Thermopower of correlated semiconductors: application to FeAs$_2$ and FeSb$_2$

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We investigate the effect of electronic correlations onto the thermoelectricity of semi-conductors and insulators. Appealing to model considerations, we study various many-body renormalizations that enter the thermolectric response. We find that, contrary to the case of correlated metals, correlation effects do not 

We use this frame-work to make general arguments on how to obtain high values for the figure of merit $ZT$. Given the sizes of the charge gaps, effective masses, and other parameters, these considerations allow us to put constraints on the possible regimes of materials of interest. In Section IV we apply realistic electronic structure tools, and show that for iron arsenide, FeAs$_2$, both the electronic structure, as well as the thermolectric response can be understood as that of a conventional semiconductor and described quantitatively by ab initio band-structure methods. For the iron antimonide on the other hand, standard density functional theory (DFT) based methods are known to be insufficient to account for the electronic structure. To show that correlation effects play an important role, we employ a hybrid functional approach, as well as Hedin's GW approximation, with the latter yielding results in good agreement with the experimental charge gap. As to the thermolectric response, however, we find qualitative agreement with experiment only at intermediate temperatures (35-70K). Our analysis shows that the low temperature Seebeck coefficient of FeSb$_2$ is incompatible with a local electronic picture, suggesting the importance of vertex corrections and non-local self-energy effects (that we neglect), or the presence of a substantial phonon drag effect, as is e.g. found in the classical example of p-type Germanium.

II. THE MATERIALS

Despite the structural similarity of FeSb$_2$, and FeAs$_2$, experimental findings point to markedly different properties, heralding a varying importance of correlation, and, potentially, electron-phonon effects. FeAs$_2$ is an insulator with a gap of 0.2-0.22 eV.
as obtained from the activation behavior in the resistivity at temperatures of 200K and higher. Below 200K, the influence of impurities is pivotal: the resistivity has a metallic slope before resuming, below 30K and down to 10K, an activation law with an energy of 0.01eV. Further, below 10K, the resistivity exhibits activation with 6K (0.5meV)3. The Hall coefficient is negative for all temperatures. Congruently, the Seebeck coefficient is negative as well: From its room temperature value \(-200\mu V/K\) it grows in magnitude upon cooling, to reach \(-7mV/K\) at 12K but it vanishes towards zero temperature.4

In the case of FeSb2, optical spectroscopy finds a small gap of 432K (37meV) at low temperatures but witnesses the development of a Drude-like peak at 70K and above. The concomitant transfer of spectral weight is found to extend over an energy range as high as 1eV, i.e. a scale that is much larger than the initial gap, a common harbinger of correlation effects.5 The resistivity of FeSb2, on the other hand, has three distinct temperature regimes that exhibit activated behavior: In the range of 50–100K the activation energy corresponds to a gap of 300K (26meV)6,7. From 20K down to 10K, the resistivity shows a shoulder-like behavior with an activation energy of \(\Delta/2 = 3meV\), while below 5K, extrinsic impurities are believed to be at the origin of a weakly temperature dependent resistivity following an activation behavior with \(0.04-0.09meV\).8 Further, below 10K, the resistivity has a metallic slope before resuming, below 30K and down to 10K, an activation law with an energy of 0.01eV.9

The system with electrons, e.g. FeSb2,10, whereas the susceptibility of FeAs2 is flat up to 350K11. That the low temperature feature in the specific heat of FeSb2 has no spin signature might indicate that its contribution to the entropy is associated with either the charge degrees of freedom or an electron-phonon effect. The importance of electron-electron effects in FeSb2 is further highlighted by the fact that various properties are very sensitive with respect to changes in the carrier density. Doping the system with electrons, e.g. FeSb2–Sn,12 or holes, e.g. FeSb2–Te,13,14, instantly metalizes the compound, sometimes generates a Curie law at low temperatures15 and reduces the Seebeck coefficient16. Thermoelectric properties of FeSb2–Te are indeed that of a correlated metal, i.e. the low temperature Seebeck coefficient is linear in T, with an enhancement factor of 15 via the effective mass.17

Despite these indications for correlation effects, some experimental findings for the antimonide are quantitatively reproducible by conventional band-structure methods: Volume and bulk-modulus are very well captured within the generalized gradient approximation (GGA) of DFT. Also, the finding of small electron and hole pockets in FeSb2–Te (see also below) within band-structure methods could simply be attributed to the well documented underestimation of charge gaps within DFT. Moreover, a calculation within the local density approximation (LDA) with a Hartree-like Coulomb interaction (LDA+U) suggested that FeSb2, while being paramagnetic, could be close to a ferromagnetic instability.18 Weak ferromagnetism was then indeed found in Fe1–xCo2xSb2.19

III. TRANSPORT FORMULAE, GENERAL & MODEL CONSIDERATIONS

Within the Kubo formalism the Seebeck coefficient – that relates the gradients of temperature and electrical field – is given by (see e.g. Ref. 10)

\[
S = -\frac{k_B A_1}{|e| A_0} \frac{\partial^2 S}{\partial T^2} (\mu) \Xi(\omega) \tag{1}
\]

where the current-current (current-heat current) correlation function \(A_0 \, A_1\) is given by

\[
A_n = \int d\omega \beta^n (\omega - \mu)^n \frac{\partial f_\mu}{\partial \omega} \Xi(\omega) \tag{2}
\]

Here, \(f_\mu\) is the Fermi function, \(\mu\) is the Fermi level, and \(\Xi\) is the transport kernel. If vertex corrections are neglected, the transport kernel can be expressed (in matrix notation) as

\[
\Xi(\omega) = \sum_k \text{Tr} [\psi(k) A(k, \omega) \psi(k) A(k, \omega)] \tag{3}
\]

with the Fermi velocity \(v_{ij}(k) = \frac{\hbar}{m} \langle \psi_{ki} | \nabla | \psi_{kj} \rangle\), and the spectral function \(A_{ij}(k, \omega)\), where \(\psi_{ki}\) is a complete set of one electron basis functions, such as Kohn-Sham
orbits. Using transport coefficients Eq. [2], we can further express (see e.g. Ref. [6]) the dc conductivity, the thermal conductivity, and the figure of merit as

\[ \sigma = \frac{2\pi e^2}{h V} A_0 \]  
\[ \kappa = \kappa_L + \frac{2\pi k_B T}{h V} \left( A_2 - \frac{A_1^2}{A_0} \right) \]  
\[ ZT = \frac{S^2 \sigma T}{\kappa} \]

where \( V \) is the unit-cell volume, and \( \kappa_L \) the thermal lattice conductivity.

The chemical potential, \( \mu \), is obtained by the requirement of the charge neutrality,

\[ n - p - n_{D^+} = 0 \]  
where

\[ n \]
\[ p \]
\[ = \sum_k \int_{-\infty}^{\infty} d\omega \left\{ A^c(k,\omega) f_p(\omega) - A^v(k,\omega) [1 - f_p(\omega)] \right\} \]

is the number of electrons (holes). Here, \( A^{c,v}(k,\omega) \) is the valence/conduction spectral function. We also allowed for the presence of ionized donor impurities \( n_{D^+} = n_D \left[ 1 + 2e^{-\beta(E_D - \mu)} \right]^{-1} \) of concentration \( n_D \) at an energy \( E_D \).

### A. model considerations

Here we find it instructive to extend on the usual textbook considerations (see e.g. Ref. [36]) and generalize to include carrier dependent masses, renormalizations, as well as finite (yet energy-independent) scattering amplitudes. This makes it possible to investigate the important effects of particle-hole asymmetry, carrier coherence, and allows us later to discuss the consistency of a purely diffusive thermopower for a given material.

Assuming a Lorentzian line shape of the conduction (c) and valence (v) spectral functions

\[ A^{c,v}_\text{coh}(k,\omega) = \frac{Z^{c,v}_k}{\pi} \frac{\Gamma^{c,v}_k}{(\omega - \xi^{c,v}_k)^2 + (\Gamma^{c,v}_k)^2} \]

i.e. we limit the discussion to the coherent part, \( A^{\text{coh}} \), (of weight \( Z_k \)) of the full spectrum, \( A = A^{\text{coh}} + A^{\text{incoh}} \), with a quasi-particle dispersion \( \xi_k \), and an elastic scattering of amplitude \( \Gamma_k \). Within this approximation, one finds for the number of electrons \( n \)

\[ n = \sum_k \left\{ \frac{1}{2} - \frac{1}{\pi} Im\psi \left[ \frac{1}{2} + \frac{2\beta}{2\pi} \left( \left( \Gamma_k + \imath \xi_k - \mu \right) m_0 \right) \right] \right\} \]

where \( \psi(z) \) is the digamma function, and \( \beta = 1/(k_B T) \) the inverse temperature. The above expression reduces to the usual \( \sum_k f(\mu - \xi_k) \) in the coherent limit \( (\Gamma = 0) \).

The response functions, Eq. [2], can be also be expressed analytically (here, we restrict ourselves to \( n = 0, 1 \) as

\[ A_n = \frac{\beta^{n+1}}{4\pi^3} \sum_k \frac{Z^2_k}{\Gamma_k \xi_k^2} \left( \xi_k - \mu \right)^n \text{Re}(\psi(x)) \]

\[ -\frac{\beta}{2\pi} \left( \xi_k - \mu \right)^n \Gamma_k \text{Re}(\psi''(x)) + n\Gamma_k^2 \text{Im}(\psi''(x)) \right\} \]

where the arguments of the derivatives of the digamma function are

\[ x = \left[ \frac{1}{2} + \frac{\beta}{2\pi} \left( \Gamma_k - \imath (\xi_k - \mu) \right) \right] \]

The first contribution in the curly brackets of Eq. [11] is the leading term in the coherent limit \( (\Gamma \to 0) \). Indeed with \( \beta/(2\pi^2) \text{Re}(\psi''(1/2 + i\beta x/(2\pi)) = -f'(x) \) one recovers the Boltzmann expressions for transport coefficients \( (\beta \to 0) \) in which appears an \textit{ad hoc} lifetime \( \tau = 1/(2\Gamma) \). In the above expression, however, the influence of finite scatterings is not restricted to the pre-factor, but since the spectrum broadens, a wider energy range becomes activated for supplying charge carriers, and, as a result, the Fermi statistics assumes the digamma form, and, also, higher order terms appear. Therewith, contrary to the Boltzmann description at small \( \Gamma \), coherence effects do not in general cancel in the ratio \( A_1/A_0 \) of the Seebeck coefficient.

For illustrative purposes, we now analyze Eq. [11] in terms of special cases for simple quadratic dispersions : We consider bare bands \( E^c,v = \frac{\hbar^2 k^2}{2m^{c,v}_0} \), and an interacting dispersion

\[ \xi^{c,v}_k = \pm \Delta/2 \pm \frac{\hbar^2 k^2}{2m^{c,v}_0} \]

where \( \Delta \) is the charge gap, \( m^* \) the effective mass of the carriers, and the origin of the chemical potential is chosen at the mid-gap point. Further, we assume the Fermi velocities to be given by the group velocity \( v_k \)

\[ v_k = \frac{1}{\hbar} \partial_k E_k \]

and weights, \( Z \), and scattering rates, \( \Gamma \), that are independent of momentum.

#### 1. the large gap coherent semi-conductor

In the limit of a coherent system \( (\Gamma \ll 1) \) with a large gap \( (\beta \Delta/2 - \mu \gg 1) \), Eq. [11] can be simplified to

\[ A_n^{c,v} = (\pm 1)^n \frac{3\lambda}{\sqrt{2\pi^3\beta}} e^{-\beta \Delta/2} \left\{ (\beta(\pm \mu - \Delta/2))^n - \frac{5}{2} \right\} \]

where all carrier specific parameters have been gathered in

\[ \lambda^{c,v} = \frac{Z^2 m^{5/2}}{\Gamma m^0} e^{\pm \beta \mu} \]
Therewith the Seebeck coefficient becomes

\[ S = \frac{k_B}{|e|} \left( \frac{1}{T} \left( \mu - \frac{\Delta}{2} \delta \lambda \right) - \frac{5 k_B}{2 |e|} \delta \lambda \right) \]

(16)

\[ = \frac{1}{|e| T} \left( \mu - \frac{\Delta}{2} \delta \lambda \right) - \frac{5 k_B}{2 |e|} \delta \lambda \]

(17)

where the asymmetry parameter \( \delta \lambda \) (that depends on \( \mu \) and \( T \)) is given by

\[ \delta \lambda = \frac{\lambda_c - \lambda_v}{\lambda_c + \lambda_v} \]

(18)

Hence, a large Seebeck coefficient can be achieved by an interplay of the gap, \( \Delta \), the anisotropy or asymmetry \( \delta \lambda \) in the transport function – stemming from either the densities of states \( (n_{c,v}^{0}) \), different bandwidth narrowings \( (m_{c,v}) \), scattering amplitudes \( (\Gamma_{c,v}) \) or quasiparticle weights \( (Z_{c,v}) \).

2. Upper limit for thermopower in a semiconductor

The position of the chemical potential plays an important role in maximizing the thermopower of a semiconductor. Let us take the example of two equivalent bands. In this case, the term in brackets in Eq. (17) becomes

\[ \mu - \frac{\Delta}{2} \tanh(\beta \mu) \]

(19)

This coefficient of the 1/\( T \) behavior is displayed in Fig. 1(a) as a function of the chemical potential for a few different temperatures. At high temperatures the optimal chemical potential, which maximizes thermopower, is near the gap edges. At low temperature \( k_B T < \Delta / 2 \), the 1/\( T \) coefficient shows a local extremum. The Seebeck coefficient vanishes at the point of particle-hole symmetry (\( \mu = 0 \) in the symmetric case considered here), and the optimal location of the chemical potential at low temperature is in the direct vicinity of that point, hence very close to the center of the gap.

Fig. 1(b) displays the value of the 1/\( T \) coefficient in this extremum and the value at the gap edge as a function of temperature. For \( k_B T \gtrsim 0.3 \Delta \) the maximum value of the thermopower is achieved when the chemical potential is at the gap edge, and for lower temperature, it is achieved close to the middle of the gap, where the thermopower can reach the maximum value of \( S = \Delta / (2eT) \).

In an asymmetric case, the thermopower can be larger than this maximum value, however, for a given charge gap \( \Delta \), there is always an upper bound for the Seebeck coefficient, namely

\[ |S(T)e| \leq \frac{\Delta}{T} + \frac{5}{2k_B}. \]

(20)

This is because the asymmetry is bounded to an absolute value of one \( |\delta \lambda| \leq 1 \). This extremal value corresponds to the fictitious system in which only one type of the two carriers contributes to the thermoelectricity, e.g. the conduction electrons, and with the chemical potential being, in that case, at the edge of the valence band.

Thus the correlation effects, such as small \( Z \) in the conduction band and large \( Z \) in the valence band, can enhance the thermopower of a semiconductor. However, this effect is limited by the form of Eq. (11), allowing maximum \( S \) bounded by Eq. (20). The possible merits of electron-hole-asymmetry for the case of metals is discussed in Ref. 11.

3. Model semi-conductor in the presence of donor impurities

With the goal of understanding the thermopower and the figure of merit in a renormalized semiconductor in a very general setup, including the presence of impurities, we now numerically study the model based on the response functions Eq. (11). As before, we assume...
parabolic dispersions Eq. (12), with the band structure depicted in the inset of Fig. 2(b): excitations of different effective masses are separated in energy by a gap $\Delta$, and we allow for the presence of donor impurities, situated at an energy $E_D$, as measured from the middle of the gap. We again assume transition matrix elements to be given by the group velocity, Eq. (13).

We choose the parameters compatible with the band structure of FeAs: we consider a gap $\Delta = 0.2\text{eV}$, and, unless stated otherwise, an impurity level at $E_D = 95\text{meV}$, as inferred from the low temperature activation behavior of the resistivity. In our current treatment, we assume that the impurity carriers have vanishing Fermi velocities, and thus their only effect is to shift the chemical potential.

To fix the particle–hole asymmetry, we note that at high temperatures, the number of ionized impurities are irrelevant with respect to the number of conduction and valence carriers, and the chemical potential follows the intrinsic behavior. In the coherent limit of the large gap semiconductor (see above) one finds that

$$\mu = 3kBT/4\ln(\eta_c/\eta_v), \quad \text{with } \eta = m^*/m_0. \quad (21)$$

In this regime the resistivity shows an activation law with the activation gap $\Delta/2$. Of course, the situation in a real material can be much more complicated (several types of impurities, temperature dependence of the gap, etc.). Using our ab initio data for FeAs2 (that is presented below in Section IV and Section V A), and assuming the approximate validity of Eq. (21) for non-parabolic dispersions, one finds the ratio of the valence and conduction effective mass $\eta_c/\eta_v = 2.5$ for FeAs2, which we will use for all the following model calculations.

Further, we note that a uniform weight-factor $Z$ cancels in the Seebeck coefficient, whereas in the figure of merit it can be seen as a scaling factor of the thermal lattice conductivity, for which we assume $\kappa_L/Z^2 = 250\text{W/(Km)}^{-1}$\text{cm}^3. Moreover, we use a unit-cell volume of 80\text{Å}^3.

Having thus fixed the size of the gap and the asymmetry, the principle parameters to vary in this setup are the concentration of impurities $n_D$ and the scattering rate $\Gamma$ (that we assume to be orbital independent). We will also study the dependence on the position of the impurity band $E_D$ away from the value motivated by the experimental resistivity.

a. \textit{Impurity concentration.} With these parameters, we display in Fig. 2(a) the temperature dependence of the chemical potential, the Seebeck coefficient, and the figure of merit for various impurity concentrations $n_D$, and for a constant scattering rate $\Gamma = 5\mu\text{eV}$.

In the intrinsic case, $n_D = 0$, the chemical potential is indeed linear above a certain temperature that is related to the scattering rate $\Gamma$. Below this regime, the chemical potential is almost temperature independent. Since $\Gamma$ is small, the point of charge neutrality at zero temperature is very close to the midgap point (where it is in the coherent case). The point of particle-hole compensation for the thermopower, however, is higher in energy than the midgap point, as rationalized, above, for the symmetric semiconductor. As a consequence, the chemical potential is actually moving through the symmetry point of the Seebeck coefficient, and we expect it to change sign as a function of temperature. This is clearly seen in Fig. 2(a).

At finite dopings, and at low temperatures, the donor level plays the role of the valence band, and the chemical potential (top panel) is between $E_D$ and the conduction band. In a coherent semiconductor the chemical potential is $\mu \approx (\Delta/2 - \delta/2) - k_BT\ln(n/n_D^*)$, where $\delta = \Delta/2 - E_D$ is the impurity activation energy, seen in the resistivity of this regime ($\rho \sim e^{-\beta\delta/2}$). At very low temperatures, $\mu$ is pinned to $E_D$. At intermediate temperatures the chemical potential smoothly connects with the intrinsic high temperature slope, as shown in Fig. 2(a). For large enough concentration of impurities ($n_D > 10^{16}$/cm$^3$), the chemical potential can even inside the conduction band at some intermediate temperature, which can result in a shoulder, or even a metallic slope in the resistivity (cf. the mentioned transport measurements on FeSb2, and FeAs2).

For finite impurity concentration ($n_D > 0$), the Seebeck coefficient displayed in Fig. 2(a) may or may not be enhanced at a given temperature, depending on whether or not the additional carriers bring the chemical potential closer to its optimal value. As explained above for the case of two particle hole symmetric bands, at fixed temperature there exists a value of the chemical potential, which maximizes the Seebeck coefficient. In our asymmetric setup with $\eta_c/\eta_v > 1$, the optimum chemical potential is located above the mid gap point. In the limit of vanishing impurity concentration, the midgap remains the point of charge neutrality at zero temperature, while the optimal chemical potential is very near that point, hence the Seebeck coefficient is a very strong function of temperature in this limit, and can even change sign, as seen in the middle panel of Fig. 2(a). We note however, that the fundamental extremum, established by the size of the gap, is always respected as is evident in Fig. 2(a).

The efficiency of the thermoelectrical material is determined by its figure of merit $ZT$, which we also plot in the lower panel of Fig. 2(a). $ZT$ can be greatly enhanced by the presence of impurities, and its maximum is not necessarily in close vicinity of the thermopower maximum. Indeed the largest $ZT$ for the current parameters is achieved at about $T = 220\text{K}$ for a concentration $n_D = 2 \cdot 10^{16}$/cm$^3$. The Seebeck coefficient at this point is actually smaller than in the intrinsic limit. The position of the impurity with respect to the gap edge has a large effect on the optimal impurity density. While for our specific choice of parameters, $\Delta/2 - E_D = 5\text{meV}$, the optimal density is $n_D = 2 \cdot 10^{16}$/cm$^3$, we notice that for larger gaps and/or larger separation of the impurity level from the gap edge, the optimal impurity density can reach a value as large as $10^{20}$/cm$^3$. To elucidate the origin of an optimal density further, we show in Fig. 2(b)
FIG. 2: Thermoelectric properties of the model semi-conductor with parameters inspired from FeAs$_2$: (a) from top to bottom: chemical potential, Seebeck coefficient, and figure of merit $ZT$ as a function of temperature for different impurity concentrations. (b) ingredients to the figure of merit $ZT$ as a function of temperature for different impurity concentrations. The inset shows the general setup of asymmetric valence and conduction dispersions, with an impurity level $E_D$ in the gap $\Delta$. (c) effect of the scattering amplitude $\Gamma$ (top) and the position $E_D$ of the impurity (bottom). ($\Delta=0.2eV, n_D=2 \cdot 10^{16}/cm^3$). Inset: $S(T)$ for various scattering rates. (d) $ZT$ with and without lattice contributions (top), carrier concentration that maximizes $S$ or $ZT$ (bottom left), $ZT$ as a function of $n_D$ (bottom right). ($\Delta=0.2eV, \Gamma=5\mu eV$)

the dependence of the quantities entering the expression of $ZT$ as a function of the particle density ($n = p + n_D^+$) at
fixed $T = 220K$, the temperature which maximizes $ZT$ in Fig. 2(a). We use here the total particle density, because in this case the description becomes independent of the bare impurity concentration and the level position, and in particular the additional carriers can find their origin from multiple impurity sources.

For the given gap, and $T = 220K$ no smaller densities than $10^{15}/\text{cm}^3$ can be accessed. The optimum $n_D$ found in Fig. 2(a) translates into $n = 1.3 \cdot 10^{16}/\text{cm}^3$, mainly as a trade-off between decreasing $S^2$ (less entropy per carrier) and increasing $\sigma$ (larger conductivity with more carriers). In particular, we note that $S$ achieves its maximum for smaller concentration of carriers then the figure of merit.

The thermal conductivity in this range varies very slowly with concentration, which is not surprising since we fixed the lattice contribution to thermal conductivity to a fixed value of $\kappa_L/Z^2 = 250\text{W/(Km)}^{-1}$.

Having chosen parameters to represent FeAs$_2$, we note that the experimentally measured carrier concentration in this compound, as inferred from Hall measurements, is $5 \cdot 10^{17}/\text{cm}^3$ in the range of 60-170K which is higher than the density that optimizes $ZT$ in our model. Thus, it seems conceivable that by a deliberate change in the impurity concentration or position, an increase in the figure of merit of the specimen can be achieved.

b. lifetime effects. Next we pick the impurity concentration which maximized $ZT$, and we investigate the role of the scattering rate for the figure of merit and the Seebeck coefficient in Fig. 2(c). The life-time has two effects: through the change in chemical potential, and directly through the dependence of the response functions Eq. (11) on scattering rate $\Gamma$. It is this latter effect that causes the Seebeck coefficient to vanish at low temperatures for a sufficiently large scattering amplitude, as can be seen in the inset of the top panel of Fig. 2(c). The increase in scattering rate reduces the absolute value of both the Seebeck coefficient and the figure of merit, hence long lifetimes are preferred in thermoelectric materials.

The upper limit of the Seebeck coefficient has been discussed above. As a function of the scattering rate $\Gamma$, figure of merit $ZT$ is limited as well. If lifetimes are long, the dependence of the response functions thereof is linear, and thus cancels in the dimensionless ratio $ZT$ if there are no lattice contributions to the thermal conductivity. Therefore, with decreasing $\Gamma$, the figure of merit converges towards the purely electronic limit in which $\kappa_L = 0$.

c. position of the impurity level. In Fig. 2(c) we show the dependence of the figure of merit on the position of the impurity level $E_D$. We fix the impurity concentration to $n_D = 2 \cdot 10^{16}/\text{cm}^3$ and the scattering rate to $\Gamma = 5\mu\text{eV}$. It is clear from Fig. 2(c) that a maximum $ZT$ is achieved when the donor level is very close to the conduction band, which is located at $\Delta/2 = 100\text{meV}$.

This can again be understood from the optimal number of carriers: Indeed, in order to reach the ideal electron density of $1.3 \cdot 10^{16}/\text{cm}^3$, the chemical potential must be rather close to the conduction band. Since at $T = 200K$ and a gap of $\Delta = 0.2\text{eV}$, the compensating holes cannot come from the valence electrons, they have to be supplied from the impurity band, hence, for an impurity concentration $n_D = 2 \cdot 10^{16}/\text{cm}^3$ that is larger than the needed number (if completely ionized), also $E_D$ must be very close to the conduction band.

Next we study the sensitivity of the figure of merit to the lattice thermal conductivity. With thick line in Fig. 2(d) we show the $ZT$ in the absence of lattice thermal conductivity $\kappa_L$ and with a thin line is shown $ZT$ for a constant value of $\kappa_L/Z^2 = 250\text{W/(Km)}$. The figure of merit is clearly enhanced when the lattice conductivity is reduced, hence the desire for a “phonon glass” (see e.g. the review), i.e. a solid which has a low phonon mean free path such as to prevent substantial heat conduction by lattice vibration modes. The effect of lattice thermal conductivity is most enhanced at low temperature and for small impurity concentration.

In Fig. 2(d) lower panel we plot the optimal carrier concentration, which maximizes $S$ or $ZT$, as a function of temperature. The $ZT$ curve is monotonically increasing function of temperature, hence for best performance at higher temperature we need more impurity carriers. To maximize $ZT$ we need larger impurity concentration than we need to maximize $S$. Finally, we also display the value of $ZT$ as a function of impurity concentration for a few representative temperatures. For our setup of parameters, the figure of merit is very sharply peaked at room temperature around a carrier concentration $(n_D \approx 10^{17}/\text{cm}^3)$.

In conclusion, these model considerations give guidance as to where to look for promising thermoelectric materials. In particular we showed that – unless vertex corrections or strongly frequency dependent lifetimes are of pivotal importance – electronic correlation effects are not in the position to enhance the thermopower of a gapped system, they can only shift the asymmetry of the contributions for electrons and holes. Indeed we found the thermopower (of purely electronic origin) to have an upper bound that is given by the size of the gap.

IV. ELECTRONIC STRUCTURE OF FeSb$_2$ AND FeAs$_2$

FeSb$_2$ and FeAs$_2$ both crystallize in the (regular) marcasite structure, have the orthorhombic space group Pmmm, and there are two formula units per unit cell (see Ref. 38). The iron ions are surrounded by distorted pnicogen octahedra, that share corners along the c-axis (see e.g. Ref. 33). In the ligand-field picture, the compounds have a Fe$^{3d^6}$ configuration and the 3d-orbitals are split into $e_g$ and lower lying $2g$ orbitals. The inequivalence of Fe–pnicogen distances causes the $2g$ to split further into two degenerate lower and one higher orbital. In this picture, the compounds are in an insulating low spin state with the two degenerate $2g$ orbitals filled. From this perspective, metalization of FeSb$_2$ is driven by a tem-
temperature induced population of the third \(2g\) orbital. \cite{33,38}

Previous band-structure calculations, however, suggest a more covalent picture, \cite{18}, in the sense that stabilization occurs for \(d\)-orbitals that point towards the ligands, i.e. in particular lowering the \(e_g\) orbitals. In FeSb, this happens to the extend that LDA calculation actually yield a metallic ground-state,\cite{17,18,29,66}.

From the perspective of electronic structure methods, at zero temperature, the challenge is hence to obtain an insulting ground state for FeSb. We therefore compare how three different approaches, GGA, Hybrid functionals and the GW approximation perform in this problem. In preparation for future work which should include correlations beyond GW, and to clarify what would be the starting Hamiltonian to describe these materials, we obtain transfer matrix elements and estimate the values of the interaction using the constrained random phase approximation (cRPA) method.

d. band-structures. Our results for the band-structure of FeSb given by the GGA of DFT as implemented in the Wien2k package\cite{42} are displayed in Fig. 3a. We used the atomic positions at room-temperature (\(a=5.83\) Å, \(b=6.54\) Å, \(c=3.20\) Å)\cite{41,67}. Congruent with previous work\cite{17,18,29}, the GGA ground-state is metallic with small electron pockets half-way between the \(\Gamma\) and \(Z\) symmetry points, and corresponding hole-pockets at all corners, \(R\), of the orthorhombic Brillouin zone. Crucial to the understanding of the gap mechanism within more sophisticated techniques (see the GW discussion below) is to note the different orbital characters of the pockets.\cite{17,18} To quantify this, we transform the local coordinate system of the \(d\)-orbitals into a basis, in which the local projection of the \(d\)-block of the (GGA) Hamiltonian is as diagonal as possible. In this coordinate system, the \(x\) and \(z\) axes point (almost) towards the antimonide atoms,\cite{68} and the \(e_g\) orbitals exhibit the expected bonding/anti-bonding splitting. In this basis, the electron pocket is mainly of \(d_{xy}\) character, and the hole pocket is formed by the now degenerate \(d_{xz}\) and \(d_{yz}\) \((2g)\) orbitals. The respective occupations (within the muffin spheres) are shown in Tab. I.

\begin{table}[h]
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\begin{tabular}{|c|c|c|c|c|c|}
\hline
FeSb & \(d_{z^2}\) & \(d_{x^2-y^2}\) & \(d_{xz}\) & \(d_{yz}\) & \(d_{xy}\) \\
\hline
n & 1.27 & 1.42 & 1.42 & 1.42 & 1.06 \\
\hline
\end{tabular}
\caption{Occupations of the \(d\)-orbitals (in the muffin spheres) within the transformed local coordinate system (see text for details). The \(2g\) orbitals \(d_{xz}\) and \(d_{yz}\) are degenerate and mainly account for the hole pocket. The electron pocket is of \(d_{xy}\) character.}
\end{table}

The GGA band-structure of FeAs\cite{23} (we use \(a=5.3\) Å, \(b=5.98\) Å, \(c=2.88\) Å)\cite{23} is shown in Fig. 3c. With respect to FeSb, the chemical pressure of the larger As atoms is almost isotropic, and the \(c/a\) ratio remains virtually constant (as a function of external pressure, the ratio slightly decreases).\cite{33} In consequence, and as is apparent from the graph, the bands of FeAs are much akin to those of FeSb, and could have roughly been obtained from a rigid band-shift.\cite{69} While within GGA the gap of FeSb was underestimated (no gap at all), the value...
0.28 eV for FeAs₂ is just slightly too large with respect to the experimental 0.2 – 0.22 eV\(^{14,23}\). On a qualitative level, one could thus say that a DFT calculation seems to work rather well for FeAs₂. Below we will explain why we believe this to be a mere coincidence.

e. Maximaly localized Wannier function and cRPA. From a conceptual point of view, we find it insight-full to note, and compare, the hierarchy of transfer matrix elements and the magnitude of local Hubbard interactions. Starting from a full-potential (FP) LMTO (LDA) computation\(^{22}\), we construct maximally localized Wannier functions for the subsystem consisting of the Fe-3d and Sb-5p orbitals, as described e.g. in Ref. \(^{43–45}\) and find that the (largest) nearest neighbor hopping amplitudes in FeSb₂ are \(t_{dd} \sim 0.2 – 0.3\) eV, \(t_{pd} \sim 0.95\) eV, and \(t_{pp} \sim 0.7\) eV. The significance of both the Sb-5p dispersion and the large hybridization was heralded already in the band-structure, Fig. 3\(^a\), and Refs. \(^{17,18}\) as well as in the strong mixed orbitals characters\(^{17,18}\) see also the recent work Ref. \(^{44}\). The two transfers \(t_{pd}, t_{pp}\) being of comparable magnitude, the system thus lies between the canonical Anderson model (\(t_{pp} \gg t_{pd}\)) and cuprate like compounds (\(t_{pp} \ll t_{pd}\)). The centers of gravity of the \(d\) and \(p\) bands are separated by \(\Delta \epsilon_p = 4.2\) eV. In FeAs₂ the \(pd\)-transfers are larger, as expected from the chemical pressure and the covalent/hybridization-like character of the gap. We find \(t_{dd} \sim 0.25 – 0.3\) eV, \(t_{pd} \sim 1.1\) eV, and \(t_{pp} \sim 0.8\) eV, and \(\Delta \epsilon_p = 5.0\) eV.

To investigate the strength of local interactions, we compute the Hubbard \(U\) using the constrained RPA\(^{15,47}\) technique. Since in FeSb₂ and FeAs₂ the eigenvalues of the \(pd\)-subspace are entangled with higher energy bands, we employ the scheme presented in Ref. \(^{48}\).

In relation to the rather small energetic shift needed to deplete the (GGA) pockets in FeSb₂, we find, see Tab. \(^{11}\) that the orbital dependence of the Hubbard \(U\) within the \(d\)-orbitals is of notable 5% – an effect hitherto mostly neglected in methods for correlated materials that start from a parametrized Hubbard-like Hamiltonian. Before, we alluded to the different orbital characters of the pockets. Here, we indeed find for the \(d_{xz}\) and \(d_{yz}\) orbital that mainly account for the hole pocket a value of \(U = 8.6\) eV, while for the orbital corresponding to the character of the electron pocket the interaction is slightly larger with \(U_{dxz} = 8.8\) eV. This differentiation in principle favors a charge transfer towards a gap opening. Further matrix elements of the interaction are \(U_{pd} \sim 2.5\) eV, and \(U_{pp} \sim 4\) eV. The hierarchy of the interaction strength within the \(d\)-shell is already seen in the bare, i.e. unscreened, Coulomb interaction and is thus linked with the construction of the localized orbitals. On general grounds, larger matrix elements of the unscreened interaction are indeed expected for orbitals that have stronger hybridizations with other orbitals, thus are more spatially delocalized\(^{48}\). In analogy with the pressure dependence of Coulombic interactions in a localized basis\(^{19,50}\), the bare interaction of FeAs₂ is larger than that of FeSb₂, in the cubic reference frame we find e.g.

\[
\begin{array}{cccccc}
\text{FeSb₂} & d_{x^2} & d_{y^2} & d_{yz} & d_{xz} & d_{xy} \\
- & - & - & - & - & - \\
\text{FeAs₂} & 8.5 & 7.0 & 7.3 & 7.3 & 7.0 \\
\end{array}
\]

\[d_{x^2} \quad 7.0 \quad 8.8 \quad 7.2 \quad 7.2 \quad 7.5 \\
\]

\[d_{yz} \quad 7.3 \quad 7.2 \quad 8.6 \quad 7.0 \quad 7.3 \\
\]

\[d_{xy} \quad 7.0 \quad 7.5 \quad 7.3 \quad 7.3 \quad 8.8 \\
\]

TABLE II: constraint RPA values for the Hubbard \(U\) (in eV) of FeSb₂ for the Fe₃d-orbitals in the pd-setup of maximally localized Wannier functions in the local coordinate system.

\[V_{dz^2} = 22.7\) eV and \(V_{dz^2-\gamma^2} = 22.6\) eV (the corresponding values for FeSb₂ are \(V_{dz^2} = 22.0\) eV, \(V_{dz^2-\gamma^2} = 21.5\) eV). Moreover, with respect to the antimonide, the Kohn-Sham eigenvalues of the arsenide move towards higher energies, therewith reducing screening strengths, and causing significantly larger values also for the Hubbard \(U\) : \(U_{dz^2} = 11.0\) eV, \(U_{dz^2-\gamma^2} = 10.7\) eV.

f. Hybrid functional approach. Previous attempts to produce an insulating band-structure for FeSb₂ were made within the LDA+U scheme\(^{52}\), where the paramagnetic state (LDA) was found to be stable below a critical U=2.6eV with respect to a ferromagnetically ordered phase (LDA+U)\(^{18}\). Here, we use a hybrid functional approximation (HYB)\(^{20}\), and Fig. 3\(^b\) displays the resulting band-structure, where the B3PW91 functional was used for the \(d\)-orbitals of the iron atoms\(^{70}\). We also note that, while given the freedom, the system does not develop any magnetic moment within this setup (in LDA+U it necessarily does). The band-structure features an indirect gap of about 0.6eV, i.e. it is by far larger than in experiment. This points towards, both, a serious underestimation of static correlations within the previously used GGA, and the lacking of dynamical effects in the hybrid functional approach that will work to reduce the size of the gap.

g. GW approximation. To investigate the dynamical effects of electronic correlations in our compounds, we applied Hedin’s (non-selfconsistent) GW approximation\(^{52}\), which has proven to be quite successful for semiconductors\(^{52,53}\), in its FP-LMTO realization\(^{54}\) to both, FeSb₂ and FeAs₂. In Fig. 4\(^a\), we display, besides the FP-LMTO (LDA) Kohn-Sham energies \(\epsilon_{KS}\), the band-structure obtained by taking into account (perturbatively) the energy shifts as provided by the GW self-energy

\[
\epsilon_{GW} \approx Z \left[ \epsilon_{KS} + \text{Re} \Sigma (\epsilon_{KS}) \right]
\]

with \(Z^{-1} = 1 – \partial_\epsilon \text{Re} \Sigma |_{\epsilon=\epsilon_{KS}}\). In the case of FeSb₂ this indeed opens a charge gap in agreement with experiment. We note that, consistent with the above hybrid functional calculation, as well as with the discussed orbital-dependent interaction strength, the static part of the GW self-energy, i.e. setting \(Z = 1\) in Eq. (22), yields a too large gap of \(\sim 0.2\) eV. Thus, it is the dynamics of the self-energy, therewith a true correlation effect, that
scales down the gap size with respect to Hartree-like approaches – a situation quite akin to that of correlated band insulators.\textsuperscript{55,56} Indeed, the real part of the diagonal matrix-elements of the self-energy are linear in frequency over an extended energy range of up to 10 eV. While the derivative of these elements are basically orbitally independent within the Fe-3d, and Sb-5p orbital subsets, respectively, the different off-diagonal elements yield for the antimonide, in the Kohn-Sham basis, a minimal value (eigenvalue of the self-energy derivative matrix) of $Z \approx 0^{52}$ for "bands" near the Fermi level, and $Z \approx 0^{52} - 0^{57}$ for higher lying "bands"\textsuperscript{71}. Concomitant with the linear slope of the real-part, the imaginary part of the self-energy is basically quadratic, but notably asymmetric with respect to the Fermi level, see Tab. III.

Despite the larger values of the Hubbard $U$, the correlation dynamics is less pronounced in FeAs\textsubscript{2}, and values of $Z$ reach a minimum of $Z \approx 0.6$ for excitations closest to the Fermi level. Also, lifetime effects are both smaller in magnitude, and less asymmetric, see Tab. III. While for FeSb\textsubscript{2} the GW approach has the correct trend with respect to experiments, the gap of FeAs\textsubscript{2} also slightly increases from its LDA value of 0.28 eV to the experimental 0.2 eV. The calculation of the transport coefficients uses a small, frequency and momentum independent scattering rate/self-energy ($\Gamma \sim 20 \text{meV}$), which therewith practically cancels out in the Seebeck coefficient (see the discussion above)\textsuperscript{72} and, at this point, we do not attempt to introduce effects of impurities.

In Fig. 5 we show our theoretical Seebeck coefficient of FeAs\textsubscript{2} and FeSb\textsubscript{2}.
FeAs$_2$ as a function of temperature (green dashed curve), and compare it to experimental results\cite{Sun} for the same polarization. The agreement is excellent in the intrinsic, i.e. not impurity dominated, temperature regime ($T > 12K$).

Also shown is a simple fit, using the formula Eq. (17) for the large gap semiconductor. The individual determination of the parameters $\delta \lambda$ and $\mu$ ($\Delta$ given by experiment) is ambiguous, given the scale of the low temperature thermopower (mV/K) with respect to the high temperature Heikes limit (which is of the order of $k_B/e = 86\mu V/K$). In Fig. 5 we show results for the large gap model for $\Delta/2\delta \lambda - \mu = 85\text{meV}$ (blue dotted curve), which is compatible with the constraints $|\delta \lambda| \leq 1$, and $\mu \leq \Delta/2$.

The decrease of the Seebeck coefficient at low temperature can be understood from our model considerations. This can both be an effect of the scattering rate as well as the presence of impurities, as seen e.g. in Fig. 2(a,b). In the current \textit{ab initio} case, our limited numerical precision (mostly caused by the finite k-mesh) prevents us from endeavoring to include these effects.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{thermopower.png}
\caption{Thermopower of FeAs$_2$. Shown are the theoretical Seebeck coefficient for x-polarization, using a constant self-energy, and the GGA band-structure, with the gap scissored to 0.2eV. Experimental results are of Sun \textit{et al.}\cite{Sun} (x-polarization). Also displayed is a simple 1/T fit corresponding to the large gap semi-conductor, Eq. (17), yielding $\Delta/2\delta \lambda - \mu \approx 86\text{meV}$. Further indicated is the largest possible purely electronic Seebeck coefficient for FeAs$_2$.}
\end{figure}

\subsection{FeSb$_2$}

The situation is entirely different for FeSb$_2$. The maximal measured Seebeck coefficient $S(T = 10K)$ is $-45\text{mV/K}$\cite{Sun}. If one takes the maximum possible asymmetry parameter, $\delta \lambda = 1$, and one assumes the chemical potential to be at the most favorable position, i.e. $\mu = -\Delta/2$, the charge gap must be larger than $\Delta = 0.45\text{eV}$ to explain the value of the measured thermopower in terms of our purely electronic model. The experimental charge gap, however, is only $\Delta \approx 30\text{meV}$. We are thus led to suspect that the large thermopower of FeSb$_2$ at low temperature is not purely of electronic origin.

A possible scenario, mentioned in the literature, is that the very large Seebeck coefficient is mainly caused by a substantial phonon drag, i.e. by an electron drift induced by a scattering with phonons. While there is no conclusive evidence that this effect is operational in FeSb$_2$, there are several reasons why it is more likely to be present in this material than in FeAs$_2$. Since the thermopower is a measure for the entropy per carrier, the phonon contribution to the Seebeck coefficient will be proportional to the lattice specific heat times the electron-phonon coupling constant divided by the electron density. Given Debye temperatures of 348 K for FeSb$_2$\cite{Sun}, and 510 K for FeAs$_2$\cite{Sun}, the specific heat of FeSb$_2$ will be larger than that of FeAs$_2$. The charge carrier concentration at temperatures where the thermopower is maximal, on the other hand, is larger for FeSb$_2$: for the best sample $n \approx 8 \cdot 10^{14}/\text{cm}^3$ whereas for FeAs$_2$ $n \sim 5 \cdot 10^{14}/\text{cm}^3$\cite{Sun}.

The electron-phonon coupling is the least accessible ingredient from the theoretical point of view. Experimentally there are some insinuations: First of all, the low temperature feature seen in the specific heat of FeSb$_2$, that has no analogue in the spin response, and is absent in the arsenide,\cite{Sun} could originate from a substantial electron-phonon coupling, charge ordering, excitonic or polaronic effects from an enhanced coupling to the lattice. Also the nuclear spin-lattice relaxation rate increases below $40K$\cite{Sun}, i.e. in the regime where the thermopower starts its huge magnification, whereas an activation law decrease ($\Delta = 473 \text{ K}$) is found above $50K$\cite{Sun}. Recently, also polarized Raman scattering experiments gave indications for a notable electron-phonon coupling, that is strongly temperature dependent below $40K$\cite{Sun}. Further, we note that, as expected for substantial phonon drag contributions to the Seebeck coefficient, the magnetothermopower of FeSb$_2$ is very low for those samples that exhibit the largest response without magnetic field\cite{Sun}. A decrease in the phonon mean free path by non-electronic scattering (i.e. in particular by imperfections) is expected to lower the respective effect in the thermopower, and indeed the Seebeck coefficient of polycrystalline samples\cite{Sun} and thin films\cite{Sun} was found to be significantly smaller than for single crystals, while having the same high temperature behavior\cite{Sun}. Recently, also substituted FeSb$_{2-x}$As$_x$ was investigated\cite{Sun}. Interestingly, it was found that the above mentioned increase in the susceptibility, starting at around $50K$, is stable with respect to the substitution, whereas the shoulder in
the resistivity at 10 – 20K is flattened out, and the Seebeck coefficient decreases. In the phonon-drag picture this would, again, be owing to a decrease in the phonon mean free path for non-electronic scattering due to the presence of the As “impurities”.

![Graph showing the thermopower of FeSb](image)

**FIG. 6:** Thermopower of FeSb. Shown are our theoretical together with experimental results from Ref. [4] for measurements along the crystallographic orientations x, y, z, as indicated. We can expect reasonable agreement in the temperature range indicated by the gray area. See text for details.

Comparing the thermopower of the antimonide and the arsenide (see Fig. 1 in Ref. [14]) one notes that the Seebeck coefficient of FeAs$_2$ is larger than that of FeSb$_2$ at 35K and higher. This might indicate – if the phonon drag picture holds – that the effective electron-phonon coupling in FeSb$_2$ has sufficiently decreased (by umklapp and phonon-phonon scattering) so that the thermopower is now dominated by the electronic degrees of freedom, i.e. the larger gap in FeAs$_2$ causes a larger response. Yet, we note that optical spectroscopy and some transport measurements see metallic behavior above 70K, or already above 40K, respectively, an effect not captured by our one-particle approach. Hence we will focus on the temperature range from 35K upwards to, at best, 70K.

Since the GGA Kohn-Sham spectrum is metallic, we opted for using the hybrid functional calculation (see Fig. 3(b)), albeit with a gap scissored to the experimental value $\Delta = 0.03\text{eV}$, to compute the theoretical Seebeck coefficient. Moreover, we assume the presence of donor impurities at $E_D = 9\text{meV}$, corresponding to an activation energy $\delta = \Delta/2 - E_D = 6\text{meV}$ as is seen in the resistivity in the range of 5-15K and we use an impurity concentration $n_D = 10^{17}/\text{cm}^3$. This concentration yields $\nu^c_p \approx 7 \cdot 10^{16}/\text{cm}^3$ at 20K, in rough accordance with the respective hole concentration of $4 \cdot 10^{17}/\text{cm}^3$ found in Hall measurements.

We again limit the influence of impurities to their effect on the chemical potential. At high temperature, the latter is linear in $T$ as expected, and, using Eq. (21), we find an effective mass ratio $\eta_c/\eta_v = m^*_v/m^*_c = 0.23$ when using a constant scattering rate, and the very similar $\eta_c/\eta_v = 0.25$ when using the imaginary parts of the self-energy from the GW calculation, i.e. the anisotropy is mainly propelled by the spectral weight and the Fermi velocities, and the GW scattering actually slightly reduces the particle–hole asymmetry in the current case.

We further note that the asymmetry is opposite to that of FeAs$_2$, where we found $\eta_c/\eta_v = 2.5 > 1$.

Thus obtained Seebeck coefficient is displayed in Fig. 6 along with experimental results on single crystals for the three polarizations along the crystallographic axes. In the limited range (discussed above), starting at 35K, and extending towards 70K (indicated by the gray gradient in Fig. 6), we find good agreement with experiments: Both, the order of magnitudes, as well as the hierarchy of polarizations is captured within our approach. Below 35K, the single crystal experiment reaches stellar magnitudes of up to $-45\text{mV/K}$, that we argued to be beyond our approach which neglects vertex corrections. Measurements (not shown) using a polycrystalline sample and on films with preponderant $< 101 >$ orientation display Seebeck coefficients that at low temperatures never surpass $-500$ and $-200\mu\text{V/K}$, respectively, while having the exact same high temperature behavior, advocating a disorder or decoherence induced lowering of the electron drift. At intermediate temperatures, those experiments agree qualitatively with both the single crystal measurements, and our theoretical results.

VI. CONCLUSIONS

In conclusion we have considered the problem of thermoelectricity in correlated insulators and semiconductors. We developed a simple toy model to study how the various many body renormalizations enter the thermoelectric response. We used LDA, hybrid density functional theory and GW methods to carry out a comparative study of two systems of current experimental and theoretical interest FeAs$_2$ and FeSb$_2$. The ratio between strength of the Hubbard $U$ and the bandwidth of FeAs$_2$ and FeSb$_2$ are comparable and so is the correlation strength. In FeAs$_2$ DFT is qualitatively correct, while in FeSb$_2$ correlation effects beyond DFT are essential for obtaining an insulating ground state, and the one shot GW approximation succeeds in that respect. Indeed, using this method, we obtained good agreement with the experimental values of the gap for both materials.

The tools developed in this work were sufficient to describe the thermoelectric response of FeAs$_2$ quantitatively. This framework is not as successful for the FeSb$_2$ compound, and in particular it fails to explain the remarkably high low temperature thermopower discovered by Bentien et al. Our work implies that the latter cannot be understood in the context of local correlations, and one should focus either on vertex corrections to the transport coefficients, or on non local self energy...
effects characteristic to the proximity to a quantum critical point. In this context we notice that within LDA+U this material is close to a ferromagnetic instability. 

An important form of vertex corrections describe the phonon drag effect. A framework to estimate quantitatively these effects in conjunction with \textit{ab initio} methods, are currently not available. Above, we mentioned several experimental findings that suggest the presence of this mechanism in FeSb$_2$, providing a strong incentive to further development in this vein.

Future work should include explicit calculations on correlated insulators using LDA+DMFT to compare with the results of the toy model calculations. Furthermore, the investigation of vertex corrections on the thermoelectricity together with the effects of non local self energies that go beyond the quasiparticle approximation should be considered.

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As is the usual custom when not knowing the incoherent part of the spectrum (as e.g. in slave boson techniques), we omit the Z-factors in the determination of the particle numbers.

for a discussion on this kind of Peierls approximation see e.g.6061

We note that (even in the absence of impurities) the point of charge neutrality and the particle-hole symmetric point for the thermoelectric response are distinct from each other in the asymmetric case, in particular the latter is largely influenced by the Fermi velocities.

This is a typical order of magnitude for our compounds of interest, see e.g.44. For simplicity we are neglecting the sizable temperature dependence of the thermal lattice conductivity.

In Ref. 13 a gap of 0.3eV was extracted from the density of states. Yet, the displayed bandstructure is metallic.

Results (not shown) for low temperature position do not qualitatively differ as far as the following discussion is concerned.

The new unit vectors are x = (−0.627, 0.326, 0.707), y = (0.627, −0.326, 0.707), z = (0.462, 0.887, 0.)

We note that the former hole pockets on the Brillouin zone corners are slightly surpassed as the uppermost valence band by momentum regions on the way from the midpoint of the xz-face (Y-point), up z-wards towards the edge point T.

In Wien2k, the hybrid functional approach is implemented only within the atomic spheres. Therewith one can choose to apply corrections to specific atomic characters.

We stress that the parameter Z occurs here only formally as parameter in the frequency expansion of the self-energy, and should not be confounded with the quasi-particle weight in a Fermi liquid.

There is however an apparent influence of the line broadening on the determination of the chemical potential. Lacking for finite k-mesh sampling (without the tetrahedron method) the numerical precision to avoid spurious in gap spectral weight, we extract from the realistic data the linear high temperature evolution, and, via Eq. (21), use the realistic asymmetry in the semi-conductor model, Eq. (21), (along with a reduced broadening of Γ = 5µeV) for finding the chemical potential at low temperatures.

To avoid pathologies introduced by the necessary numerical broadening in the spectral function, we eliminate spurious weight of valence and conduction bands inside the gap.