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Anti-ferromagnetism enables electron-phonon coupling in iron-based superconductors

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It is shown that a generic form of an anti-ferromagnetic wavefunction opens strong electron-phonon coupling channels in the iron-based superconductors. In the non-magnetic state these channels exist locally on a single iron atom, but are canceled out between the two iron atoms in the primitive unit cell. Our findings are based on symmetry and presence of \(xz/yz\) Fermi surface near the M point and thus should be relevant for the known iron-based pnictide or chalcogenide superconductors.

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There is much evidence that the superconducting state occurs in parallel with the anti-ferromagnetic state in iron-based superconductors. Initially it was suggested in\(^{1}\) that fluctuations of the anti-ferromagnetic state induce electron pairing, and these electrons condense in the superconducting state. Unlike conventional electron-phonon induced pairing, this interaction is repulsive so it requires the superconducting gap to change sign on the Fermi surface. However, experiments on the latest generation\(^{2,3}\) of iron-based materials find a single pocket of carriers with anisotropic but node-less superconducting gap.\(^{4,5}\) While this finding does not rule out all unconventional pairing symmetries, it is consistent with a conventional electron-phonon pairing. In addition, scanning tunneling microscopy features found\(^{6}\) in the FeSe monolayer above the superconducting gap are consistent with the calculated phonon spectral function\(^{7}\) as well as with the kinks in the angle resolved photoemission spectra on a related material\(^{8}\). Therefore, it is possible that electron-phonon interaction can’t be ignored in the iron-based superconductors, and that early\(^{9,10}\) theoretical calculations may have underestimated its strength, as suggested in more recent calculations\(^{7,11}\).

If the electron-phonon interaction is important for pairing, it begs the obvious question: was a parallel occurrence of superconductivity and anti-ferromagnetism just a coincidence? We explore this question here.

As shown below, the electron-phonon matrix element \(g\) in these materials can be decomposed into contributions from individual atoms in the unit cell. Representing with \(\pm 1\) the contribution of a single iron atom to \(g\), we find in the non-magnetic state,

\[
g \simeq 1 - 1 = 0
\]

as the two iron atoms in the cell contribute to \(g\) with an opposing sign (see Fig. 1). However, in an anti-ferromagnetic state, the iron \(d\)-like wavefunction \(\Phi\) of a state of specific spin orientation is localized on only one out of the two iron atoms in the cell. Therefore only one iron atom per cell contributes to \(g\) of this electronic state and the cancellation is prevented, resulting in

\[
g \simeq 4 - 0 = 4.
\]

FIG. 1. Contribution of two iron atoms to the electron-phonon matrix element \(g\) cancel each other in the non-magnetic state, but not in the anti-ferromagnetic state.

The four-fold increase of contribution of the active iron atom to \(g\) is another consequence of the weight transfer in \(\Phi\), and will be discussed later.

Our discussion here is mostly based on symmetry, but we rely on an earlier first-principles calculation\(^7\) to obtain quantities that can’t be inferred from symmetry alone (for example the orbital character of states near the Fermi level). Our earlier first-principles calculations were done for a FeSe monolayer on a SrTiO\(_3\) substrate and were focused on phonons in FeSe, not in the SrTiO\(_3\) substrate. Therefore, our results likely extend to most, if not all, iron-based superconductors, as they all contain layers of FeSe (or FeAs). In fact, several earlier studies\(^{10,12}\) on bulk materials found that electron-phonon matrix elements calculated in the anti-ferromagnetic state of the iron-based superconductors are larger than those in the non-magnetic state (see also Fig. S.1 in the supplement\(^{13}\)). However, the microscopic origin of this increase remained unclear.

The focus of our paper is on the interplay between anti-ferromagnetism and electron-phonon interaction, not on the pairing symmetry. In general one expects that pairing may have contributions not only from electron-phonon interactions but also from other interactions such as magnetic or orbital fluctuations.\(^{14,15}\) Therefore, while our finding about electron-phonon interactions is general, it does not specifically address the resulting pairing symmetries across the families of iron-based superconductors.

We would also like to point out that the arguments
presented in this paper assume that the xz/yz bands are crossing the Fermi level near the M point of the Brillouin zone. Nevertheless, as far as we are aware, xz/yz bands cross the Fermi level near the M point in all the known iron-based pnictide or chalcogenide superconductors.

In what follows we first discuss two specific electronic states of an FeSe monolayer with the smallest primitive unit cell: the non-magnetic (NM) and the checkerboard anti-ferromagnetic (cAFM) state. Later we generalize our findings to nearly any ordered or disordered static anti-ferromagnetic state (with correlation length exceeding the unit cell dimension). We leave for future work the role of dynamic effects of the anti-ferromagnetic order.

We work here within the density functional theory framework where formally the exact ground state electron density \( \rho \) is written in terms of the effective (Kohn-Sham) electron orbitals \( \Phi_I \) that solve the Schrödinger-like equation with an effective one-body potential \( V \). Index \( I \) on \( \Phi_I \) refers both to the band index \( n \) and the crystal momentum \( k \). These orbitals characterize the quasiparticle states of the system. In the spin-polarized variant of the theory orbitals for up and down spins may have different spatial dependence, \( \Phi_{I\uparrow}(r) \neq \Phi_{I\downarrow}(r) \). In particular, a (collinear) anti-ferromagnetic state has occupied electron orbitals for up and down spin located on a different subset of magnetic atoms in the unit cell. For the simplest order, cAFM discussed earlier, the wavefunction \( \Phi_{I\uparrow}(r) \) is mostly confined to one of the two iron atoms in the primitive unit cell while the corresponding \( \Phi_{I\downarrow}(r) \) is on the other. In the NM state, orbitals of both spin types exist in equal amplitudes on both Fe atoms in the cell, as sketched out in Fig. 2.

\[
\begin{array}{c}
\text{a) Non-magnetic} & \text{b) Anti-ferromagnetic} \\
\Phi_{I\uparrow}^2 & \Phi_{I\uparrow}^2 \\
\Phi_{I\downarrow}^2 & \Phi_{I\downarrow}^2 \\
\end{array}
\]

FIG. 2. Sketch of the amplitude of electron orbital of up and down spin in the non-magnetic (left) and the anti-ferromagnetic (right) state. In our first-principles calculation for the anti-ferromagnetic state, the weight of the electron wavefunction crossing the Fermi level is 85% on one of the iron atoms, 5% on the other iron atom, and the remaining weight is mostly on selenium p-states.

Even though the electron orbital \( \Phi_I \) might be localized on only one of the Fe atoms in the unit cell, with a well-defined crystal momentum it must periodically extend over all unit cells in the crystal. Similarly, a phonon with a well defined momentum corresponds to a displacement of all atoms in the crystal. Therefore as atoms vibrate around their equilibrium positions they change the effective potential \( V \rightarrow V + \partial V_J \) across the entire crystal. Here the phonon index \( J \) replaces phonon momentum \( \mathbf{q} \) and the branch index \( \nu \).

The overlap integral of these three extended quantities,

\[
g = \langle \Phi_I | \partial V_J | \Phi_J' \rangle,
\]

defines the electron-phonon matrix elements, one of the main ingredients that determines the electron pairing strength within the BCS mechanism of superconductivity. However, representing \( g \) in terms of extended quantities is not convenient for understanding the microscopic reason for the magnitude of \( g \). Therefore we rewrite the calculated extended electronic state \( \Phi_I \) as a sum of functions \( \phi_i \) highly localized on a single atom in the crystal, \( \Phi_I = \sum \phi_i \). Later we will give an explicit expression for \( \phi_i \). Similarly, we rewrite the calculated extended potential change \( \partial V_J \) due to a phonon as the sum of potential changes arising from the movement of individual atoms in the crystal, \( \partial V_J = \sum_j \partial v_j \).

Now, following Ref. 16, an extended matrix element \( g \) is rewritten as a sum of products of localized quantities \( \phi_i, \partial v_j, \) and \( \phi_{J'} \),

\[
g = \langle \Phi_I | \partial V_J | \Phi_J' \rangle
= \left( \sum_i \langle \phi_i | \right) \left( \sum_j \partial v_j \right) \left( \sum_{J'} \phi_{J'} \right).
\]

Since all three components \( \phi_i, \partial v_j, \) and \( \phi_{J'} \) may be constructed using Wannier function concept so that they are exponentially localized in real space, they will generally contribute to \( g \) only when they are close to each other. In the case of a FeSe monolayer we find that dominant contributions for electronic states at the Fermi level either have \( i, i', \) and \( j \) associated with the same iron atom, or \( i \) and \( i' \) on the same iron atom and \( j \) on the neighboring selenium atom (numerical values are given in the supplement\(^{13}\)). Since orbitals \( i \) and \( i' \) are associated with a single iron atom we can group them together so that \( g \) is rewritten as a sum over all iron atoms \( a \) in the crystal,

\[
g \simeq \sum_{\text{atoms}} g_a.
\]

We now make the analysis more concrete by rewriting the extended electron wavefunction \( \Phi_I = \Phi_{n\mathbf{k}} \) as a vector \( C \) in an explicit localized atomic basis \( |am\rangle \), such as a maximally localized Wannier function\(^{17}\) with an orbital character \( m \),

\[
|\Phi_I\rangle = \sum_{am} C_{am}^I e^{i\mathbf{k}\cdot\mathbf{r}_a} |am\rangle.
\]

Unless specified otherwise, the electron momentum \( \mathbf{k} \) is defined in the two-iron atom unit cell. Iron \( d \)-like orbital characters indexed with \( m \) are always defined in the Cartesian frame of the one-iron atom cell (\( z \) axis is perpendicular to the FeSe plane). Vector \( \mathbf{r}_a \) points to atom \( a \) and coefficients \( C_{am}^I \) are cell-periodic by Bloch’s theorem.
Similarly, we rewrite $\partial V_J = \partial V_{b\beta}$ in terms of a potential change $\partial \nu_{b\beta}$ due to the displacement of a single atom $b$ in Cartesian direction $\beta$,

$$\partial V_J = \sum_{b\beta} \xi_{b\beta} e^{i\mathbf{q} \cdot \mathbf{r}_b} \partial \nu_{b\beta}.$$ 

Here $\xi$ is the polarization vector for phonon $J$ specifying the cell-periodic displacement of atom $b$ in direction $\beta$.

Inserting a decomposition of $\Phi$ and $\delta V$ into Eq. 1 and using the simplification from Eq. 3 after some algebra we obtain the contribution of a single iron atom $a$ to $g$,

$$g_a = \sum_{mm'} C_{am} C_{am'} \sum_{b\beta} \xi_{b\beta} e^{i\mathbf{q} \cdot (\mathbf{r}_a - \mathbf{r}_b)} \langle am | \partial \nu_{b\beta} | am' \rangle.$$ 

(5)

It is clear from the cell-periodicity of $C$ that $g_a$ is cell periodic as well, so it will be sufficient to compute $g_a$ only in the primitive unit cell.

In our previous report, we computed $g$ from first principles in a cAFM state of a FeSe monolayer and identified two channels, namely 1 and 2, that have by far the largest $g$ among all states $I$ and $I'$ on the Fermi surface and among all phonon modes $J$. Therefore in what follows we focus our discussion only on these two channels.

Now we are ready to compute the sign of $g_a$ on both iron atoms in the primitive unit cell, for both channels (1 and 2). The sign of $g_a$ is determined by the signs of the $C$’s, $\xi$, the exponential factor, and $\langle am | \partial \nu_{b\beta} | am' \rangle$ appearing in Eq. 5. We start by analyzing channel 2 in the NM state.

i) Channel 2 scatters states originating from $d$-like $xz$ states to those of $yz$ character, and vice versa. In the two-iron unit cell both $xz$ and $yz$ states cross the Fermi level near the $K = (\pi, \pi)$ point, but in the unfolded one-iron unit cell one state is near $(0, \pi)$ and another near $(\pi, 0)$. Therefore, as illustrated in Fig. 3, in the two-iron unit cell $\Phi$ either has opposite sign on two iron atoms in the cell, or the same sign. Which is which depends on the choice of the basis atoms in the two-iron cell, and will not be relevant for the following discussion. Recalling the definition of the coefficients $C$ from Eq. 4 and using $K = (\pi, \pi)$ we conclude that for one of the states the relative sign of $C$ is the same on both iron atoms, and opposite for the other state.

ii) The phonon eigenvector $\xi$ in channel 2 has an opposite sign on two iron atoms in the unit cell, as it consists of an out of phase vertical displacement of iron atoms, with first neighboring iron atoms moving in opposite directions. In the one-iron unit cell this mode would have $\mathbf{q} = (\pi, \pi)$, consistent with the fact that this channel scatters state $K = (0, \pi)$ to state $(\pi, 0)$.

iii) The exponential factor $e^{i\mathbf{q} \cdot (\mathbf{r}_a - \mathbf{r}_b)}$ equals 1 as a vertical displacement of an iron atom dominantly affects the localized orbital on the same iron atom. Therefore $a = b$, so $\mathbf{r}_a = \mathbf{r}_b$, and $e^{i\mathbf{q} \cdot (\mathbf{r}_a - \mathbf{r}_b)} = 1$.

iv) Finally, since channel 2 couples a $d$-like $xz$ to $yz$ state, the matrix element $\langle am | \partial \nu_{b\beta} | am' \rangle$ is dominated by the induced potential $\partial \nu_{b\beta}$ with $xy$ character (since $\langle xz | xy \rangle \neq 0$). By symmetry, displacement of the iron atom perpendicular to the FeSe plane can create this kind of potential only through an interaction with neighboring selenium atoms, as shown in Fig. 5 and discussed later in more detail. However, selenium tetrahedra formed around two iron atoms in the unit cell are inverted images of each other (compare red and blue tetrahedra in Fig. 1) so the induced $xy$ potential has an opposite sign as well, as confirmed by an explicit calculation of the matrix element given in the supplement.

Since the relative sign on the two iron atoms is opposite for an odd number of factors (i, ii, and iv) we conclude that the relative sign of $g_a$ is opposite as well. Therefore, the electron-phonon matrix element $g$ for channel 2 vanishes in the non-magnetic state.

In the cAFM case the wavefunction coefficient $C$ for a state at the Fermi level is zero on one of the Fe atoms in the unit cell (see Fig. 2), so the corresponding $g_a$ is zero as well. This prevents cancellation of the contributions between two atoms in the unit cell. The remaining iron atom has $g_a$ four times larger than in the NM case. The reason for this increase is the two-fold increase in a specific spin density on the iron atom in the cAFM state compared to the NM state. Therefore the square of the wavefunction coefficient $|C|^2$ appearing in Eq. 5 is increased by a factor of two. Another factor of two originates from a two-fold increase in the $xy$-like induced potential $\delta \nu_{b\beta}$ upon vertical displacement of an iron atom (see also for numerical values).

Generalization to nearly any kind of anti-ferromagnetic order is now straightforward. Let us consider a large $N \times N$ supercell of FeSe with an arbitrary anti-ferromagnetic ordering of spins (as shown in panels a, b, and c of Fig. 4). For any such order, we can construct a pattern of atom displacements in which all up-spin iron atoms move vertically above the FeSe plane, and all down-spins move into the plane (or vice-versa). One can easily check from our previous analysis that the electron-phonon matrix element $g$ for this displacement pattern will be as large as in the cAFM state. For example, it would be enough to
The discussion for channel 1 reaches a similar conclusion to that for channel 2: in the NM state, two iron atoms in the supercell contribute to $g$ with opposite signs. Phonon coupling in channel 1 is a soft mode responsible for a condensation of the so-called orthorhombic (nematic) ground state in bulk FeSe\textsuperscript{18}. Therefore this mode consists of an in-plane displacement of both iron and selenium atoms. However, only the in-plane displacement of selenium atoms contributes to $g_a$. In particular, coupling is large only when selenium atom $b$ moves towards or away from the neighboring iron atom $a$ (see supplement\textsuperscript{15} for explicit numerical values of $g_a$). Since $a \neq b$ we have $\mathbf{r}_a \neq \mathbf{r}_b$ and the exponential term appearing in Eq. 5 leads to the dependence of the matrix element $g$ on the phonon momentum $\mathbf{q}$. We find that $g \sim |\mathbf{q}|$ for small $\mathbf{q}$ and therefore conclude that the forward scattering is greatly suppressed over the backward scattering in channel 1. Another difference with respect to channel 2 is that states coupled in channel 1 have the same orbital character ($d$-like $xz$ state couples to $xz$, and $yz$ to $yz$).

Our findings rely on having $xz/yz$ bands crossing the Fermi level near the M point as found experimentally. However, some DFT calculations result in electronic structure that is inconsistent with experiment. In these cases, our mechanism may or may not apply. One such example is the DFT calculated band structure for the striped phase. Nevertheless, the calculated electron-phonon interaction with the incorrect band structure is also stronger than in the non-magnetic case. Sorting out these theoretical observations is worthy of a future study, but it is beyond the focus and scope of our current work.

In closing, we discuss the role of selenium height on the magnitude of the electron-phonon matrix element $g$. In channel 1 the dominant contribution to $g$ comes from the displacement of selenium atoms, so it would not be surprising that the position of selenium atoms would be relevant for $g$ in that channel. Somewhat less expected is our finding that the position of a selenium atom is crucial for coupling in channel 2, since it involves displacement of iron atoms. As mentioned earlier, this channel scatters a $d$-like $xz$ state to $yz$ and therefore the potential induced by the upward movement of the iron atom must have a $xy$-like component. However, iron atoms are all in the same plane, so the $xy$-like potential component must originate from the interaction of an iron atom with neighboring tetrahedrally bonded selenium atoms. We confirmed this finding with an explicit first-principles calculation on a $3 \times 3$ FeSe unit cell. Upward displacement of a single iron atom in this enlarged cell transfers electron’s charge into the Fe-Se bond above the iron plane (making the bond more covalent), and out of the Fe-Se bond below the plane. Therefore, the induced potential has some $xy$ component as shown in Fig. 5.

![FIG. 5. The iso-surfaces of the potential induced by an upward displacement (along $z$) of the iron atom inside the tetrahedral selenium cage. The ellipsoidal asymmetry in the iso-surfaces along the $x$ and $y$ axes indicate presence of a $d$-like $xy$ component of the induced potential. The Cartesian labels are given in the one-iron unit cell convention. The red (blue) iso-surface is drawn at a constant value of the induced potential. This constant value is set at 2% of the maximal (minimal) value of the induced potential.](image)
This means that $g$ in channel 2 is maximized when the direction of the iron-selenium bond is aligned with the extremal points of product $\Phi^*_I \Phi_I$. Since the electron wavefunctions have $d$-like $xz$ and $yz$ character, $g$ is maximal when Fe-Se bonds form an ideal tetrahedron. This is consistent with the empirical finding that an ideal tetrahedral environment gives the highest superconducting transition temperature\textsuperscript{19}.

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