Thermoelectric cooling materials

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Solid-state thermoelectric devices can directly convert electricity into cooling or enable heat pumping through the Peltier effect. The commercialization of thermoelectric cooling technology has been built on the Bi₂Te₃ alloys, which have had no rival for the past six decades around room temperature. With the discovery and development of more promising materials, it is possible to reshape thermoelectric cooling technology. Here we review the current status of, and future outlook for, thermoelectric cooling materials.

In 1834, Jean Peltier discovered that there is a heating or cooling effect when electric current flows through the junction of two dissimilar metals¹. The principle of the Peltier effect is that when current flows across an interface, heat is absorbed (cooling) or rejected (heating) owing to the difference in the thermal energy transported by charge carriers (electrons or holes) in the two materials. Using the Peltier effect as a means of refrigeration was first seriously considered in 1911 by Altenkirch⁶. Early attempts to devise thermoelectric coolers, however, were unsuccessful because of the lack of high-performance materials¹. It was not until the 1950s that an appreciable Peltier cooling effect was realized in semiconductors⁴,⁵. Subsequently, there was an explosion of interest in thermoelectric cooling, as it was expected to be as efficient as vapour compression cooling once the materials were improved. However, it was soon realized that such refrigerators could not compete against the existing mechanical refrigerators in terms of price and efficiency. Consequently, most of the work on thermoelectric cooling stopped in the mid-1960s⁶. Interest re-emerged in the 1990s with the awareness of the environmental impact of the fluids used for vapour compression refrigeration⁷. Today, thermoelectric coolers are used in specific applications (see Box 1), and the market for thermoelectric modules is growing.

In the past two decades, however, the vast majority of research has focused on materials that are promising for power generation at medium and high temperatures, and there have been few reports on cooling below room temperature. Whereas the commercialization of thermoelectric cooling has been realized for several decades, thermoelectric power generation is still in its infancy. Thermoelectric modules for power generation demand high thermal stability of the materials at elevated temperatures, minimum diffusion of the elements under an extreme temperature gradient (several hundred kelvin across a couple of millimetres), and an excellent match of the coefficients of thermal expansion between the n- and p-type legs for minimum thermal stress, in addition to high thermoelectric performance. In contrast, thermoelectric coolers are often used near ambient temperature, and the temperature difference between the hot and cold sides is usually less than 100 K. Thermal stability, thermal stress and the diffusion of elements are thus not severe issues, and the engineering and manufacturing processes for thermoelectric cooling modules are well established. With the discovery of high-performance materials, it is possible to advance thermoelectric cooling technology.

In this Perspective, concepts underlying the thermoelectric cooling performance will be introduced first. Subsequently, we will summarize state-of-the-art materials and discuss Bi₁₋ₓSbx, Bi₂Te₃ alloys and Mg₂BiₓSbx in detail. Additionally, we will discuss ideas for identifying promising materials. We conclude with a discussion of the outlook and challenges of thermoelectric cooling materials and technology. As this Perspective is not intended to be comprehensive, readers interested in additional details are directed to other reports⁸-¹¹.

### Cooling performance

#### Coefficient of performance

The cooling power ($Q_C$) for a thermoelectric unicouple consisting of an n-type leg and a p-type leg is¹²

$$Q_C = (S_p - S_n) IT_C - \frac{1}{2} I^2 R - K (T_H - T_C)$$

(1)

where $I$ is the electrical current; $T_H$ and $T_C$ are the hot- and cold-side temperatures, respectively; $S_p$ and $S_n$ are the respective Seebeck coefficients for the p-type and n-type thermoelectric legs; and $R$ and $K$ are the total electrical resistance and thermal conductance of the unicouple, respectively. To maximize the cooling power, the unicouple is expected to have a large $(S_p - S_n)$ and small $R$ and $K$ values. As a result, the thermoelectric performance of an ideal unicouple can be measured by the figure of merit $Z$, where $Z = (S_p - S_n)^2 R^{-1} K^{-1}$.

The voltage of the external power source applied to the unicouple should be equal to the sum of the voltage drop due to the electrical resistance and the Seebeck voltage induced by the temperature difference. Therefore, the input electrical power ($P$) for a unicouple is¹³

$$P = I [RI + (S_p - S_n)(T_H - T_C)]$$

(2)

The cooling efficiency of a unicouple is quantified by the amount of cooling divided by the electrical power input. This efficiency is known as the coefficient of performance (COP). Since the cooling power and the input power are both functions of electrical current, COP can be maximized at an optimal electrical current. The maximum COP at given $T_H$ and $T_C$ values is¹³

$$\text{COP} = \frac{Q_C}{P} = \frac{T_C}{T_H - T_C} \sqrt{1 + \frac{Z T_w}{T_H - T_C} - \frac{T_H}{T_C}}$$

(3)

where $Z T_w$ is the averaged $Z$ between $T_H$ and $T_C$. The COP, which is dependent on temperature difference ($\Delta T = T_H - T_C$), is calculated (the solid lines) and shown in Fig. 1a. The symbols represent the experimental COP values of thermoelectric cooling systems.

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Refrigerants, as used in the vapour compression cycles, can deplete the ozone layer and are extremely potent greenhouse gases. In contrast, solid-state thermoelectric coolers are much more environmentally friendly. Thermoelectric cooling devices have the advantages of being reliable, compact in size, silent in operation, relatively lightweight, capable of providing rapid heating and cooling, and able to realize precise control of temperature. Additionally, a thermoelectric system has much more flexible cooling power, which can be easily tuned by electrical current, than a system using vapour compression refrigeration. A high COP can be achieved for a vapour compression refrigerator only when its cooling capacity is rather large; in practice, the COP is reduced with the lowering of cooling capacity, in contrast to the thermoelectric cooler, which has a COP that is more or less independent of its cooling capacity.

**Small-scale cooling.** Generally, conventional vapour compression refrigeration systems are not competitive, in terms of cost and efficiency, with thermoelectric cooling when the heat load is under 25 W (ref. 6). Currently, thermoelectric coolers have been mainly used in applications that require a small cooling power, such as portable refrigerators for carrying medical supplies and for food storage; cooling laser diodes and infrared, X-ray and gamma-ray detectors; thermal cyclers for DNA synthesizers; and temperature control of car seats.

**Large-scale cooling.** In large systems, in which the electrical power consumption is an essential consideration due to the cost of power, the COP of the cooling technology is critical. Because of the low COP of the thermoelectric cooling system (Fig. 1a), it is challenging for it to be economically viable for large-scale cooling applications. Nevertheless, large thermoelectric cooling systems with high cooling power have been developed for specific applications, such as air conditioning for large buildings, railway coaches and submarines. The use of large thermoelectric cooling systems has proved their technical reliability, but not their economic feasibility.

Hot-spot cooling. A refrigerator is normally used to maintain a heat source at a temperature below that of its surroundings. When the heat source must be prevented from rising above a temperature that is greater than ambient, a heat sink is usually employed rather than a refrigerator. Passive cooling is often realized by heat spreading using materials with high thermal conductivity. A copper heat spreader, for example, can remove heat equalling \( \kappa_{\text{Cu}} A\Delta T/L \), where \( \kappa_{\text{Cu}} \) is the thermal conductivity of copper and \( A \) and \( L \) are the cross-section and thickness of the copper plate, respectively. Thermoelectric cooling has been considered as a potential thermal management solution for electronics. The maximum heat extraction by a thermoelectric cooler (neglecting the heat removed by thermal conduction down the thermoelectric elements) is

\[
Q = \frac{A T_{\text{H}}^2}{2L} \left( \frac{S_p - S_n}{\sqrt{\rho_p} + \sqrt{\rho_n}} \right)^2 \tag{5}
\]

and the thermoelectric elements are assumed to have the same dimensions as the copper plate. If the thermoelectric device is to remove more heat than a copper plate, the following conditions should be satisfied:

\[
\frac{T_{\text{H}}^2}{2} \left( \frac{S_p - S_n}{\sqrt{\rho_p} + \sqrt{\rho_n}} \right)^2 > \kappa_{\text{Cu}} \Delta T \tag{6}
\]

Given the currently available materials, the thermoelectric device would perform better than the copper plate only when \( \Delta T < 2 \) at the \( T_{\text{H}} \) of 300 K (refs. 76, 77). Furthermore, owing to Joule heating of the thermoelectric legs, more heat is added to the hot side that must be carried to the environment. In fact, for hot-spot thermal management of electronics by lateral heat spreading, one should look for materials with not only a large power factor but also a high thermal conductivity, which is different from the traditional high-\( ZT \) materials.

A maximum temperature difference (\( \Delta T_{\text{max}} \)) between the hot and cold sides depends on the \( ZT \), and it can be expressed as

\[
\Delta T_{\text{max}} = \frac{1}{2} ZT_C^2 \tag{4}
\]

The \( ZT_C \)-dependent \( \Delta T_{\text{max}} \) of a unipolar is calculated and shown in Fig. 1b, where the \( \Delta T_{\text{max}} \) increases monotonically with \( ZT_C \). The \( ZT_C \) is ~0.7 for a unipolar of the commercial Bi\(_2\)Te\(_3\) alloys, and it corresponds to a \( \Delta T_{\text{max}} \) of ~70 K at the hot-side temperature of 300 K. The hot-side-temperature-dependent \( \Delta T_{\text{max}} \) for a single-stage cooler using Bi\(_2\)Te\(_3\) alloys is calculated and shown in Fig. 1c. Since the \( T_{\text{H}} \) is reduced with decreasing hot-side temperature, thermoelectric coolers become inefficient in the low-temperature range (\( T_{\text{H}} < 150 \) K, the grey shaded area). Even with 50% enhancement in \( ZT_C \) of both n-type and p-type Bi\(_2\)Te\(_3\) alloys, which would be a formidable challenge, the improvement in \( \Delta T_{\text{max}} \) is still insignificant in the low-temperature range. It should be noted that the \( \Delta T_{\text{max}} \) is achievable only at zero cooling power. In other words, for a given thermal load, which requires a finite cooling power, the temperature difference will be smaller than \( \Delta T_{\text{max}} \) (ref. 18). Although thermoelectric cooling at very low temperatures (in the liquid helium range or less) has been considered, the idea remains to be validated experimentally. In practice, most thermoelectric cooling devices are used in the vapour compression cycles, can deplete the ozone layer and are extremely potent greenhouse gases. In contrast, solid-state thermoelectric coolers are much more environmentally friendly. Thermoelectric cooling devices have the advantages of being reliable, compact in size, silent in operation, relatively lightweight, capable of providing rapid heating and cooling, and able to realize precise control of temperature. Additionally, a thermoelectric system has much more flexible cooling power, which can be easily tuned by electrical current, than a system using vapour compression refrigeration. A high COP can be achieved for a vapour compression refrigerator only when its cooling capacity is rather large; in practice, the COP is reduced with the lowering of cooling capacity, in contrast to the thermoelectric cooler, which has a COP that is more or less independent of its cooling capacity.

### Box 1: Thermoelectric cooling applications

In addition to COP, the capability to achieve a low temperature at the cold side for a given hot-side temperature (that is, the temperature difference) also quantifies the cooling performance of a unipolar. (red symbols) and vapour compression refrigerators and air conditioners (blue symbols). At a given temperature difference, COP increases with a higher \( ZT_C \). Therefore, improving the performance of the unipolar is crucial for promoting the efficiency of a thermoelectric cooler. It should be noted, however, that the actual COP values for the system are lower than the prediction based on \( ZT_C = 0.5 \), although the thermoelectric elements have higher measured \( ZT_C \). This discrepancy is mainly attributed to parasitic losses such as thermal resistances between the thermoelectric module and fluids and other heat leakages within the module. Compared with refrigerators and air conditioners based on the vapour compression cycle, thermoelectric cooling systems exhibit much lower COP values. To achieve energy efficiencies that compete with those of conventional refrigerators, an extraordinary leap in the performance of materials is required (\( ZT_C \approx 4 \)). Discouragingly, it is difficult to see how such a high \( ZT_C \) could be realized in the foreseeable future. As a result, thermoelectric cooling should be regarded not as a competitor to the conventional methods, but rather as a complementary technique for specialized applications.

### Maximum temperature difference

In addition to COP, the capability to achieve a low temperature at the cold side for a given hot-side temperature (that is, the temperature difference) also quantifies the cooling performance of a unipolar.
applications are close to room temperature, except for specific electronic cooling applications that achieve the required low temperatures by using multistage coolers (see Box 2).

State-of-the-art materials
The thermoelectric performance of a single n-type or p-type material is measured by $ZT = S^2\rho^{-1}k^{-1}T$, where $\rho$ is the electrical resistivity and $k$ is the thermal conductivity. Although many materials with $ZT$ above unity at medium and high temperatures have been discovered over the past two decades, progress in developing materials for thermoelectric cooling has been unsatisfactory. The state-of-the-art low-temperature and room-temperature thermoelectric materials, including n-type YbAl$_3$ (ref. 19), Cu$_{33,6}$Ni$_{66,4}$AgSe (ref. 20), Bi$_2$Te$_3$,Se$_x$ (ref. 21), Mg$_4$Bi$_2$Sb$_{15/3}$ (ref. 22) and Bi$_2$Te$_3$,Sb$_x$ (ref. 23), and p-type Bi$_{1-x}$Sb$_x$Te$_3$ (ref. 24), CePd$_{2.95}$ (ref. 25) and Ce(Ni$_{0.6}$Cu$_{0.4}$)$_2$Al$_3$ (ref. 26), are summarized in Figs. 2a and 2b, respectively. A further comparison of the figure of merit $zT$, as the materials’ intrinsic property, among several high-performance materials is shown in Fig. 2c. As can be seen, only Bi$_2$Te$_3$ alloys, Cu$_{33,6}$Ni$_{66,4}$AgSe and Bi$_2$Te$_3$,Sb$_x$ exhibit high performance near room temperature, and only Bi$_{1-x}$Sb$_x$ excels below 150 K. There is an obvious reason that realizing high performance near room temperature, and only Bi$_{1-x}$Sb$_x$, Mg$_4$Bi$_2$Sb$_{15/3}$ and the following discussion will be limited to these materials.

Bi$_{1-x}$Sb$_x$. Bismuth and antimony are semimetals with small band overlap energies (the overlap energy of the conduction and valence bands). Bismuth has a low density-of-states effective mass $(m^*_{n})$ of $-0.1m_0$ (where $m_0$ is the free electron mass) for the conduction band$^{27}$, thus leading to an exceptionally high room-temperature Seebeck coefficient of about $-70 \mu$V K$^{-1}$, and for single-crystalline bismuth it is about $-100 \mu$V K$^{-1}$ along the trigonal axis$^6$. Such favourable electronic properties give rise to large power factor $(S^2\rho^{-1/2})$ values of $\sim 77 \mu$W cm$^{-1}$ K$^{-2}$ at 300 K and $\sim 200 \mu$W cm$^{-1}$ K$^{-2}$ at 100 K in single-crystalline bismuth$^2$. Substitution of bismuth by antimony can greatly alter the band structure of bismuth, and Bi$_{1-x}$Sb$_x$ compositions with $0.07 < x < 0.22$ are semiconductors with a small bandgap that exhibits a maximum at around 15$\sim$17 at% antimony$^4$. The semiconducting Bi$_{1-x}$Sb$_x$ shows a large Seebeck coefficient and a substantially reduced thermal conductivity, and thus its thermoelectric performance is effectively improved$^{22,26}$. Single-crystalline Bi$_{1-x}$Sb$_x$ exhibits the highest $zT$ among all of the reported materials below 150 K (Fig. 2c), and it is promising for low-temperature thermoelectric cooling. Further substantial enhancement in the thermoelectric performance of Bi$_{1-x}$Sb$_x$ can be achieved by applying a transverse magnetic field (with a peak $B$ of 8.6$\times$10$^{-1}$ T at 100 K)$^{25}$, benefiting from the thermomagnetic effect. The Ettingshausen effect, one such thermomagnetic effect, appears when a transverse magnetic field acts on electric current and produces a temperature gradient in the mutually perpendicular direction. Unlike the Peltier effect, in which the contributions from electrons and holes counteract one another, electrons and holes jointly contribute to the Ettingshausen effect. Consequently, semimetals are ideal candidates for Ettingshausen cooling, and single-crystalline bismuth showed a large $\Delta T$ of $\sim 100$ K at the hot-side temperature of $\sim 300$ K on application of a magnetic field of $\sim 11$ T (ref. 20). Further details on Bi$_{1-x}$Sb$_x$ for thermoelectric and thermomagnetic cooling can be found elsewhere$^{22,24}$.

Bi$_2$Te$_3$ alloys. The Peltier effect was initially demonstrated by freezing water droplets into ice at the junction of bismuth and antimony$^6$. These semimetals were then used for early measurements of Peltier cooling, with a unipolar consisting of Bi$_4$Sb$_3$ and Bi$_4$Sn$_4$, and showed a small $\Delta T$ of only 10 K (ref. 14). It was not until the 1950s that semiconductors were recognized as promising materials for thermoelectric cooling, with Bi$_2$Te$_3$, as the best-in-class candidate. The unipolar of p-type Bi$_2$Te$_3$ and n-type Bi was found to exhibit a $\Delta T_{\text{max}}$ of 26 K at the hot-side temperature of 285 K (ref. 14). Shortly afterward, the n-type Bi$_2$Te$_3$ was produced, and a $\Delta T_{\text{max}}$ of 40 K was observed for the unipolar of n- and p-type Bi$_2$Te$_3$.

Fig. 1 | Thermoelectric cooling performance. a, A comparison of the COP values between thermoelectric cooling systems and vapour compression refrigerators and air conditioners$^{22,24}$. The solid lines are calculated by assuming that the hot-side temperature is 300 K, and the filled symbols are experimental values. The COP is calculated using constant $ZT_m$ values, which implicitly assume that the thermoelectric transport properties are independent of temperature. b, Relationship between the $ZT_m$ of the unicouple and the maximum temperature difference at the hot-side temperature of 300 K. c, Relationship between the maximum temperature difference and the hot-side temperature for the unicouple based on the state-of-the-art Bi$_2$Te$_3$ alloys. The measured temperature-dependent thermoelectric properties of commercial Bi$_2$Te$_3$ alloys are used for this calculation. The blue line shows the theoretical $\Delta T_{\text{max}}$ for those alloys given a 50% increase in $ZT_m$. 

The thermoelectric performance of a single n-type or p-type material (Fig. 2d). Polycrystalline bismuth exhibits a room-temperature Seebeck coefficient of about $-70 \mu$V K$^{-1}$, and for single-crystalline bismuth it is about $-100 \mu$V K$^{-1}$ along the trigonal axis$^6$. Such favourable electronic properties give rise to large power factor $(S^2\rho^{-1/2})$ values of $\sim 77 \mu$W cm$^{-1}$ K$^{-2}$ at 300 K and $\sim 200 \mu$W cm$^{-1}$ K$^{-2}$ at 100 K in single-crystalline bismuth$^2$. Substitution of bismuth by antimony can greatly alter the band structure of bismuth, and Bi$_{1-x}$Sb$_x$ compositions with $0.07 < x < 0.22$ are semiconductors with a small bandgap that exhibits a maximum at around 15$\sim$17 at% antimony$^4$. The semiconducting Bi$_{1-x}$Sb$_x$ shows a large Seebeck coefficient and a substantially reduced thermal conductivity, and thus its thermoelectric performance is effectively improved$^{22,26}$. Single-crystalline Bi$_{1-x}$Sb$_x$ exhibits the highest $zT$ among all of the reported materials below 150 K (Fig. 2c), and it is promising for low-temperature thermoelectric cooling. Further substantial enhancement in the thermoelectric performance of Bi$_{1-x}$Sb$_x$ can be achieved by applying a transverse magnetic field (with a peak $B$ of 8.6$\times$10$^{-1}$ T at 100 K)$^{25}$, benefiting from the thermomagnetic effect. The Ettingshausen effect, one such thermomagnetic effect, appears when a transverse magnetic field acts on electric current and produces a temperature gradient in the mutually perpendicular direction. Unlike the Peltier effect, in which the contributions from electrons and holes counteract one another, electrons and holes jointly contribute to the Ettingshausen effect. Consequently, semimetals are ideal candidates for Ettingshausen cooling, and single-crystalline bismuth showed a large $\Delta T$ of $\sim 100$ K at the hot-side temperature of $\sim 300$ K on application of a magnetic field of $\sim 11$ T (ref. 20). Further details on Bi$_{1-x}$Sb$_x$ for thermoelectric and thermomagnetic cooling can be found elsewhere$^{22,24}$.
Box 2 | Thermoelectric cooling modules

Most thermoelectric modules contain a horizontal array of thermoelectric elements: that is, a single-stage module. When several such modules are stacked in a vertical array, a multistage module can be fabricated. Typically, a multistage module is a pyramid shape because the heat dissipated by the upper stage should be removed by the lower stage, and thus more legs are required for the lower stage. The possibility of using such a cascade cooler for augmenting cooling performance was initially proposed by Altenkirch \( ^2 \) and experimentally confirmed later \( ^6 \).

The COP of a cascade cooler is \( ^{18} \)

\[
\text{COP} = \frac{1}{\Pi_i (1 + \frac{1}{\text{COP}_i})} - 1
\]

where \( \text{COP}_i \) is the COP of the \( i \)th stage, and \( n \) is the total number of stages. This reaches a maximum \( (\text{COP}_{\text{max}}) \) when the temperature differences at each stage are chosen so that each stage has an identical COP. The \( \text{COP}_{\text{max}} \) is

\[
\text{COP}_{\text{max}} = \frac{1}{(1 + \frac{1}{\text{COP}_1})^n - 1}
\]

The temperature difference achievable by a cascade cooler is determined by the separate working temperature intervals at each stage \( ^{18} \):

\[
\Delta T = \sum_{i=1}^{n} \Delta T_i
\]

where \( \Delta T_i \) is the temperature difference of the \( i \)th stage.

When the temperature difference is large, the COP can be effectively improved for multistage modules compared with single-stage modules (part a of the figure). Additionally, the \( \Delta T_{\text{max}} \) of a cascade cooler can be notably enhanced with an increasing number of stages, according to Eq. (9). A comparison between the calculated and measured \( \Delta T_{\text{max}} \) for modules with different numbers of stages is shown in part b of the figure. A single-stage thermoelectric module can obtain a temperature difference of \( \sim 70 \) K at the hot-side temperature of \( 300 \) K. In comparison, a six-stage module can achieve a temperature difference of up to \( 130 \) K at the same hot-side temperature \( ^{19} \). The cooling performance can be further improved by adopting different compositions or materials at each stage — for example, using Bi\( \text{\textsubscript{x}} \)-Sb for the upper stage and Bi\( \text{\textsubscript{x}} \)-Te alloys for the lower stage. Additionally, it is possible to use optimally graded or segmented legs to augment the thermoelectric cooling performance \( ^{8} \). However, the cooling capacity of a multistage module is usually limited, so it is primarily used for spot cooling, particularly for detectors, in which the lowered temperature effectively reduces the noise and increases the sensitivity \( ^{14} \).

Comparison of cooling performance between single-stage and multistage modules. a. The calculated temperature-difference-dependent COP for single-stage and multistage modules. b. Comparison between calculated and experimental \( \Delta T_{\text{max}} \) (refs. 67,79) for modules with different numbers of stages. The error bars in the experimental data are obtained by averaging the performance of different commercial thermoelectric modules 67,79. All of the calculations are based on the temperature-dependent thermoelectric properties of commercial Bi\( \text{\textsubscript{x}} \)-Te\( \text{\textsubscript{3}} \) alloys, and it is assumed that the hot-side temperature is \( 300 \) K.

(Ref. 18). Better cooling performance was later reported for zone-melted Bi\( \text{\textsubscript{x}} \)-Te\( \text{\textsubscript{3}} \) and was attributed to its improved homogeneity, adequately controlled impurities and preferred orientation 22. By this time, the solid-solution alloying strategy for reducing lattice thermal conductivity had just been identified 18. This was applied to Bi\( \text{\textsubscript{x}} \)-Te\( \text{\textsubscript{3}} \) and improvements in \( zT \) were obtained. As a result, \( \Delta T_{\text{max}} \) above 60 K was realized for uniconicals of Bi\( \text{\textsubscript{x}} \)-Te\( \text{\textsubscript{3}} \) alloys 34.

In the past two decades, improvements in the thermoelectric performance of Bi\( \text{\textsubscript{x}} \)-Te\( \text{\textsubscript{3}} \) alloys have been reported 18. However, such \( zT \) improvements have not yet been engineered into applications, and the performance of commercial coolers has seen little change over this time. At this point, further significant improvement in \( zT \) of the Bi\( \text{\textsubscript{x}} \)-Te\( \text{\textsubscript{3}} \) alloys seems to be an ambitious goal. For commercial applications, the large-scale synthesis of Bi\( \text{\textsubscript{x}} \)-Te\( \text{\textsubscript{3}} \) alloys with improved mechanical properties using a cost-effective method with high quality control is as important as \( zT \) enhancement. Further details on the development of Bi\( \text{\textsubscript{x}} \)-Te\( \text{\textsubscript{3}} \) alloys for thermoelectric cooling can be found elsewhere 38. Commercial thermoelectric cooling has been exclusively based on Bi\( \text{\textsubscript{x}} \)-Te\( \text{\textsubscript{3}} \) alloys, with n-type Bi\( \text{\textsubscript{x}} \)-Se\( \text{\textsubscript{3}} \) exhibiting slightly inferior \( zT \) to that of p-type Bi\( \text{\textsubscript{x}} \)-Sb\( \text{\textsubscript{3}} \) Te. Thus, identifying new n-type materials with high thermoelectric performance is of great importance for potentially replacing the n-type Bi\( \text{\textsubscript{x}} \)-Se\( \text{\textsubscript{3}} \).
Compounds based on MgSb have long been regarded as persistent p-type semiconductors. However, n-type MgSb2 was successfully obtained in the 1950s, and the importance of Mg stoichiometry in determining the conduction type was noted. Unfortunately, this result had been overlooked for decades, and n-type MgSb2-based materials with high thermoelectric performance were reported only recently. Initially, it was the successful conversion from p-type to n-type conduction and the high zT in the mid-temperature range that made this compound highly interesting. Subsequently, chemical doping and optimization of processing conditions greatly improved its thermoelectric performance around room temperature, making n-type Mg1−xSb x Te comparable to the state-of-the-art n-type Bi2Te1−xSex (ref. 22). Most recently, further improvements in the average zT of Mg1−xSb x Te were reported (ref. 23). In contrast to the semiconducting MgSb, MgBi is a semimetal with a small band overlap energy. Unexpectedly, the band overlap energy is 0.05 eV, the conduction-band density-of-states effective mass is 0.5m0, and the carrier concentration is 2.5×10^{19} cm^{-3} (2×10^{19} cm^{-3}). A very similar solid line can be calculated by using the alternative values in the brackets. A comparison of the Seebeck coefficient among several semimetals: single-crystalline bismuth (along the trigonal direction), polycrystalline bismuth, polycrystalline Mg3Bi2 (ref. 22), CoSi (ref. 22), and single-crystalline Cd3As2 (ref. 22). The relationship between temperature and optimal bandgap energy. Values of band overlap energy for several semimetals — MgBi1−xSb x Te, bismuth (~30 meV), and CoSi (~40 meV) — are also shown. The band overlap energy of MgBi is estimated from the temperature-dependent resistivity, and the corresponding temperature is assumed to be 300 K.

Recent, single-crystalline MgBi1.25Sb0.75 was reported to have an improved low-temperature thermoelectric performance around room temperature, making n-type Mg3Bi2 highly interesting. Subsequently, chemical doping and optimization of processing conditions greatly improved its thermoelectric performance around room temperature, making n-type Mg1−xSb x Te comparable to the state-of-the-art n-type Bi2Te1−xSex (ref. 22). Most recently, further improvements in the average zT of Mg1−xSb x Te were reported (ref. 23). In contrast to the semiconducting MgSb, MgBi is a semimetal with a small band overlap energy. Unexpectedly, the band overlap energy is 0.05 eV, the conduction-band density-of-states effective mass is 0.5m0, and the carrier concentration is 2.5×10^{19} cm^{-3} (2×10^{19} cm^{-3}).

**Future research in thermoelectric cooling materials**

Re-evaluation of existing materials and strategies. Investigating the low-temperature thermoelectric properties of existing materials and strategies is also important for practical application, and thus more effort should be devoted to related research.
medium- and high-temperature materials would be a meaningful step toward identifying promising cooling materials. Indeed, certain materials that exhibit high $zT$ at medium temperature also demonstrate promising thermoelectric performance below 300 K, examples being PbTe and Mg$_2$Bi$_2$Sb$_2$. It is surprising that the low-temperature thermoelectric properties of many medium- and high-temperature materials have not yet been extensively studied. In addition, since the optimized carrier concentration reduces with decreasing temperature, materials with maximum $zT$ at low temperatures should be less heavily doped. It is essential to minimize contamination by impurities during the synthesis and to reduce the concentration of defects by controlling the composition and preparation conditions.

Additionally, re-evaluation of established strategies that can enhance the $zT$ of medium- and high-temperature materials is essential. As the temperature declines below 300 K, the dominant scattering mechanisms for electrons (acoustic phonon scattering) and phonons (umklapp scattering) can be overshadowed by defect scattering. Thus, it is pivotal to determine whether certain phonon scattering strategies can still improve the ratio of mobility to lattice thermal conductivity at low temperatures. Isoelectronic alloying has proved to be effective in reducing the lattice thermal conductivity and enhancing $zT$, as is the case in Bi$_{1-x}$Sb$_x$, Bi$_2$Te$_3$ alloys and Mg$_2$Bi$_2$Sb$_2$. However, defects such as dislocations or charged point defects (vacancies and interstitials) can scatter electrons, substantially reducing the carrier mobility at low temperatures. Nanostructures with dimensions in between the electron and phonon mean free paths should scatter more phonons than electrons. However, designing specific nanostructures without introducing additional defects is essential. Currently, the effect of defects on thermoelectric transport properties is not fully understood, so investigating the defect scattering of electrons and phonons is of particular importance. Additionally, band engineering, which is considered an effective approach to improve the power factor, should be re-evaluated. Maximizing the valley degeneracy by converging multiple electronic bands (that is, aligning the band edges within the energy difference of $\sim 2k_B T$) becomes more difficult as the temperature decreases. In addition, intervalley scattering, which is often overlooked, could be strong in this case, partially offsetting the benefit of band convergence.

### Exploration of semiconductors with very narrow bandgap

Thermal activation of electrons from the valence band into the conduction band will result in the production of equal numbers of electrons and holes (that is, bipolar conduction) and could greatly reduce $zT$. To minimize this deleterious effect, a finite bandgap has always been regarded as indispensable for high thermoelectric performance. The optimal bandgap energy ($E_g$) is predicted to be between $6k_B T$ and $10k_B T$ (refs. [12,13,16]), as shown in the grey shaded area in Fig. 2f. The optimized $E_g$ is in the range 0.16–0.26 eV at 300 K, which explains why narrow-bandgap semiconductors have long been regarded as the most promising candidates for thermoelectric applications. This optimal $E_g$ was predicted by assuming that the electrons and holes have comparable mobilities and density-of-states effective masses — that is, an electron-to-hole weighted mobility ratio of unity. This assumption is not always valid, since the electron-to-hole weighted mobility ratio can be considerably higher or lower than unity. In other words, optimal $E_g$ in the range between $6k_B T$ and $10k_B T$ should not be considered the golden rule, and expanding the exploration to semiconductors with a very small $E_g (< 6k_B T)$ is also meaningful.

### Discovery of promising semimetals

The concept of optimal $E_g$ also implies the exclusion of semimetals (with zero or negative $E_g$) as promising thermoelectric materials. Admittedly, semimetals are unlikely to exhibit a high $zT$ at medium or high temperatures, owing to their notable bipolar conduction. Nevertheless, semimetals such as bismuth[22], Mg$_2$Bi$_2$ (ref. [23]), CoSi (ref. [24]) and Cd$_2$As$_2$ (ref. [25]) exhibit promising thermoelectric properties near room temperature and at low temperatures. The asymmetry in electronic band structures, for example, the disparities in the band effective mass and valley degeneracy between the conduction and valence bands, and the accompanying asymmetry in the electron and hole mobilities, as quantified overall by the electron-to-hole weighted mobility ratio, are responsible for their thermoelectric performance. Such asymmetries have been observed in various semiconductors[26] and are expected to exist in many more semimetals[27]. In semiconductors, the effect of the electron-to-hole weighted mobility ratio on the transport properties of semiconductors can be overshadowed by the greatly increased majority carrier concentration through chemical doping. For semimetals, in contrast, owing to the simultaneous presence of the comparable concentration of electrons and holes, the weighted mobility ratio has great importance for the thermoelectric properties. Thus, this parameter can guide the search for promising semimetals. In principle, the weighted mobility ratio could be manipulated if one type of carrier can be preferentially scattered, and thus it is possible to further enhance the $zT$ (ref. [28]).

The band overlap energy is another critical parameter that needs to be taken into account, and it should be small enough ($< 0.1 \text{ eV}$) to limit the carrier concentration. A lower carrier concentration ($< 10^{20} \text{ cm}^{-3}$) is essential for a large Seebeck coefficient and a low electronic thermal conductivity. Alloying a semimetal with a semiconductor enables fine-tuning of the band overlap energy or can even open a small bandgap that allows the shifting of the peak performance across a certain temperature range. Such an alloying effect is also beneficial for reducing the lattice thermal conductivity and improving the $zT$.

### Looking ahead to improved materials and systems

Only a small fraction of research effort has been focused on thermoelectric cooling materials in the past two decades, and this partially accounts for the sluggish developments in the field. The slow progress can be easily seen in the dwindling numbers of publications per year on thermoelectric cooling, and it is also evidenced by the difficulty in finding the materials’ low-temperature thermoelectric properties among the literature. More focused research effort will be required for future advances.

Bi$_2$Te$_3$ alloys have been the state-of-the-art thermoelectric cooling materials for six decades, and commercially available ingots exhibit $zT$ of unity around room temperature. Although improved values of room-temperature $zT$ have been reported for Bi$_2$Te$_3$ alloys, the low-temperature properties have often been overlooked, and the cooling performance cannot be assessed. Currently, it is challenging to improve the Bi$_2$Te$_3$ alloys further, so discovering promising materials that exhibit higher $zT$ is critical. The recently discovered n-type Mg$_2$Bi$_2$Sb$_2$ represents encouraging progress for such an exploration, but its average $zT$ remains similar to that of commercial n-type Bi$_2$Te$_3$-Se$_2$, and further improvement of performance is necessary. As a promising candidate, n-type Ag$_2$Se with high thermoelectric performance around ambient temperature was reported[29], but its low-temperature properties have been much less studied, and its cooling performance has not yet been characterized. In addition, since the cooling efficiency of a thermoelectric module hinges on both its n-type and its p-type legs, it is also vital to identify high-performance p-type materials. Bi$_{1-x}$Sb$_x$Te$_3$ has remained the state-of-the-art p-type material, exhibiting a room-temperature $zT$ superior to that of all other p-type materials. The p-type CsBi$_2$Te$_3$ is the only material that outperforms Bi$_{1-x}$Sb$_x$Te$_3$ below 250 K (ref. [30]) (Fig. 2b), and thus it is a good starting point in the search for new p-type materials. But further studies on CsBi$_2$Te$_3$ have rarely been reported, and its cooling performance has not yet been evaluated.
Since it is the efficiency of a thermoelectric cooling system that eventually determines its applications, optimizing the system architecture is of equal importance to the development of materials. Minimizing the thermal resistance and improving the performance of heat-exchanging components are crucial for improving the COP. The relatively high cost of thermoelectric cooling systems also limits its wide application\(^9\), so reducing the related expenses is of great economic importance. Thermoelectric materials that are inexpensive and abundant will be favourable, and lowering the cost of heat exchangers and other components is also critical. With concerted progress in materials, devices and systems, both COP improvement and cost reduction are expected and could eventually propel advances in thermoelectric cooling technology.

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