Metallic–covalent bonding conversion and thermoelectric properties of Al-based icosahedral quasicrystals and approximants

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Received 31 January 2014
Accepted for publication 11 June 2014
Published 2 July 2014

Abstract
In this article, we review the characteristic features of icosahedral cluster solids, metallic–covalent bonding conversion (MCBC), and the thermoelectric properties of Al-based icosahedral quasicrystals and approximants. MCBC is clearly distinguishable from and closely related to the well-known metal–insulator transition. This unique bonding conversion has been experimentally verified in 1/1-AlReSi and 1/0-Al12Re approximants by the maximum entropy method and Rietveld refinement for powder x-ray diffraction data, and is caused by a central atom inside the icosahedral clusters. This helps to understand pseudogap formation in the vicinity of the Fermi energy and establish a guiding principle for tuning the thermoelectric properties. From the electron density distribution analysis, rigid heavy clusters weakly bonded with glue atoms are observed in the 1/1-AlReSi approximant crystal, whose physical properties are close to icosahedral Al–Pd–TM (TM: Re, Mn) quasicrystals. They are considered to be an intermediate state among the three typical solids: metals, covalently bonded networks (semiconductor), and molecular solids. Using the above picture and detailed effective mass analysis, we propose a guiding principle of weakly bonded rigid heavy clusters to increase the thermoelectric figure of merit ($ZT$) by optimizing the bond strengths of intra- and inter-icosahedral clusters. Through element substitutions that mainly weaken the inter-cluster bonds, a dramatic increase of $ZT$ from less than 0.01 to 0.26 was achieved. To further increase $ZT$, materials should form a real gap to obtain a higher Seebeck coefficient.

Keywords: quasicrystal, icosahedral cluster, MEM/Rietveld, electron density, chemical bond, metal-insulator transition, thermoelectric material

1. Introduction
Since the discovery of icosahedral quasicrystals in liquid-quenched Al–Mn alloys by Shechtman et al [1], extensive studies have been performed on the structural [2], electrical [3, 4], thermal [5], and magnetic properties [6] of many stable quasicrystals and related approximant crystals [7]. Quasicrystals and their approximants are recognized as complex metallic phases constructed from clusters with icosahedral symmetry, which cannot coexist with periodicity. In particular, Al-based quasicrystals, which are called Al-based icosahedral cluster solids (ICSs), exhibit non-metallic behavior. This can be understood by the formation of a deep pseudogap near the Fermi energy and the electron localization effect arising from the quasiperiodic arrangement [8]. The origins of the pseudogap and the stabilization mechanisms of...
quasicrystals are the Fermi surface–Brillouin zone interactions, the Hume-Rothery mechanism [9–12], and covalent bonds including sp–d hybridization [13, 14]. From comparative studies of Al- and B-based ICSs, the concept of metallic–covalent bonding conversion (MCBC) has been proposed by Kimura et al [13, 15], and this is the first topic of this review article. For the related approximants, band structure calculations are a powerful tool because they provide beneficial information about atomic orbital hybridization, the electron density distribution, and pseudogap formation in the electronic density of states (DOS) of both real materials and hypothetical structure models [16, 17].

Al-based icosahedral quasicrystals represented by Al–Pd–Re and Al–Pd–Mn ternary systems exhibit a semiconductor-like transport property with high Seebeck coefficient (S) [18]. Because of their giant unit cells and complex crystal structures, their thermal conductivities are close to 1 W/(m K), which is comparable to the thermal conductivity of glass materials as demonstrated by Chernikov et al [5]. Therefore, one of the possible applications of quasicrystals is as thermoelectric materials, which is the second topic of this review article. It is possible to obtain a high dimensionless figure of merit (ZT), $ZT = S^2\sigma T/k$, where $\sigma$, $k$, and $T$ are the electrical conductivity, thermal conductivity, and temperature, respectively, by optimizing the individual parameters [19]. Pope et al first reported the $ZT$ for the Al–Pd–Mn quasicrystal in 1999 [20]. In 2002, the composition dependence of the thermoelectric properties of Al–Pd–Re quasicrystals was systematically investigated by Kirihara and Kimura [21]. Maciá performed theoretical calculations of various types of quasicrystals and approximants using an analytical model, and recently published a review [22]. For Al-based approximant crystals, Takeuchi et al reported the band structure and phonon dispersion calculations, and discussed the intrinsic properties caused by a deep pseudogap [23, 24].

This review contains three parts. First, we outline the characteristic features of both Al- and B-based ICSs and discuss their possible applications, such as thermoelectric [21], superconducting [25], and high-resistive chip [26] materials. Next, we explain the bonding nature in the related approximants revealed by the maximum entropy method and Rietveld refinement for powder x-ray diffraction (MEM/Rietveld) analysis. Discussions include the relationship between the bonding nature, electronic structure, and electrical and thermal properties. Finally, we overview the thermolectric properties and $ZT$ enhancement using the guiding principle of weakly bonded rigid heavy clusters (WBRHCs) proposed by Kimura et al [27].

### 2. ICS: characteristic features and potential applications

Figure 1 shows an outline of the characteristic features and potential applications of Al- and B-based ICSs constructed by Kimura et al [15].

1. ICSs have complex structures, such as quasicrystals or giant-unit-cell crystals, because the icosahedral symmetry cannot coexist with periodicity.

2. B and Al are located at both sides of the boundary between elements, which have covalent and metallic bonds. Among the group-13 elements, an elemental crystal of only boron has covalent bonds and is a semiconductor. Of course, pure Al has metallic bonds and its crystal is a metal. However, the 1/1-AlReSi approximant, which consists of Al$_{12}$ icosahedra without a central atom like B$_{12}$, exhibits a covalent bonding nature and shows semiconductor-like transport properties. On the other hand, the isolated Al$_{13}$ icosahedral cluster and the Al$_{12}$Re icosahedron in the 1/0-Al$_{13}$Re approximant have metallic bonding nature with an atom in the center of the icosahedra.

The latter crystal has metallic transport properties. Therefore, MCBC is expected. The details of the chemical bonding nature of Al-based approximants are discussed in section 3.

3. ICSs can have a very deep pseudogap near the Fermi energy because of the high isotropy of the icosahedral symmetry according to the Hume-Rothery mechanism. The opposite of (3): ICSs can have a high density of states at the Fermi energy because of high degeneracy of electronic states caused by the high symmetry of the structure.

Feature (4) is advantageous for high superconducting transition temperature materials. Indeed, a superconducting transition was recently discovered in Li-doped α-boron [25]. One of the possible applications using the bonding conversion (feature (2)) is high-resistive chip materials [26]. V-doped amorphous boron can simultaneously have a relatively high electrical resistivity and a very small temperature dependence of the electrical resistivity. The strong covalency would deepen the pseudogap, which results in a high $S$. Using this bonding conversion can control the transport properties, such as thermoelectric materials.
as the coexistence of high $\sigma$ and large $S$ values [18]. Furthermore, five-fold symmetry conflicting with the periodicity decreases $\kappa$ because of a giant unit cell and its complex crystal structure. These features are favorable for thermoelectric materials [21], which can generate power through the direct conversion of heat energy into electrical energy.

Table 1 lists the differences between structurally simple and complex solids. Structurally simple solids have small unit cells and only one atomic site for each element, while structurally complex solids are giant-unit-ell crystals, quasicrystals, and amorphous solids with many types of atomic site for each element. These structurally complex solids are considered to be nanoscale, cluster-based composite materials and there is the possibility for the coexistence of functions that cannot coexist in structurally simple solids, as shown in Figure 2. For example, (i) magnetism and superconductivity, (ii) high electrical and low thermal conductivity, (iii) metallic and covalent bonds, and (iv) electron donors and electron acceptors. Feature (ii) agrees with the well-known guiding principle of phonon glass, electron crystals (PGECs) proposed by Slack [28] for high-performance thermoelectric materials such as clathrates [29] and skutterudites [30]. ICSs possess features (iii) and (iv), and (iii) is favorable for high-performance thermoelectric materials, as described in section 4.

**Table 2.** Comparison between (a) the metal–insulator transition and (b) metallic–covalent bonding conversion.

| (a) Metal ($\sigma \neq 0$) | Insulator (semi-conductor) ($\sigma = 0$) |
|---------------------------|----------------------------------------|
| Density of states at $E_F$, $D(E_F)$ | Finite ($n$ or $D(E_F)=0$) and or |
| Wave function at $E_F$ | Extended or or or Localized ($\mu$ or $\nu_F=0$) |

(b) Metallic bond Covalent bond

| Share of electrons | Many atoms | 2 atoms |
| Bond direction | Isotropic | Directed |
| Packing fraction | High ($\sim 0.75$) | Low ($\sim 0.34$) |
| Typical structure | Closed-packed structure fcc, hcp | Diamond structure |
| Icosahedral cluster | 13 atoms, Al$_{13}$ | 12 atoms, Al$_{12}$ |

The electrical conductivity ($\sigma$) is expressed as $\sigma = n e u = \frac{e^2}{3} \nu_F D(E_F)$, where $n, e, \mu, \nu_F, D(E_F)$ are the carrier density, elemental charge, mobility, relaxation time, Fermi velocity, density of states, and Fermi energy, respectively.

To experimentally investigate MCBC, we performed a MEM/Rietveld analysis using the BL02B2 beamline at the SPring-8 synchrotron facility (Hyogo, Japan). This method is widely used to visualize the electron density distribution between atoms in real-space, and was developed by the group of Sakata and Takata [31–33]. First, we investigated the pure elements Si and Al as typical examples of covalent and metallic bonding networks, respectively. Figure 3 shows the equidensity surfaces for pure Si and Al by MEM/Rietveld analysis. Covalently bonded (sp$^3$-hybridized) electrons can be clearly seen between the atoms in Si even in the high equi-density surface ($0.40 \text{ e A}^{-3}$). On the other hand, such bonding electrons are not observed and the electron density distribution is completely spherical for Al even if the equidensity surface level is lowered to 0.25 e A$^{-3}$. Because the average valence electron densities of Si and Al are 0.20 and 0.18 e A$^{-3}$, respectively, the above situation does not change when the charge densities are normalized using the average valence densities, i.e., that of Al increases by only about 10%.

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From this result, MEM/Rietveld analysis can clearly show the difference between covalent and metallic bonds.

In figure 3(a), there is a small charge density maximum at the bond midpoint, which is an artifact and originates from the insufficient d-spacing range of the experimental structure factors [33]. Although this maximum completely disappears using the d-spacing range \(d > 0.343\) [33], \(d > 0.76\) was used in figure 3 because a similar range (\(d > 0.80\)) was used in figure 4. The magnitude of this maximum was estimated to be less than 0.03 e Å\(^{-3}\) [32, 33].
and does not affect the qualitative discussion of figures 3 and 5(b).

Next, we discuss the results of icosahedral approximants containing an Al_{12} icosahedron without the central atom (1/1-AlMnSi and 1/1-AlReSi icosahedral approximants) and the Al_{12}Re icosahedron with a central atom (1/0-Al_{12}Re icosahedral approximant) [13, 34, 35]. Figure 4 shows the equidensity surfaces of the first-shell cluster of the 1/1-AlReSi and 1/0-Al_{12}Re approximants. Both compounds have the same Al_{12} icosahedron building block, although the Al_{12} cluster is slightly strained because of the cubic symmetry of the crystalline structure. The structural difference between the two structures arises from the occupation of the cluster center site. In the 1/1-AlReSi approximant, the center of the first-shell Al_{12} icosahedron is vacant, whereas the center is occupied with Re in the 1/0-Al_{12}Re approximant, as confirmed by Rietveld analysis. For the 1/1-AlReSi approximant, there is high electron density between Al atoms in the first-shell cluster and also between Al atoms in the first-shell cluster and Re atoms in the second-shell cluster at 0.35 \( e/\text{Å}^3 \) (figure 5(a)). The latter bonds are the very thick bonds directed towards the outside of the icosahedron along the five-fold direction in figure 4(a). This covalent bond is considered to be sp- (Al) and d-electron (Re) hybridization. On the other hand, 1/0-Al_{12}Re does not have such strong bonds between Al atoms in the first-shell cluster, and the electron density distribution is almost spherical at 0.35 \( e/\text{Å}^3 \). Because the average valence electron densities of 1/1-AlReSi and 1/0-Al_{12}Re are 0.24 and 0.20 \( e/\text{Å}^3 \), respectively, the above situation, whether a covalent bond exists between Al atoms in the icosahedron or not, does not change when the charge densities are normalized using the average valence densities, i.e., that of 1/1-AlReSi decreases by about 20%. From the above comparison between Al_{12} and Al_{12}Re icosahedra, the occupation of the central site with a Re atom induces the bonding conversion from more covalent to more metallic, which was called MCBC by Kimura et al [13, 15, 36].

It is mentioned that most features of the electron density obtained by \textit{ab initio} calculations of 1/1-AlMnSi [37] are in qualitative agreement with the experimental result [34], which was obtained from the same MEM/Rietveld analysis as that used for figures 4 and 5 [13]. It has also been confirmed that MCBC occurs for 13 atom (with a central atom and metallic) and 12 atom (without a central atom and covalent) icosahedral clusters of Al and B using semi-empirical [36] and \textit{ab initio} molecular orbital calculations [38, 39]. In the latter two cases of [38] and [39], not only the electron density but also its Laplacian and the electron localizability indicator, which were proposed to study the chemical bonding nature by Bader [40] and Kouhout [41, 42], respectively, were investigated. MCBC from the 13-atom metallic to the 12-atom covalent icosahedron is consistent with the packing fractions of 0.726 and 0.604, respectively. The former value is close to 0.744 for the close-packed structure and the latter value approaches 0.340 for the diamond structure.

Figure 5(a) shows the equidensity surfaces of the electron density for the first- and second-shell clusters of the 1/1-AlReSi approximant. It was found that the bond thicknesses, i.e., strengths, are distributed, as mentioned for figure 4(a) in the second previous paragraph. To determine the bond strength distribution, the charge (electron) densities at various bond midpoints are plotted in figure 5(b) [13], together with those for the 1/0-Al_{12}Re approximant, and pure Si and Al. The bond strength is widely distributed from strong covalent bonds for Si to weak metallic bonds for Al. The coexistence of covalent and metallic bonds and the wide distribution of bond strengths are characteristic features of structurally complex solids, as described in section 2. Furthermore, the bond strengths of the intra-cluster bonds are significantly stronger than those of the inter-cluster bonds, suggesting that this crystal is partly molecular-solid-like. For 1/0-Al_{12}Re, the bond strengths of Al–Re and Al–Al are much weaker than those for 1/1-AlReSi, which is consistent with the qualitative conclusion in the second previous paragraph.

The above situation, where the charge density at the bond midpoint is well-correlated with the inter-atomic distance from weak purely metallic to strong purely covalent bonds in Al, 1/0-Al_{12}Re, 1/1-AlReSi, and Si, is a simple case and is not necessarily general. For example, in the RuGa_{2}, which is a similar 13 group and transition metal elements compound, the highest electron density bond has a longer bond length than several of the other bonds [43].
Unfortunately, MEM analysis cannot be applied to quasicrystals. Instead, indirect evidence by comparing the experimental quasi-lattice constants (parameters) and the average atomic radii has been used to determine the covalent bonding nature in Al–Pd–Re quasicrystals [44]. According to this analysis, the covalent bonding nature increases with increasing transition metal concentration, because the electron density of the strong covalent bond between Al and the transition metal elements increases with increasing transition metal concentration.

Finally, we discuss the relationship between the bonding nature, electronic structure, and electronic transport properties. Although MCBC is clearly distinguishable from the metal–insulator transition, they are closely related. If the covalent bonds in the 1/1-AlReSi approximant are sufficiently strong, they should contribute to form a (narrow) band gap in the vicinity of $E_F$, such as that seen in the related RuAl$_2$ and RuGa$_2$ crystals [45, 46]. However, band structure calculations indicate that the 1/1-AlReSi approximant forms a deep pseudogap instead of a real gap, as shown in figure 6(a), which is consistent with a previous result [23] and table 3. The minimum density of states ($D(E)$) near $E_F$ is about 1/10 of the free electron value. The origin of the finite $D(E_F)$ is considered to be the weak inter-cluster bond, which is close in strength to the pure Al–Al bonds as shown in figure 5(b). On the other hand, although the 1/0-Al$_{12}$Re approximant has weak metallic bonds between the Al atoms of the first shell, the bonds between the Al atoms and the central Re atom are slightly stronger, which can be seen by the weak bonds in the equidensity surfaces $\varepsilon=0.30 \, e \, \text{Å}^{-3}$ in figure 4(b). The weak Al–Re bonds may cause formation of a shallow pseudogap at $E_F$, as shown in figure 6(b) and table 3. The minimum $D(E)$ near $E_F$ of the 1/0-Al$_{12}$Re approximant is about 1/2 of the free electron value and about five times larger than that of the 1/1-AlReSi approximant. From comparison of the 1/0-Al$_{12}$Re and 1/1-AlReSi approximants, a stronger covalent bonding nature results in a deeper pseudogap.

To investigate the electrical and thermal properties in detail, band structure calculations are a powerful tool and widely used to calculate the thermoelectric properties [47]. The DOS calculation can qualitatively explain the experimental electrical resistivity measurement results shown in figure 6(c). The 1/1-AlReSi approximant with a deep pseudogap exhibits a negative temperature coefficient of resistivity (TCR) with high electrical resistivity ($\rho_{300K}$) of 6500 $\mu\Omega$ cm at 300 K. The origin of the negative TCR is considered to be the localization tendency, which is usually stronger for smaller $D(E)$. On the other hand, the 1/0-Al$_{12}$Re approximant with a shallow pseudogap shows a positive TCR with a low value of $\rho_{300K}=26 \, \mu\Omega \, \text{cm}$ because of the large $D(E_F)$. Both the intra- and inter-cluster bond strengths in the 1/1-AlReSi approximant are significantly stronger than those in the 1/1-AlMnSi approximant [13, 34, 35]. Indeed, the 1/1-AlMnSi approximant forms a shallower pseudogap than the 1/1-AlReSi approximant from the band structure calculation by Takeuchi [48]. Thus, a relatively low value of $\rho_{300K}=3600 \, \mu\Omega \, \text{cm}$ and shelving TCR was observed in the 1/1-AlMnSi approximant. Among the examples shown in figure 6(c), the icosahedral Al–Pd–Re quasicrystal exhibits the highest room temperature electrical resistivity of $\rho_{300K}=15000 \, \mu\Omega \, \text{cm}$ and the most negative TCR, indicating that the Al–Pd–Re quasicrystal should have the deepest pseudogap because of the strong covalent bonds between the Al and Re atoms, as mentioned above.

It should be noted that some arc-melted and annealed Al–Pd–Re quasicrystals and 1/1-AlReSi approximants have porous microstructures. Although the absolute value of $\rho$ is affected by pores and cracks in the sample microstructure, the...
Table 3. Minimum electronic density of states near $E_F$ and calculated density of states at $E_F$ using the free electron model for 1/1-AlReSi and 1/0-Al$_2$Re approximants.

|               | $D(E_{\text{min}})$  | $D(E_F)$          |
|---------------|-----------------------|--------------------|
|               | (states/eV atom) near $E_F$ | (free electron model) (states/eV atom) |
| 1/1-AlReSi    | 0.04                  | 0.40               |
| 1/0-Al$_2$Re  | 0.20                  | 0.40               |

TCR is hardly affected by them. $S$ is also not greatly affected by such porous microstructures [49], but strongly depends on the local electronic structure near $E_F$. $S$ can be approximated by

$$S \propto \frac{1}{D(E_F)} \left[ \frac{dD(E)}{dE} \right]_{E=E_F}.$$  

(1)

From equation (1), a large $S$ is obtained when $D(E)$ is small and the gradient of $D(E)$ is large at $E_F$. Therefore, a deep pseudogap can increase $S$. Indeed, large $S$ values of 90–100 and 50–60 $\mu$V K$^{-1}$ were obtained for Al–Pd–Re quasicrystals [21] and 1/1-AlReSi approximants [23], respectively. On the other hand, the metallic 1/0-Al$_2$Re approximant with a shallow pseudogap has an $S$ value of 2 $\mu$V K$^{-1}$ [50]. Among these materials, the Al–Pd–Re quasicrystal with strong covalency should form the deepest pseudogap, which is suitable for obtaining a large $S$. Details of the effect of improving the microstructure on the thermoelectric properties is described in section 4.

Although the electronic transport properties are well explained by the electronic structure near $E_F$, it is not clear how to control it. Controlling the carrier concentration is easy and frequently effective to control the bulk thermoelectric properties when the rigid band scheme is applicable [23, 51]. Because the bonding nature greatly affects the pseudogap formation near $E_F$, which has a large influence on the bulk thermoelectric properties, it can be an easily controllable factor, when the rigid band scheme is not applicable. We have proposed a new guiding principle to increase the thermoelectric performance by controlling the bonding nature, and it will be explained in section 4.

### 4. Thermoelectric properties of Al–Pd–(Re, Mn) icosahedral quasicrystals and a guiding principle to increase ZT: WBRHCs

As described in section 2, Al-based ICSs have several advantages for thermoelectric materials. In particular, Al–Pd–(Re, Mn) icosahedral quasicrystals are the most promising candidates because they have the strongest bonds, the deepest pseudogap, and the largest $S$, as discussed in section 3. Furthermore, as described in section 3, 1/1-AlMnSi and 1/1-AlReSi approximants have a continuous range of bond strengths from covalent to metallic. Overall, the bond strengths of intra-cluster bonds are stronger than those of inter-cluster bonds. This mean that these approximants are considered to be an intermediate state among the three typical solids: metals, covalently bonded networks (semiconductors), and molecular solids, as shown in figures 7(a) and (b). This unique situation is considered to be the same for Al–Pd–(Re, Mn) icosahedral quasicrystals because they have similar icosahedral cluster building blocks to Mackay icosahedral (MI) clusters.

To further improve the thermoelectric performance of the Al–Pd–(Re, Mn) quasicrystals, we have proposed a guiding principle for increasing $ZT$: WBRHCs [27]. To increase $ZT$, it is essential to simultaneously optimize $\sigma$, $S$, and the electron and phonon thermal conductivities ($\kappa_e$ and $\kappa_p$). Although there are several suggested guides for increasing $ZT$ represented by PGEcs [28], WBRHCs are different from the other cases and was also introduced in the News and Views of Nature Materials by Thiel [52]. In PGEcs, phonon transfer is difficult (like in glass) but electron transfer is easy (like in crystals), such as in clathrates and filled-skutterudites, as described in section 2. Figure 7(b) shows a schematic diagram of the WBRHCs scheme with varying intra- and inter-MI-cluster bonds strengths ($V_i$ and $V_c$, respectively). At the limit of the molecular solid, $V_i$ and $V_c$ determine the band interval and band width, respectively. The effective mass ($m^*$) decreases with increasing band width, i.e., $V_i$. On the other hand, $m^*$ decreases with decreasing $V_c$ because of the small $m^*$ of typical (nearly free electron) metals. Increasing $V_i$ and decreasing $V_c$ (WBRHCs, the direction of thick black arrow in figure 7(b)) increases $m^*$ and enhances the thermoelectric performance (i.e., increases $ZT$) as follows. For the nearly free electron model with $m^*$, the power factor

$$S^2 \sigma = \frac{n^{2/3}}{3^{5/3} \pi^2} \frac{k_B^4}{\hbar^2} T^2 \left( \frac{m^* \tau}{n^{1/3}} \right),$$  

(2)

which is the numerator of $ZT$, is simply proportional to $m^*$ and $\tau$, and inversely proportional to $n^{1/3}$. $\kappa_p$ can be expressed as

$$\kappa_p = \frac{1}{3} C a^2 K \tau_p,$$  

(3)

where $C$, $a$, $K$, $M$, and $\tau_p$ are the specific heat, the distance between clusters, the force constant of bond between clusters, the mass of a cluster, and the relaxation time of a phonon, respectively. In the WBRHCs scheme, $m^*$ increases by weakening the inter-cluster bonds and strengthening the intra-cluster bonds, and thus $S^2 \sigma$ should increase. Simultaneously, $K$, i.e. $\kappa_p$, decreases by weakening of the inter-cluster bonds. By both increasing $S^2 \sigma$ and decreasing $\kappa_p$, $ZT$ should increase through weakening of the inter-cluster bonds and strengthening of the intra-cluster bonds. Additionally, $M$ increases through heavy element substitution, and $\tau_p$ can be decreased by an alloying effect. Heavy metal substitution and alloying effect also decrease $\kappa_p$ and increases $ZT$.

From $m^*$ analysis using the two-band model of the ternary Al–Pd–Re and quaternary Al–Pd–Re–Ru quasicrystals, we confirmed that the WBRHCs was effective [53].
Figure 8 shows the temperature dependences of $\sigma$ and $S$ for $\text{Al}_{71}\text{Pd}_{20}(\text{Re}_{1-x}\text{Ru}_x)_9$. The solid lines are fitted curves obtained using the two-band model. From the bond strength analysis of the isostructural 1/1-$\text{AlMnSi}$ and 1/1-$\text{AlReSi}$ approximants described in section 3, the intra-cluster bonds are stronger than inter-cluster bonds (i.e., $V_i > V_o$), and both the intra- and inter-cluster bonds of the 1/1-$\text{AlReSi}$ approximant are stronger than those of the 1/1-$\text{AlMnSi}$ approximant [13, 34, 35]. This indicates that the bond strengths of 5d transition metal elements are higher than those of 4d transition metal elements, and that the bond strengths of 4d transition metal elements are higher than those of 3d transition metal elements, as shown in figure 9.

(i) In the ternary Al–Pd–Re quasicrystal, transition metals are considered to first occupy the intra-cluster transition metal sites and then the inter-cluster site, because the former sites have stronger bonds, like in figure 5(b). Accordingly, $m^*$ (i.e., $ZT_{\text{max}}$) has been experimentally confirmed to increase with increasing $V_i$ (A → B in figures 7(b), (c) and (e)), where the number of transition metal atoms in the intra-cluster sites increases, and then to decrease with increasing $V_o$ (B → C in figures 7(b), (c) and (e)), where the number of transition metal atoms in the inter-cluster sites increases.

(ii) In the quaternary Al–Pd–Re–Ru system, starting from the highest $ZT_{\text{max}}$ composition of the ternary system (B in figure 7), Ru is considered to first substitute at weak inter-cluster Re sites with low Ru concentration and decrease the bond strength $V_o$ (B → D in figure 7(b)).
Figure 9. Transition metal elements that are candidates for substitution in the Al-Pd–Re quasicrystal. The bond strength will decrease with decreasing atomic number in congener. Route (1) is Ru and Fe substitution in the Re site, and route (2) is full replacement of Mn in Re sites.

because Ru forms a weaker bond than Re. At high Ru concentration, after the inter-cluster sites are fully occupied by Ru, Ru substitutes at intra-cluster Re sites, and $V_1$ decreases ($D \rightarrow E$ in figure 7(b)). In this case, $m^*$ has been experimentally confirmed to first increase and then decrease with increasing Ru concentration ($B \rightarrow D \rightarrow E$ in figures 7(d) and (f)).

(iii) The higher $m^*$ than that of D should be obtained in the direction of decreasing $V_1$, and increasing $V_1$ (the thick black arrow in figure 7(b)).

According to the WBRHCs scheme, we performed two more substitutions, as shown in figure 9. One was Fe substitution following Ru substitution for Re, and the other was full replacement of Mn for Re. From the composition optimization for high $ZT$ with Al–Pd–Re quasicrystals as a starting material [21], we selected Al$_{17}$Pd$_{30}$Re$_9$ as the composition (B in figure 7). In the Al–Pd–Re quasicrystals, $ZT$ greatly increased from less than 0.01 to 0.11 by composition optimization. However, one remaining issue is that $\sigma$ is difficult to reproduce because the sample contains a lot of pores and cracks. The effects of pores and cracks depend on the measurement sample area, as will be discussed later. Through Fe substitution for Re in the Al$_{17}$Pd$_{30}$Re$_9$ quasicrystal [54], it is expected that the inter-cluster bond strength decreases, which then decreases the intra-cluster bond strength at high Fe concentration, similar to the Ru case. In these cases, $\sigma$ and $S$ significantly increased at medium fourth-element concentrations, that is, substitution of $x = 0.55$ Ru and $x = 0.65$ Fe. Thus, $S^2\sigma$ significantly increased from an increase in $m^*$. Substitution with a fourth element decreased the phonon thermal conductivity because of the decrease of both $K$ by weakening the inter-cluster bonds and $\tau_p$ by an alloying effect. As a result, maximum $ZT$ values of 0.15 and 0.22 were obtained for optimized Al$_{17}$Pd$_{30}$(Re$_{1-x}$Ru$_x$)$_9$ ($x = 0.55$) [53] and Al$_{17}$Pd$_{30}$(Re$_{1-x}$Fe$_x$)$_9$ ($x = 0.65$) [54], respectively, as shown in figures 10(a) and (b).

Before discussing the effect of Mn replacement for Re, we will briefly mention the improvement of the porous microstructure in Al–Pd–Re quasicrystals. After arc-melting and annealing the Al–Pd–Re quasicrystal, a large number of pores and cracks are observed, as shown in figure 11(a). In this case, the relative density is less than 70%. These extrinsic factors result in poor electrical properties and anisotropy, which is often disadvantageous for thermoelectric materials in terms of the difficulty in controlling the microstructure and brittleness. Therefore, various $ZT$ values have been reported for the same composition of the Al–Pd–Re quasicrystal [21, 53, 54], as shown for $x = 0$ in figures 10(a) and (b). To avoid the influence of the extrinsic factors on the electrical conductivity, we used the spark plasma sintering (SPS) method to obtain samples with no cracks and fewer pores as shown in figure 11(b) [49]. After successfully synthesizing a nearly 100% relative density sample, we found that both the electrical and thermal conductivities increased, but the pores and cracks did not have a large influence on $S$ [49]. As a result of increasing $\sigma$ by increasing the electron mobility, a $ZT$ value of 0.12 was obtained for the sintered Al$_{17}$Pd$_{30}$Re$_9$ quasicrystal as shown in figure 10(c). Al–Pd–Mn quasicrystals do not contain a large number of pores, but they do contain some cracks. The relative density of Al–Pd–Mn quasicrystals is typically greater than 95%, indicating that it is not necessary to improve the microstructure.

Figure 10(c) shows the temperature dependence of $ZT$ for Al$_{17}$Pd$_{30}$Mn$_9$ and sintered Al$_{17}$Pd$_{30}$Re$_9$ quasicrystals [55]. Note that, the composition of Al$_{17}$Pd$_{30}$Mn$_9$ shows the highest $S^2\sigma$, which is the same situation as in the Al–Pd–Re system.
The Al$_{71}$Pd$_{20}$Mn$_9$ quasicrystal exhibited a relatively high ZT value of 0.16 at 373 K, which is much higher than that of the different composition Al$_{70.8}$Pd$_{20.9}$Mn$_{8.3}$ (0.08 at room temperature) [20]. Similarly, in Al–Pd–Re quasicrystals, the thermoelectric properties are sensitive to the sample’s composition, which can be understood by the average valence electron number per atom (e/\(a\)) [49]. This is mainly because of a deep pseudogap near \(E_F\) in both Al–Pd–Re and Al–Pd–Mn quasicrystals. The Al$_{71}$Pd$_{20}$Mn$_9$ quasicrystal shows a higher ZT value than that of the Al$_{71}$Pd$_{20}$Re$_9$ quasicrystal mainly because of its high \(\sigma\). The differences in \(\sigma\) and \(e/\(a\)) between the Al–Pd–Mn and Al–Ga–Pd–Mn (4 at.% Ga substituted for Al) systems. However, there is no significant difference in the DOS (figure 13). On the other hand, \(\kappa\) (especially \(\kappa_p\)) benefically decreased by Ga substitution, as shown in figure 12(b) and table 4, by the combination of weakening of the inter-cluster bonds and an alloying effect (phonon scattering by the disorder lattice induced by the alloying) [55]. \(\kappa_p\) near room temperature was at the level of the theoretical minimum thermal conductivity (\(\kappa_{\text{min}}\)). \(\kappa_p\) was obtained by subtracting the electron thermal conductivity (\(\kappa_e\)) determined using the Wiedemann–Franz law from \(\kappa_e\).

To confirm the weakening of the inter-cluster bonds by Ga substitution for Al, we performed a speed of sound measurement because MEM/Rietveld analysis cannot be directly applied for quasicrystals. The average speed of sound (\(v_s\)), the Debye temperature (\(\theta_D\)), the relative change in the relaxation time of the phonon (\(\Delta\tau_{\text{ph}}/\tau_{\text{ph}}\)), and \(\kappa_p\) for Al$_{71-x}$Ga$_x$Pd$_{20}$Mn$_9$ quasicrystals (\(x = 0, 2, 3\)) are listed in table 4 [55]. The \(v_s\) and \(\theta_D\) values of the samples with \(x = 0\) and 2 are almost identical, but those of the sample with \(x = 3\) significantly decrease, indicating that the force constant (\(K\)) should be lower for \(x = 3\) than for \(x = 0\) and 2. \(v_s\) can be expressed as

\[
\frac{1}{v_s} = \sqrt{\frac{\alpha^2 k}{M}}.
\]  

(4)

To confirm the decrease of \(K\) by the Ga substitution, we substituted the parameters \(v_s\) (or \(\theta_D\), \(-7.6\%\)), \(a\) (\(+0.1\%\)), and \(M\) (\(+2.8\%\)) for the \(x = 3\) sample into equation (4) and obtained \(-15\%\) of \(K\), which is in good agreement with the WBRHCs picture. The experimental value of \(\kappa_p\) decreased by more than 50% after Ga substitution. Therefore, from equation (3), \(\Delta\tau_{\text{ph}}\) should decrease by about 35% through the alloying effect.

Finally, we briefly mention the recent progress in Ga-substituted Al$_{70}$Pd$_{21}$Re$_9$ quasicrystals. Although very small amounts of the Al$_{11}$Re$_4$ and Al$_{12}$Pd$_2$ binary phases are observed in sintered samples, \(\sigma\) and \(S\) increase with Ga substitution [60], which is a different trend to Al–Ga–Pd–Mn quasicrystals. Ga substitution leads to a decrease in the phonon thermal conductivity, and results in ZT increasing from 0.12 to 0.18 [60]. However, this value does not exceed the maximum value of 0.26 for the Al$_{70}$Ga$_2$Pd$_{20}$Mn$_9$ quasicrystal.

The above experimental results demonstrate that our established guiding principle (WBRHCs) provides a route for increasing ZT in cluster-based complex structure solids, especially by weakening the inter-cluster bonds. However, this principle does not yet offer the upper ZT limit of a target material. As frequently discussed in the thermoelectric...
society, it is essential to have a suitable electronic structure, such as a narrow band gap, to obtain a large power factor. Since quasicrystals have complex crystal structures, the phonon thermal conductivity is less than 1 W/(m K) at room temperature, which is almost the theoretical minimum thermal conductivity. Therefore, to obtain a high-ZT quasicrystal material, the materials should possess a real gap rather than a pseudogap; that is, it should be a semiconductor quasicrystal. We are now furthering the search for semiconductor quasicrystal based on theoretical and experimental studies [61].

Acknowledgement

The authors acknowledge the collaborators Prof. M Sakata at Nagoya University, Prof. M Takata at the RIKEN SPring-8...
Center, Prof. E Nishibori at Tsukuba University, and Dr K Kirihara in our laboratory for MEM/Rietveld analysis. We also thank Dr K Kirihara, Dr T Nagata, Dr J T Okada, Dr S Hosoi, Mr T Hamamatsu, and Mr T Kamimura in our laboratory for sample preparation, thermoelectric properties measurement, and their analysis.

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