temperatures. Our exploration of multi-anion antiperovskites X_6 NFSn₂ (X = Ca, Sr) realizes the "phonon-glass, electron-crystal" concept within the antiperovskite structure.

INTRODUCTION

Thermoelectric (TE) materials allow for a direct energy conversion between heat and electricity, and they have potential applications in heat pumping and solid-state power generators.^{1–3} The conversion efficiency of TE materials can be assessed by the dimensionless figure of merit (ZT) $ZT = \frac{S^2 \sigma}{\kappa_1 + \kappa_e} T$. A combination of a large Seebeck coefficient (S), a high electrical conductivity (σ), and a low thermal conductivity (κ ; sum of electronic conductivity [κ_e] and lattice thermal conductivity [LTC] [κ_l]) is highly desired for high-performance TE materials.⁴ However, the complex interrelationship between electrical and thermal properties makes it challenging to achieve an ultrahigh ZT.⁵ For decoupling electrical and thermal properties, the concept of "phononglass, electron-crystal" (PGEC), viz. designing TE materials in structures allowing for efficient transport of charge carriers, as in a good semiconductor crystal, and largely blocking thermal transmission by the lattice, as in a glass, has been proposed.⁶ Indeed, the PGEC concept redirects the materials design strategy to intrinsic features of structure and bonding characteristics, and PGEC behavior was originally observed in cage-like compounds such as skutterudites and clathrates, where heavy atoms trapped in the cages "rattle" and thus strongly scatter phonons.⁷⁻⁹ Besides skutterudites and clathrates, the perovskite structure provides a platform to validate the PGEC concept as the heavy-element or molecular group sits at the A site of the AMX₃ perovskite structure and forms a large cuboctahedral cage with 12 atoms at

PROGRESS AND POTENTIAL

The concept of "phonon-glass, electron-crystal" (PGEC) has been proposed to decouple electrical and thermal properties and thus to enhance the figure of merit (*ZT*) of thermoelectric (TE) materials. PGEC behavior was originally observed in cage-like compounds such as skutterudites and clathrates. In addition to skutterudites and clathrates, the perovskite structure provides a platform to validate the PGEC concept. Nevertheless, the thermoelectric performance of existing perovskites lags far behind that of the state-of-the-art thermoelectric materials such as SnSe. Here, we design multi-anion antiperovskites X_6 NFSn₂ (X = Ca, Sr, and Ba). Ca₆NFSn₂ and Sr₆NFSn₂ exhibit superior electrical and thermal properties, leading to high ZTs of \sim 1.9 and \sim 2.3 at 1,000 K, which are higher than the reported perovskite oxides and halides. This design realizes the PGEC concept within the vast family of perovskiterelated compounds and provides a new strategy for promising TE materials.



the X site. Additionally, the A and X interactions are normally ionic and weak, with large vibrational amplitudes for the cations, which provides a chance for A-site element rattling at their equilibrium positions.¹⁰

Oxide perovskites have a long history of being studied as TE materials because of their high-temperature air stability, environmental friendliness, and straightforward synthesis.¹¹ However, the main limitation of these perovskites is their high κ (>10 W m^{-1} K⁻¹).^{11,12} State-of-the-art bulk TE materials like SnSe and PbTe exhibit κ s of less than 3 W m^{-1 K-1} at the same temperature (300 K).¹³ The well-studied SrTiO₃ perovskite with dopants La and Nb has shown promise in recent years and achieves a maximum ZT of 0.6 at over 1,000 K.¹⁴ Nonetheless, it is much inferior to bulk TE materials such as SnSe, PbTe, and BiCuSeO and typical half-Heusler alloys with demonstrated ZTs >1 at high temperature in a laboratory setting.^{15–18} Halide perovskites have drawn great attention with a view on application in photovoltaics, lightemitting diodes, and photodetectors due to their extraordinary optoelectronic properties.^{19–23} Generally, metal halides are not seriously considered as TE materials. However, a large S combined with an ultralow κ have been claimed for both hybrid and all-inorganic halide perovskites,²⁴⁻²⁶ which spurs the investigation of TE properties of halide perovskites.^{27,28} Despite the inherent high S and the ultralow κ , it is reported that intended doping for increasing conductivity in CH₃NH₃PbX₃ (X = Cl, Br, I) and CsPb X_3 (X = Cl, Br, I) remains challenging, which is attributed to the defect-tolerant electronic structure and ionic compensation of charged point defects.^{29–31} In contrast to the lead analogs, $CsSnX_3$ (X = Cl, Br, I) perovskites exhibit the possibility of using self-doping processes of Sn²⁺ to Sn⁴⁺ to achieve p-type electrical conductivities.^{32,33} Nevertheless, the σ is still low and seriously limits its TE performance; that is, current maximum ZT values of 0.08 and 0.175 are attained in CsSnl₃ at room temperature (RT) and 500 K, respectively.^{34,35}

In addition to perovskites, nitride and oxide antiperovskites have been investigated as TE materials.^{36–40} Antiperovskites maintain the perovskite structure, while their ionic types are inverted.^{41–45} Both nitride and oxide antiperovskites can show enhanced stability compared to halide perovskites. Similar to the cases of traditional binary nitrides,^{46–48} nitride antiperovskites exhibit high κ s, which are detrimental to the TE performance; for example, the κ s of Sr₃AsN are larger than 10 W m^{-1 K-1} at RT.³⁶ Compared to nitride antiperovskites, oxide antiperovskites have lower κ s, while they suffer from a low *S* due to narrower band gaps.^{37,38} Some of them (Ba₃OGe, X₃OSn, and X₃OPb [X = Ca, Sr, Ba]) even show band-gap closing, which is unfavorable for high-performance TE materials due to the cancellation of the electron and hole carrier transport and the loss of the n- or p-type character.³⁷ Thus far, many issues remain to be solved for realizing the full potential of existing perovskite-related materials for TE applications.

In light of the complex crystal structure with heavy elements favoring low LTC, here, we design multi-anion antiperovskites (double-antiperovskites) X_6 NFSn₂ (X = Ca, Sr, and Ba), which can be regarded as the B-site anion mutation in antiperovskite X_3 OSn (X = Ca, Sr, and Ba). Crystal structure predictions for X_6 NFSn₂ (X = Ca, Sr, and Ba) are performed using *ab initio* random structure searching (AIRSS) and particle swarm optimization.^{49–52} All of the low-energy structures of X_6 NFSn₂ (X = Ca, Sr, and Ba) appear in antiperovskite structures. Based on first-principles calculations, we demonstrate the phase stabilities, including dynamic stability and thermodynamic stability, and investigate the electronic structure of Ca₆NFSn₂ (space group [SG] $Fm\overline{3}m$), Sr₆NFSn₂ (SG R $\overline{3}$), and Ba₆NFSn₂ (SG R $\overline{3}$). The electrical and TE properties of Ca₆NFSn₂ and Sr₆NFSn₂ are further studied because of their larger band gaps

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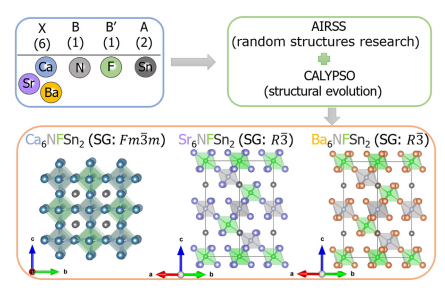


Figure 1. A schematic diagram of the crystal structure prediction of X₆**NFSn**₂ (**X** = **Ca**, **Sr**, and **Ba**) Conventional cells of compounds are displayed. Corresponding space group (SG) are given in parentheses. Color codes are given in the formula of the compounds.

(>0.2 eV), which are essential for maintaining a single carrier type and therefore a high ZT at high temperature. Ca_6NFSn_2 and Sr_6NFSn_2 both display low LTCs and remarkably enhanced electrical conductivities compared to halide perovskites, thus resulting in the predicted high maximum ZTs of p-type Ca_6NFSn_2 (1.89) and Sr_6NFSn_2 (2.28) at 1,000 K. Notably, the extraction of high-order force constants (FCs) for computing the LTC is achieved by advanced machine learning algorithms and first-principles calculations, which avoids the extremely high cost for extracting high-order FCs via substantial computations for the compounds with lower symmetry and large unit cells.

RESULTS AND DISCUSSION

Crystal structure prediction

We performed global structure searches for X_6 NFSn₂ (X = Ca, Sr, and Ba) to identify whether their low-energy structures can crystallize in the double-antiperovskite structure via AIRSS and crystal structure analysis by particle swarm optimization (CALYPSO). Using AIRSS, for X_6NFSn_2 (X = Ca, Sr, and Ba), 2,114, 2,078, and 2,075 random structures were generated, respectively, which were subsequently relaxed by CASTEP. As shown in Table S1, a low-energy structure with the symmetry of Fm3m of Ca_6NFSn_2 is found, which is identified by CALYPSO searching as well. Sr_6NFSn_2 and Ba_6NFSn_2 with the symmetry of $R\overline{3}$ exhibit the lowest energies. Explicit crystal parameters of these compounds are summarized in Table S2. Ca_6NFSn_2 (SG Fm3m), Sr_6NFSn_2 (SG R3), and Ba_6NFSn_2 (SG R3) form a double-antiperovskite structure with X-site cations (X = Ca, Sr, and Ba) at the corners of the octahedra, B/B'-site anion (N/F) at the center of octahedra, and Sn located at the A site, as displayed in Figure 1. $[X^{2+}_{6}N^{3-}]$ and $[X^{2+}_{6}F^{-}]$ (X = Ca, Sr, and Ba) octahedra alternate along the three crystallographic axes and form the rock-salt-type ordering. Heavy-atom Sn at the A site forms cuboctahedral cages with 12 Ca, Sr, and Ba atoms at the X site, and the distances between them are larger than 3.48, 3.55, and 3.63 Å, respectively, providing a chance for Sn atoms to act as cage rattlers. As the X-site element becomes heavier from Ca to Sr or Ba, the crystal structure evolves from cubic to trigonal symmetry while maintaining the corner-sharing octahedra. In contrast to high-symmetry Ca₆NFSn₂ without distortion, Sr₆NFSn₂ and Ba₆NFSn₂ show





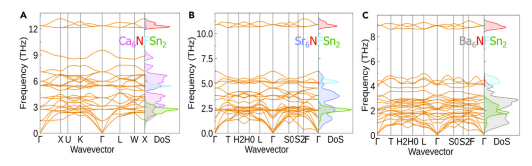


Figure 2. Phonon spectra together with atom-projected density of states (DoS) curves of X_6NFSn_2 (A–C) Phonon spectra together with atom-projected density of states (DoS) curves (A–C) of X_6NFSn_2 (X = Ca, Sr, Ba).

 $a^{-}a^{-}a^{-}$ (Glazer's notation) octahedral tilting, which is consistent with observations in the oxide perovskites, i.e., when the X-site radius increases, the tilting of octahedra can appear.^{53,54} The phase stabilities of X_6 NFSn₂ (X =Ca, Sr, and Ba) are evaluated in the next section.

Phase stability and electronic structure

The mechanical stability of X_6NFSn_2 (X = Ca, Sr, and Ba) is firstly evaluated. As summarized in Table S3, all compounds fulfill the Born-Huang stability criteria⁵⁵ and are mechanically stable. The thermodynamic stability of Ca₆NFSn₂, Sr₆NFSn₂, and Ba6NFSn2 was evaluated via convex hull analysis. All three compounds present on the hull, suggesting their thermodynamic stability. Via convex hull constructions, the predominant decomposition pathway of X_6 NFSn₂ into corresponding binary and ternary materials is identified and is given in the supplemental information. Moreover, a stable polyhedron of Ca₆NFSn₂ in the phase diagram is identified (see Figure S1), and the chemical potential region limits are given in Table S4, which demonstrate its thermodynamic stability. All the reported competing phases from the Materials Project, OQMD, and ALFOW databases were considered for the phase diagram of Ca_6NFSn_2 .^{56–58} It is displayed as a representation because the phase diagram of multinary compounds calculated by hybrid functional is time consuming to create. With regard to experimental results, it is noteworthy that several alkaline-earth metal nitrohalide Ca_2NI and Ba_2NX (X = F, Cl, Br) samples were successfully prepared under high temperature (~1,000 K).⁵⁹⁻⁶¹ The precursors of antiperovskite Ca₃OSn samples were heated at 1,123 K for several hours and then quenched.³⁸ Analogous reactions might enable the synthesis of the herein suggested compounds.

After validating the thermodynamic stability, their dynamic stabilities were assessed via harmonic phonon calculations. Three compounds show dynamic stability, as indicated by the absence of imaginary phonon modes (Figures 2A–2C). Thus far, the phase stability of X_6NFSn_2 (X = Ca, Sr, Ba) has been demonstrated. According to the inverse relationship between the reduced mass and frequency, the high-frequency phonon modes in the three compounds are dominated by the motions of N, the mid-frequency ones mostly comprise the motions of alkaline-earth metal (Ca/Sr/Ba) and F atoms, and the low-frequency ones are contributed by the motions of Sn and alkaline-earth metal (Ca/Sr/Ba) atoms, as shown in Figures 2A–2C. Ba₆NFSn₂ displays the narrowest frequency range among the three compounds. It can be ascribed to the heavy X atoms downshifting the frequencies. The highest acoustic mode boundary frequencies of Ca₆NFSn₂, Sr₆NFSn₂, and Ba₆NFSn₂ are 3, 2.5, and 2 THz, respectively, which are comparable to that of one of the current state-of-the-art TE materials, SnSe, and suggest a strong mode softening in these compounds.⁶² Notably, optical phonon modes appear in low-frequency regions





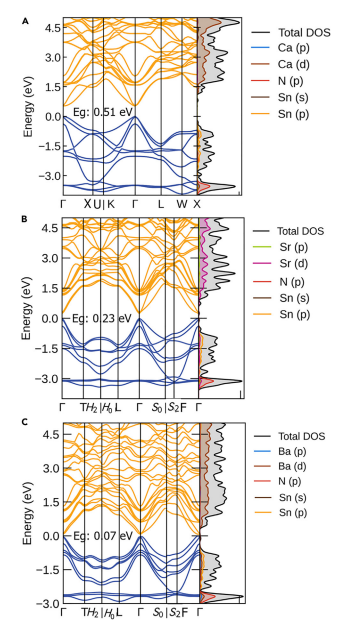


Figure 3. Band structures and projected DoS of X₆NFSn₂ calculated at the HSE06+SOC level (A–C) Band structures and projected DoS of (A) Ca₆NFSn₂, (B) Sr₆NFSn₂, and (C) Ba₆NFSn₂ calculated at the HSE06+SOC level.

(<3 THz). Taking Ca₆NFSn₂ as an example, optical phonon with a frequency of 1.42 THz cm⁻¹ at the Γ point corresponds to the [Ca₆F] and [Ca₆N] octahedra rocking motion, and the one with a frequency of 2.75 THz cm⁻¹ relates to the rocking of Sn atoms. These low-frequency optical phonon modes couple with acoustic phonons. Such a coupling would be expected to increase the scattering channels for the heat-carrying acoustic phonons, which indicates potentially low LTC. Analyses of the phonon scattering rate between acoustic and optical modes are shown in part below. Similar vibrational modes are observed in Sr₆NFSn₂ and Ba₆NFSn₂ as well.

Band structure calculations (Figure 3) show that Ca_6NFSn_2 and Sr_6NFSn_2 possess direct band gaps of 0.51 and 0.23 eV at the Γ point. Unlike semiconducting





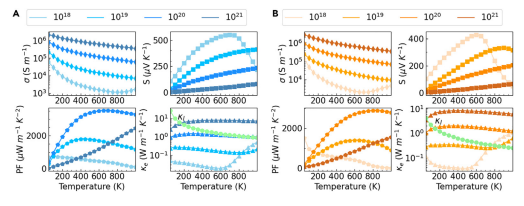


Figure 4. Calculated electrical transport properties as a function of temperature for p-type X₆NFSn₂ at four different carrier concentrations (A and B) Calculated electrical transport properties as a function of temperature for p-type (A) Ca₆NFSn₂ and (B) Sr₆NFSn₂ at four different carrier concentrations (1.0 × 10¹⁸–1.0 × 10²¹ cm⁻³): electrical conductivity (σ), Seebeck coefficient (S), power factor (*PF*), and electronic thermal conductivity (κ_e). The LTC (κ_i) is displayed on the κ_e plots with green symbols for comparison.

Ca₆NFSn₂ and Sr₆NFSn₂, the band gap of Ba₆NFSn₂ almost closes, which is generally not favorable for the TE performance. The change of element $X = Ca \rightarrow Sr \rightarrow Ba$ leads to the reduction of the band gaps due to the downshift of the conduction band edge mainly comprising X-atomic orbitals. One striking feature is that they all have 4- and 2-fold orbital degeneracies of the valence band edge (VBE) and the conduction band edge (CBE), respectively. The high orbital degeneracy of the VBE implies a potentially large *S*. Moreover, the valence band maximum (VBM) and the conduction band minimum (CBM) predominantly consist of Sn-p and nominally unoccupied X-d orbitals (see Figure 3), which suggests a cross-gap hybridization and covalent character in these compounds.⁶³ Owing to the cross-gap hybridization, these mixed-anion antiperovskites exhibit large valence and conduction band dispersion, indicative of small effective masses of the carriers.

Electrical transport and TE properties

The electrical transport properties of Ca_6NFSn_2 and Sr_6NFSn_2 were obtained via the momentum relaxation time approximation (MRTA) as implemented in AMSET under p-type- and n-type-doping conditions with carrier concentrations ranging from 10^{18} to 10^{21} cm⁻³. Compared to an accurate but highly costly method like DFPT+ Wannier, ^{64,65} AMSET has been demonstrated to produce excellent agreement for carrier mobilities and Ss in a series of semiconductors such as Si, GaN, SnSe, Bi₂Te₃, and CH₃NH₃Pbl₃ with experimental measurements at comparatively lower computational costs.⁶⁶ The TE performance of p-type Ca₆NFSn₂ and Sr₆NFSn₂ is discussed in detail here because these p-type compounds show higher ZTs in comparison to n-type. The corresponding data of n-type Ca₆NFSn₂ and Sr₆NFSn₂ are given in the supplemental information (Figures S2 and S3) for comparison.

Strikingly, Ca₆NFSn₂ and Sr₆NFSn₂ show good electrical transport properties. As shown in Figure 4A, the electrical conductivities of Ca₆NFSn₂ are in the range of 1.12×10^3 (at a carrier concentration of 10^{18} cm⁻³ and a temperature of 800 K) to 2.10×10^6 S m⁻¹ (at a carrier concentration of 10^{21} cm⁻³ and a temperature of 50 K). At the same carrier concentrations, Sr₆NFSn₂ shows higher electrical conductivities ranging from 1.99×10^3 (700 K) to 2.64×10^6 S m⁻¹ (50 K) (Figure 4B), and CsSnI₃ displays lower electrical conductivities from 1.03×10^2 to 5.10×10^5 S m⁻¹ (Figure S4). When calculating the hole mobilities, three scattering processes, namely the acoustic deformation potential (ADP), ionized impurity (IMP), and polar-optical phonon (POP) scattering, were considered. As shown in Figure S5, the





POP scattering process dominantly limits the hole mobility in Ca₆NFSn₂ and Sr₆NFSn₂ at RT and higher temperatures. Notably, the p-type electrical conductivities of Ca₆NFSn₂ and Sr₆NFSn₂ are at least an order of magnitude higher than those of CsSnI₃ at fixed carrier concentrations of 10^{18} and 10^{19} cm⁻³. As the carrier concentrations are further increased to 10^{20} and 10^{21} cm⁻³, the electrical conductivities of the two mixed-anion antiperovskites increase by 4-fold compared to those of CsSnI₃ at the same carrier concentrations. It has been experimentally pointed out that the low σ of CsSnI₃ restricts the TE performance despite its extremely low LTC; correspondingly, the obtained maximum ZT values of CsSnI₃ are 0.08 at RT and 0.175 at 550 K.^{10,34,35} Heavy hole (electron) doping to improve the electrical conductivities is thus crucial to boost the potential of CsSnI₃ as a promising TE material. We note that the calculated electrical conductivities of Ca₆NFSn₂ and Sr₆NFSn₂ are on a par with that of the current flagship chalcogenide TE material SnSe at the doping level of 10^{19} cm⁻³ using similar levels of theory.⁶⁷ We focus our discussion on this order of magnitude because the optimum ZTs of single-crystal and polycrystalline SnSe are obtained at this level and, on the other hand, the ZT maxima for Ca₆NFSn₂ and Sr_6NFSn_2 appear for carrier concentrations in the region of 10^{19} cm⁻³.^{15,68} The dopability of Ca₆NFSn₂ will be evaluated in future work. However, as we noticed, the dominant native defects (vacancies) in ternary nitride antiperovskites show low formation energies and efficient p-type or n-type self-doping.^{69,70}

In addition to high electrical conductivities, Ca₆NFSn₂ and Sr₆NFSn₂ have decent Ss. Their Ss are on a par with that of SnSe and 1.6-fold higher than that of CsSnI₃ at the doping level of 10^{19} cm⁻³. At the carrier concentration of 10^{18} cm⁻³, Ss of Ca₆NFSn₂ (Sr₆NFSn₂) increase smoothly to a peak at around 700 K (600 K) and then start to decline, as seen in Figure 4. The observed saturation of Ss at 10^{18} cm⁻³ at high temperatures can be attributed to the bipolar effect, which occurs in small-band-gap materials, especially at high temperature.⁷¹⁻⁷³ When the carrier concentration is elevated to 10^{19} cm⁻³, the saturation of Ss is relieved, to some extent, and the peaks of Ss shift to higher temperatures, namely 1,000 and 950 K for Ca₆NFSn₂ and Sr_6NFSn_2 , respectively. At even higher hole-doping levels (10^{20} and 10^{21} cm⁻³), the Ss increase monotonically as the temperature rises. The combination of high electrical conductivities and decent Ss of Ca₆NFSn₂ and Sr₆NFSn₂ leads to their p-type power factors (PFs) spanning the ranges of 40–3,470 and 5–2,886 μ W m⁻¹ K^{-2} , respectively. For Ca₆NFSn₂ (Sr₆NFSn₂), the PF computed at a fixed carrier concentration of 10²¹ cm⁻³ increases as the temperature rises, while those PFs calculated at the other three carrier concentrations (from 10^{18} to 10^{20} cm⁻³) peak at 100 (50 K), 400 (600 K), and 600 K (900 K), as shown in Figure 4.

The κ is composed of κ_e and κ_l . κ_l is often the limiting factor for the TE performance of promising candidates.⁴ The calculated κ_l of Ca₆NFSn₂ at RT is 2.94 W m⁻¹ K⁻¹, and it further decreases to 0.88 W m⁻¹ K⁻¹ at 1,000 K (Figure 4). Sr₆NFSn₂ show lower calculated κ_l s, viz. 0.83 and 0.25 W m⁻¹ K⁻¹ at RT and 1,000 K, respectively. The RT κ_l s of Ca₆NFSn₂ and Sr₆NFSn₂ are comparable to the calculated κ_l s of PbS (2.19 W m⁻¹ K⁻¹) and SnSe (0.52 W m⁻¹ K⁻¹).^{62,67,74} Notably, the ultralow LTC ($\kappa_l < 1 \text{ W m}^{-1} \text{ K}^{-1}$) of Ca₆NFSn₂ and Sr₆NFSn₂ at high temperatures is a feature of high-efficiency TEs. More discussion about the origin of the low κ_l is given in the following section. Our calculations of κ_l are based on the relaxation-time approximation (RTA), which accounts for three-phonon scattering mechanisms and neglects other scattering processes like electron-phonon and phonon-defect scattering.⁷⁵ The RTA also does not consider the collective phonon excitations that appear in a full solution of the linearized Boltzmann transport equation (LBTE)⁷⁶ and thus tends to possibly underestimate κ_l .^{77,78} However, the omission of collective phonon



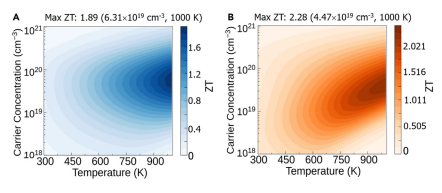


Figure 5. Predicted thermoelectric figures of merit (ZT) of p-type X₆NFSn₂ as a function of temperature and carrier concentration (A) Ca₆NFSn₂.

(B) Sr₆NFSn₂.

excitations partially compensates for the neglect of other scattering mechanisms, which often are in good agreement with experiments.^{26,79} In light of the lower LTC of Ca₆NFSn₂ and Sr₆NFSn₂, point defect scattering and nanostructuring (e.g., grain boundary effect) are not taken into consideration, as they may be not effective at reducing the LTC further, i.e., the mean-free path is already short, at less than 10 nm (see the following section).

Despite the low $\kappa_{\rm l}$, attention should be paid to the other component of the κ , $\kappa_{\rm e}$, as high σ results in high $\kappa_{\rm e}$. As shown in Figure 4, when $n_p \leq 10^{20}$ cm⁻³, the $\kappa_{\rm e}$ of Ca₆NFSn₂ is smaller than its $\kappa_{\rm l}$, and the latter thus dominates the thermal transport, which means that thermal losses through phonons would be more paramount for TE performance. When the carrier concentration exceeds 10^{20} cm⁻³, the electronic contribution becomes predominant above 210 K. For Sr₆NFSn₂, $\kappa_{\rm e}$ is smaller than $\kappa_{\rm l}$ when $n_p \leq 10^{19}$ cm⁻³ and T <810 K. At a high carrier concentration of 10^{20} cm⁻³, $\kappa_{\rm e}$ dominates above 190 K. When the carrier concentration further increases to 10^{21} cm⁻³, $\kappa_{\rm e}$ is larger than $\kappa_{\rm l}$ above 76 K, and thermal losses by charge carriers are dominant.

We combine the calculated electrical transport properties and *ks* to predict the TE ZTs for Ca₆NFSn₂ and Sr₆NFSn₂ as a function of carrier concentration and temperature (Figure 5). A maximum average ZT of 1.89 for p-type Ca₆NFSn₂ is predicted at a carrier concentration of 6.31 \times 10¹⁹ cm⁻³ and T = 1,000 K, and the maximum of the average ZT of p-type Sr₆NFSn₂ (2.28) is obtained for a carrier concentration at 4.47 × 10^{19} cm⁻³ and T = 1,000 K. ZT values of Ca₆NFSn₂ are isotropic, while those of Sr₆NFSn₂ are anisotropic. ZTs of Sr₆NFSn₂ are 2.25 and 2.34 along the a/b and c directions, respectively, which are close. These optimum hole carrier concentration and temperature values balance the electrical and thermal properties of two materials, which could be achieved experimentally based on a series of compounds with similar-composition Ca_2NI and Ba_2NX (X = F, Cl, Br) samples that are prepared at high temperature (\sim 1,000 K); Zn₂NX (X = Cl, Br) are synthesized at \sim 850 K and are thermally stable up to 1,023 K.⁵⁹⁻⁶¹ Moreover, the dominant competing phases of Ca_6NFSn_2 and Sr_6NFSn_2 are also stable at \sim 1,000 K.⁸⁰ Thus, the thermal stability of targeted compounds up to 1,000 K seems to be possible. The optimal holedoping concentration ($\sim 10^{19}$ cm⁻³) could be obtained via self-doping.^{69,70} These ZT values are the highest accessible so far for p-type perovskite-related TE materials at middle-high temperature,^{10,14,26,27,37} while they are lower than the ZT of SnSe (2.81) computed at a similar level of theory.⁶⁷ ZT values of 3.1 and 2.0 have been





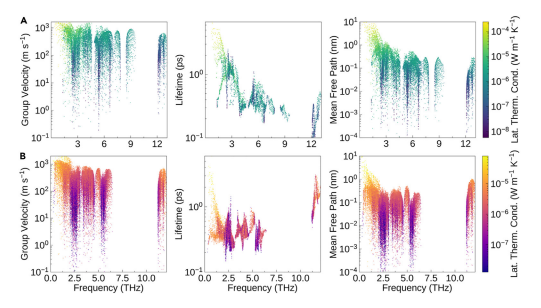


Figure 6. Analysis of the isotropically averaged modal group velocity, lifetimes, and mean-free paths of X_6NFSn_2 (A and B) Analysis of the isotropically averaged modal group velocity, lifetimes, and mean-free paths of X_6NFSn_2 (A) and Sr_6NFSn_2 (B) at 1,000 K, respectively. This analysis was performed using ThermoParser.¹⁰⁰ The data points are color-coded by the modal contributions to κ_I . κ_I value ranges are coded from blue to yellow and purple to yellow (low to high κ_I) for Ca₆NFSn₂ and Sr₆NFSn₂, respectively.

experimentally reported for polycrystalline and single-crystal SnSe at 783 and 773 K, respectively.^{15,68} The above results suggest that Ca_6NFSn_2 and Sr_6NFSn_2 are potentially promising p-type perovskite TE materials.

Origin of low LTC

FC expansions (up to the third order) were obtained by a combination of machine learning and first-principles calculations. Based on the FCs, κ_1 was computed within single-mode RTA (SM-RTA) as a sum of contributions from individual phonon modes λ following

$$\kappa_{l} = \frac{1}{NV} \sum C_{\lambda} \nu_{\lambda} \bigotimes \nu_{\lambda} \tau_{\lambda}$$
 (Equation 1)

where *N* is the number of wavevectors used for the Brillouin zone (BZ) integration and *V* is the volume of the unit cell. C_{λ} , ν_{λ} , and τ_{λ} are the modal heat capacities, the group velocities, and phonon lifetimes, respectively. Scatterplots of the three quantities against the phonon frequency can be used to evaluate the contributions of the individual modes to κ_l . The influence of C_{λ} on κ_l is not accounted for here as it is a shallow function of the phonon frequency, and the variation between phonon modes is thus minimal.

The group velocities of Ca₆NFSn₂ and Sr₆NFSn₂ are within the range of 10^{-1} to 10^{3} m s⁻¹ (left columns of Figure 6). The fastest modes (~1,000 m s⁻¹) are seen around 1–3 THz due to the dispersive phonon bands at these frequencies. The phonon lifetimes in Ca₆NFSn₂ and Sr₆NFSn₂ at 1,000 K and RT vary from 0.1 to 5 and 0.1 to 3 ps, respectively, as shown in the middle columns of the Figures 6 and S6. The modes with the longest modal phonon lifetimes (<4 ps) mainly appear in the low-frequency ranges (<3 THz). Notably, at 1,000 K, there is a group of modes below 3 THz in Ca₆NFSn₂ and Sr₆NFSn₂, of which the phonon lifetimes are shorter than 1 ps, indicative of strong phonon scattering and possibly large anharmonicity.⁸¹ The 1,000 K phonon lifetimes of the mid-frequency modes (3.5–9 THz for Ca₆NFSn₂ and 3–6.5

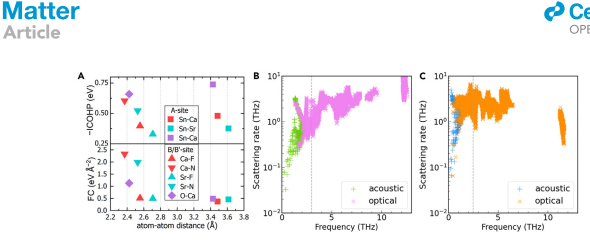


Figure 7. Analysis of the interatomic chemical bond strength

(A) Integrated crystal orbital Hamilton population (–ICOHP) (top) and force constants (FCs) (bottom) between anions centered at the A site and the B/B' site and cations at the X site.

(B) Mode-resolved scattering rates at 1,000 K for Ca_6NFSn_2 as a function of the phonon frequency.

(C) Mode-resolved scattering rates at 1,000 K for Sr_6NFSn_2 as a function of the phonon frequency.

THz for Sr₆NFSn₂) are all below 1 ps, and high-frequency N-based modes have shorter lifetimes in Ca₆NFSn₂ than in Sr₆NFSn₂. The phonon lifetime spectra at RT show a similar spread but longer lifetimes with respect to those at 1,000 K (see Figure S6). Strikingly, the overall RT spectra of ν_{λ} and τ_{λ} calculated from third-order perturbation theory are comparable to those of the hybrid perovskite CH₃NH₃Pbl₃, for which an ultralow LTC (0.086 W m⁻¹ K⁻¹) has been predicted.⁸² The combination of overall short modal lifetimes and low group velocities results in a relatively short modal mean-free path (<10 nm at 1,000 K and <100 nm at RT) (Figure 6 and S6) and thus leads to low averaged κ s of 2.94 (0.88 W m⁻¹ K⁻¹) and 0.83 W m⁻¹ K⁻¹ (0.25 W m⁻¹ K⁻¹) at RT (*T* = 1,000 K) for Ca₆NFSn₂ and Sr₆NFSn₂, respectively.

Besides the modal analysis, chemical bond characteristics, viz. chemical bonding strength and bond anharmonicity, of Ca₆NFSn₂ and Sr₆NFSn₂ are discussed. The interatomic bond strength of these new materials is evaluated based on the FCs and integrated crystal orbital Hamilton population (ICOHP) analyses, which imply the bond strength.⁸³ As a representative known antiperovskite, Ca₃OSn is taken into account as well. The bond strength generally decreases as the atom-atom distance increases. As shown in Figure 7A, Sn-X and X-N (X = Ca/Sr) bonds follow the trend, and their –ICOHPs are small with respect to SnSe,⁸⁴ indicative of the weak bond strength. However, X-F(X = Ca/Sr) bonds disobey the trend and exhibit the smallest -ICOHP among all bonds. In addition to the small -ICOHP, the FCs of F-X (X = Ca/Sr) are rather small. The FCs of X-Sn (X = Ca/Sr) are the smallest among all bonds, suggesting the role of Sn atoms acting as cage rattlers. Meanwhile, the O-Ca bonds are stiffer than X-N (X = Ca/Sr) bonds, which is consistent with the trend of LTC, i.e., the calculated LTC of Ca₃OSn (1.73 W m⁻¹ K⁻¹ at 1,000 K) is larger than those of Ca_6NFSn_2 and Sr_6NFSn_2 (0.83 and 0.25 W m⁻¹ K⁻¹ at 1,000 K). The combined analysis of -ICOHPs and FCs unravels the weak interatomic bond strength and soft lattice of Ca₆NFSn₂ and Sr₆NFSn₂. The soft lattice would impede phonon propagation and result in low LTC. Moreover, weak interatomic bond strength would introduce low-frequency vibrations, referred to as soft low-frequency optical phonons, which are indeed observed in Ca₆NFSn₂ and Sr₆NFSn₂ (Figure 2). These low-frequency optical modes do couple with acoustic phonons and contribute to the scattering rate, as shown in Figures 7B and 7C. The scattering rates of most phonon modes are within the range from 0.1 to 1 THz, which are comparable to those of SnSe and CH₃NH₃PbI₃.^{82,85} The optical and acoustic phonon resonant effect increases the scattering rate in low-frequency regions and scattering channels





for the heat-carrying acoustic phonons, which is supported by Sr_6NFSn_2 possessing a larger scattering rate but a lower κ_1 than Ca_6NFSn_2 (see Figures 7B and 7C). Additionally, as displayed in Figure S7 (cumulative LTC vs. frequency), the low-frequency modes (<3 THz), which are dominated by acoustic and optical modes, contribute to at least 65% of the LTC, further corroborating the phonon resonant scattering mechanism in these new double antiperovskites.

In addition to the weak chemical bond strength, lattice vibrational anharmonicity is another factor influencing the κ_l , as anharmonicity of the chemical bonds drives the phonon-phonon Umklapp and normal processes limiting the LTC.⁸⁶ Here, the mode Grüneisen parameter (γ) and the average Grüneisen parameter are employed to evaluate the bond anharmonicity. As displayed in Figure S8, the mode Grüneisen parameters of Ca₆NFSn₂ are positive. In contrast, the mode Grüneisen parameters of Sr₆NFSn₂ can be negative. The average RT Grüneisen parameters obtained based on the method given in Shao et al.⁸⁷ for Ca_6NFSn_2 , Sr_6NFSn_2 , and Ca_3OSn are 1.65, 1.37, and 1.54, respectively. The average Grüneisen parameters of Ca₆NFSn₂, Sr₆NFSn₂, and Ca₃OSn are on a par with some state-of-the-art TE materials such as PbTe (1.49).¹³ Compared to Ca₆NFSn₂, Sr₆NFSn₂ exhibits a lower κ_1 within the whole range of temperatures (Figure 4), and it possesses a smaller average Grüneisen parameter. Additionally, Sr has a larger atomic radius than Ca, which makes its valence shell and the electron clouds surrounding the Sr atoms larger than those of Ca. As seen in Figure S9, electrons are indeed more localized around Sr than Ca (higher electron localization function [ELF] for Sr), which possibly leads to a stronger lattice vibrational energy. Overall, the weak chemical bonding and strong bond anharmonicity are the macroscopic factors of the low κ_l in Ca₆NFSn₂ and Sr₆NFSn₂. Meanwhile phonon resonant scattering and Umklapp scattering are the dominant mechanisms affecting their κ .

Conclusion

In summary, we have discovered new multi-anion antiperovskites X_6NFSn_2 (X = Ca, Sr, and Ba) via anion mutation and global structure searches. Their phase stabilities were identified by first-principles calculations. Notably, Ca_6NFSn_2 and Sr_6NFSn_2 show superior electrical conductivities combined with low LTC at RT (<3 W m⁻¹ K⁻¹) and high temperature (<1 W m⁻¹ K⁻¹). The weak chemical bonding between the heavy-atom cage-rattler Sn and alkaline-earth metal (Ca and Sr) inducing low-frequency optical modes coupling with heat-carrying acoustic phonons in combination with strong bond anharmonicity results in the predicted low LTC in Ca_6NFSn_2 and Sr_6NFSn_2 . Owing to the high electrical conductivities and effective thermal insulation, the maximum *ZT* values of p-type Ca_6NFSn_2 and Sr_6NFSn_2 are predicted to be 1.89 and 2.28 at 1,000 K, respectively, which are currently higher than the reported perovskite oxides and halides. Our work of exploring multi-anion antiperovskites X_6NFSn_2 (X = Ca, Sr, and Ba) realizes the PGEC concept for promising TE materials within the vast family of perovskite structures.

EXPERIMENTAL PROCEDURES

Resource availability

Lead contact

Further information and requests for resources should be directed to and will be fulfilled by the lead contact, Dan Han (d.han@bham.ac.uk).

Materials availability

The materials in this study will be made available upon reasonable request.



Data and code availability

All data reported in this paper will be shared by the lead contact upon request. This paper does not report original code.

Computational methods

For the crystal structure predictions of X_6 NFSn₂ (X = Ca, Sr, and Ba), AIRSS and particle swarm optimization were utilized. AIRSS is powerful and highly parallel for locating low-energy structures with a given composition.^{49,50} The search of each composition (Ca₆NFSn₂, Sr₆NFSn₂, Ba₆NFSn₂) was constrained with the number of atoms in the reduced formula of 10. A comparative search including two reduced formula units (20 atoms) was performed for Ca₆NFSn₂, and it finds the same low-energy structures as the search with one reduced formula. For each composition, over 2,000 random structures were generated and relaxed. The plane wave density functional theory code CASTEP with core-corrected ultrasoft pseudopotentials generated as defined in the built-in QC5 library was used for the searching.⁸⁸ A plane-wave cutoff energy was set to 350 eV. The reciprocal space sampling was performed by using Monkhorst-Pack grids with a spacing of 0.07 $\times 2\pi$ Å⁻¹. CALYPSO was utilized for the crystal structure prediction of X_6 NFSn₂ (X = Ca, Sr, and Ba) as well.^{51,52} Thirty generations of structures with a population of 30 were considered. The local optimization calculations were performed three times for each structure generated by CALYPSO with a gradually increasing force convergence criteria. All the low-energy structures of X_6 NFSn₂ were found around the 20th generation, hence establishing the validity of this crystal structure search strategy.

Afterward, for further relaxations and property calculations of the low-energy structures generated by AIRSS, the Vienna Ab initio Simulation Package (VASP) was used.⁸⁹ For these calculations, a plane-wave cutoff energy of 520 eV, Γ -centered Monkhorst-Pack grids with a grid spacing of 0.03 × 2π Å⁻¹, and the projectoraugmented wave applying the parameterization of the exchange-correction potential given by Perdew-Burke-Ernzerhof (PBE) were used.⁹⁰ The convergence criteria for the energy and force were set to 10^{-5} eV and 0.01 eV/Å, respectively. The Heyd-Scuseria-Ernzerhof functional with 25% nonlocal Hartree-Fock exchange (HSE06) was employed for the electronic structure calculations.⁹¹ The spin-orbit coupling (SOC) effect was included. The postprocessing of electronic structure was carried out by sumo.⁹² Calculations of harmonic lattice dynamics (phonon spectrum and atom-projected phonon density of states) were calculated based on the frozen phonon method implemented in the Phonopy package.⁹³ 80-atom, 270-atom, and 270-atom supercells were used for phonon calculations of Ca₆NFSn₂, Sr₆NFSn₂, and Ba₆NFSn₂, respectively. The harmonic phonon spectrum of a larger supercell with 320 atoms of Ca₆NFSn₂ was calculated for testing the convergency, which shows that the phonon dispersions are reasonably converged with a small 80-atom supercell created from the primitive cell (Figure S10).

The electrical transport properties were calculated using the AMSET software package.⁶⁶ The code uses MRTA to calculate the scattering rates and carrier mobilities.⁹⁴ Here, three scattering processes, namely ADP, IMP, and POP scattering, were considered. The material parameters including high-frequency and static dielectric constants, elastic constants, polar optical phonon frequency, and deformation potentials required as input by this method were all obtained using first-principles calculations. To get the deformation potentials, calculations were performed using HSE06 functional. The high-frequency dielectric constants, static dielectric constants, and "effective polar optical phonon frequency" were obtained using density-functional perturbation theory (DFPT) with HSE06 functional.⁹⁵ The SOC effect





was not considered for deformation potential, as it is reported that the SOC effect has a negligible effect on this property of ABX_3 (A = K, Rb, Cs; B = Ge, Sn, Pb; X = Cl, Br, I) halide perovskites.⁹⁶ Elastic constants were calculated through the stress-strain approach with the PBE exchange-correlation functional. The transport properties need to be converged with respect to an interpolation factor, which controls the number of k-points in the interpolated band structures. Convergence testing results of Ca₆NFSn₂ are provided in Table S5. Dense Fourier interpolated meshes of 69 × 69 × 69 and 37 × 37 × 37 were chosen (the original input k-point meshes of the electronic band structures computed using DFT were 12 × 12 × 12 and 8 × 8 × 8) for Ca₆NFSn₂ and Sr₆NFSn₂ by balancing the computational cost and accuracy. Structures optimized with tight convergence criteria for energy (10⁻⁸ eV) and force (0.001 eV/Å) and the same tight setting were used for the calculations of harmonic phonon properties and electrical transport properties. Specific material parameters used to compute electrical transport properties are given in Table S6.

The $\kappa_{\rm I}$ was calculated by solving the phonon BTE under the SM-RTA.^{75,97} Phonon linewidths (inverse phonon lifetimes) were computed perturbatively to the third order by considering three-phonon scattering processes.⁷⁵ FCs (up to the third order) were obtained via the combination of machine learning (via the HIPHIVE package) and first-principles calculations. The utility of this approach was demonstrated via several examples like SnSe and monolayers of MoS₂.^{98,99} For Ca₆NFSn₂, 20 320-atom supercells with random displacements of standard deviation of 0.02 Å were generated. The PBE functional and the Γ point were used for calculating FCs. The cutoff radii for the two-, three-, and four-body clusters were 8, 5, and 4 Å, respectively. The linear least-squares method was used for fitting with a traintest ratio of 9:1. Similar settings were used for Sr₆NFSn₂, while larger supercells (360 atoms) were employed, and the DFT calculations were performed using the HSE06 functional. In both cases, the R^2 scores were higher than 0.99, indicating a high quality for the fittings. In addition, the phonon dispersion spectrum computed from the fitted (second-order) FCs matches well with that from standalone Phonopy calculations, as shown in Figure S11A. The LTCs calculated from the FCs that were obtained via HIPHIVE also agree well with those directly computed from the FCs obtained by Phono3py (see Figure S11B). The LTCs of Ca₆NFSn₂ were converged with respect to the q-point sampling density (Figure S11C), leading to $18 \times 18 \times 18$ and 15 x 15 x 15 meshes for Ca_6NFSn_2 and Sr_6NFSn_2 being chosen, respectively. ZT was obtained by using ThermoParser, the validation of which has been examined by literature.^{100–102}

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.matt. 2023.10.022.

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AUTHOR CONTRIBUTIONS

D.H. conceived the idea and wrote the draft. D.H. and B.Z. conducted the theoretical simulations and analyzed the data. Z.C. performed the crystal structure prediction by CALYPSO. K.B.S. provided software resources and assisted in the analysis of TE properties. S.S.R., W.S., and T.B. carried out validation and provided resources for the project. D.O.S. provided conceptualization and software resources. H.E. provided funding acquisition and project administration. All authors contributed to writing, review, and editing.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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