Dramatically Enhanced Superconductivity in Elemental Bismuth from Excitonic Fluctuation Exchange

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Motivated by the remarkable discovery of superconductivity in elemental Bismuth at ambient pressure, we study its normal state in detail using a combination of tight-binding (TB) band-structure supplemented by dynamical mean-field theory (DMFT). We show that a two-fluid model composed of preformed and dynamically fluctuating excitons coupled to a tiny number of carriers provides a unified rationalization of a range of ill-understood normal state spectral and transport data. Based on these, we propose that resonant scattering involving a very low density of renormalized carriers and the excitonic liquid drives logarithmic enhancement of vertex corrections, boosting superconductivity in Bi. A confirmatory test for our proposal would be the experimental verification of an excitonic semiconductor with electronic nematicity as a ‘competing order’ on inducing a semi-metal-to semiconductor transition in Bi by an external perturbation like pressure.

Rhombohedral Bismuth (Bi) has recently acquired prominence in a variety of contexts. Like graphite, elemental Bi also shows a magnetic field- as well as a pressure-induced metal-insulator-like transition with a large magneto-resistance, and electron fractionalization in high magnetic fields. The remarkable discovery of superconductivity (SC) at ambient pressure in Bi at very low temperature (Tc below 0.53 mK), in a lowest carrier-density elemental system to date, adds to the range of novel behaviors exhibited by this ‘simple’ system. The following experimental features of this discovery: (i) Tc ≃ 0.5 mK, while a naive BCS estimate would yield Tc ≈ 100 nano-K, and (ii) Bc(0)/Tc is experimentally found to be 9.4 T/K, greatly enhanced over the BCS estimate of 0.79 T/K, strongly point to the possibility of SC in Bi not being of the standard BCS variety and, in fact, it would be more aptly characterized as being in a non-adiabatic limit of pairing theory if electron-boson (e.g., phonon) coupling were to be invoked as the dominant pairing glue.

The unique electronic properties of Bi arise from successive distortions of a higher-symmetry simple-cubic structure, which, except for a tiny change in the b/d-parameter (here, d is the spacing of the (111)-rhombohedral planes of one sublattice, while b is the displacement of the other sublattice (111)-planes from these) remains unchanged from high (300 K) to low (4.2 K) temperature. This is a small relative displacement along the body diagonal, which doubles the unit cell. By itself, this would drive Bi into a Peierls-like band semiconductor. But additional rhombohedral shear (which exists from T= 300 K down to 4.2 K) causes further lowering of symmetry, allowing valence and conduction band overlap in good accord with band structure studies, and explains why Bi is metallic. Due to extremely small Fermi pockets (of size 10−5 of the Brillouin zone) and tiny carrier density (3 × 1017 cm−3), the long mean-free path accounts for its small low-T resistivity. One might then think that traditional one-electron band structure is an adequate framework to understand its electronic properties.

However, careful perusal of extant data points toward a much more interesting situation. Early data show that the electrical resistivity ρ0(T) ≃ ρ0 + AT2 for T < 50 K. But very unusual behavior at low T, with ρ0(T) ≃ T5 was recently seen: within an itinerant picture involving long-range coulomb interaction, an attempt was made to rationalize this behavior in a plasmaron picture, where a plasmaron is considered to be a collective excitation.

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invoking strong coupling between the charge carriers and plasmons. Optical data raise additional issues in this context. While a tiny Drude component is visible at low energy, finding of sizable mid-infra-red (mid-IR) absorption is inexplicable in a free-electron framework. Given the tiny Fermi energy \( \approx 25 \text{ meV} \), a large \( T \)-dependent transfer of spectral weight over a much larger scale \( \approx 300 \text{ meV} \) also presents a challenge for the standard (uncorrelated band) view.

On the theoretical front, first-principles density-functional theory (DFT) and Slater-Koster tight-binding fits conclusively show that (i) there are multiple tiny pockets in \( Bi \), with two being almost perfectly compensated, and the others supplying a tiny additional number of carriers. Moreover, the importance of spin-orbit coupling (SOC) is shown by the fact that the correct positions of the \( e(\hbar) \) pockets are only found when sizable SOC, \( O(1.3) \text{ eV} \), is included in band calculations. Thus, DFT + SOC calculations indeed yield good description of ground state properties. In particular, they yield the correct shape and size of the carrier pockets at the \( T \) and \( L \) points in the Brillouin zone. However, it is also well known that such a ground state theory can fall short of describing the dynamical excitation spectrum and, consequently, the finite-temperature \( (T) \) responses. This is especially true when coulomb interaction-induced bound state formation is involved: in the specific context of \( Bi \), it is long known that the dominant interaction between conduction and valence band carriers in \( Bi \) is of short-range excitonic character. This raises the possibility of exciton bound state formation, but its role in \( Bi \) has never been satisfactorily addressed. While DFT calculations work well in moderately correlated solids, they are, by construction, inadequate when such (e.g. excitonic) bound states appear due to correlations. Generically, one also expects a symmetry-adapted coupling of such interband excitons to intervalley phonons in \( Bi \): since the ratio of the phonon Debye frequency \( k_B \Theta_D \) to \( E_F \) is large (\( \gtrsim 0.5 \) in \( Bi \)), sizable self-energy effects due to this additional coupling will also influence normal state responses above \( T_c \). As far as SC is concerned, the huge enhancement of the SC transition temperature, \( T_{c \text{exp}}/T_{c \text{BCS}} \approx 10^{-3} - 10^{-4} \), and the large discrepancy between the measured and calculated (within BCS theory) ratio of the upper critical field to \( T_c \) also mandates a non-adiabatic ‘strong coupling’ SC beyond a traditional BCS view.

These limited observations must constrain theoretical modelling: both unconventional metallicity and SC must find explication in a picture based on (i) the special band structure of \( Bi \) and (ii) strong scattering (electron-hole and/or electron-phonon interaction) processes beyond DFT and, in particular, microscopic processes which generate a \( \Delta_{SC} \approx T^5 \) should also be involved in generating the SC pair glue.

Our first observation is that two of the multiple electron(hole) pockets of \( Bi \) are almost perfectly compensated, leading to a situation famously encountered in transition-metal dichalcogenides (TMD), where preformed excitonic liquid (PEL) driven charge-density-wave (CDW) states are ubiquitous. Band calculations reveal a total of four pockets in \( Bi \), of which two (electron and hole) exhibit a propensity for an excitonic instability. Were there only these two pockets, one would have expected an electron-hole attraction-mediated excitonic insulator. However, the remaining pockets now supply a tiny number of additional carriers, leading to a physical picture of an incipient excitonic insulator self-doped with a tiny number of carriers. Thus, we are led to a model of a self-doped excitonic insulator (EI) in the intermediate coupling limit, where novel physics can arise from strong scattering between carriers and preformed but uncondensed excitons. In this report, we establish this using tight-binding-plus dynamical mean-field theory (TB-DMFT) calculations as done earlier. Armed with very good accord with normal state transport, we elucidate a ‘strong coupling’ electronic mechanism, wherein strong resonant scattering between carriers and preformed excitons enhances vertex corrections, boosting the SC \( T_c \) in \( Bi \).

We begin with a tight-binding description of the DFT band structure by using the Slater-Koster (SK) fit with form factors and parameters including SOC as in earlier work (given in SI Table 1). The resulting band structure in Fig. 1 excellently reproduces all the pockets seen in full DFT calculations, constituting the appropriate band structural input for the correlation calculations. Guided by the discussion above, we focus on inter-band

**Figure 1.** Tight binding (TB) bands of \( Bi \) in the rhombohedral structure including spin-orbit coupling (SOC) as done in ref. 6. For the effective two-band model incorporating excitonic correlations within DMFT calculations, we use only the two (red and green) bands crossing the Fermi energy, \( E_F(=0) \).
excitonic correlations. The two-band Hubbard model incorporating the two bands crossing the Fermi energy is $H = H_0 + H_1$, with

$$H_0 = \sum_{(\mu, \nu = 1,2), k, \sigma} (\varepsilon_\mu(k)c_{\mu k\sigma}^\dagger c_{\nu k\sigma} + h.c.) + \Delta \sum_i (n_{i,1} - n_{i,2})$$

where $\varepsilon_\mu(k) = E_\mu + 3V_{pp\mu}(\cos(\frac{k_\mu}{2}) - \cos(\frac{k_\nu}{2})$ + $V_{pp\nu}(\cos(\frac{k_\nu}{2}) + 2\cos k_\mu)$ + $3V_{pp\mu}(\cos(\frac{k_\mu}{2}) - \cos(\frac{k_\nu}{2})$, (represented by the green line in Fig. 1), $\varepsilon_\nu(k) = E_\nu + 3V_{pp\nu}(\cos(\frac{k_\nu}{2}) + 2\cos k_\mu)$ + $3V_{pp\nu}(\cos(\frac{k_\mu}{2}) + \cos(\frac{k_\nu}{2})$, (red line in Fig. 1) where $E_\mu$ is on-site energy ($-9.643$ eV), $V_{pp\mu}$ and $V_{pp\nu}$ ($2.271$ eV and $-0.679$ eV, respectively) are the nearest neighbour hoppings and the interband matrix element $V_{\nu\mu}(k) = 2i\sqrt{3} \sin(\frac{\sqrt{3}k_\mu}{2}) \sin(\frac{\sqrt{3}k_\nu}{2})$, as extracted earlier from an SK fit. Here $c_\mu$ and $c_\mu^\dagger$ represent annihilation and creation operator and $\mu, \nu$ represent the band indices for the two bands crossing $E_F(=0)$ in Fig. 1. The interaction terms are

$$H_1 = \sum_{\mu = 1,2} U_{\mu,\mu'} \sum_\nu n_{\nu,1} n_{\nu,\mu'} + \sum_\mu U' \sum_\nu n_{\nu,\mu} n_{\nu,2}$$

taken to be dominantly local with the understanding$^8$ that the coupling between valence and conduction band carriers is of short-range excitonic character, as pointed out above. Here $U_{1,1} = U_{2,2}$ and $U'$ are the intra- and inter-band Coulomb interactions, $n_{\nu,\mu} = \sum_{\sigma} n_{\nu,\mu\sigma}$.

It is also long known that coupling of carriers to intervalley phonons is significant in $\text{Bi}_2$.$^3$ Since the dimensionless measure of the electron-phonon (e-p) coupling, defined as $g = \omega_p/E_F \approx 0.5 (\omega_p$ the Debye frequency, and $E_F$ = Fermi energy) is actually large in $\text{Bi}_2$, one is in a non-adiabatic limit of the electron-phonon problem. This intermediate coupling regime is also interesting in that in contrast to the anti-adiabatic ($g \rightarrow \infty$) or the adiabatic ($g \rightarrow 0$) limits, one cannot “integrate out” these phonons to give a further e-h attraction as done in earlier work.$^{13, 14}$ Technically, coupling of inter-band excitons to phonons is described by adding a term $H_{e-p} = g \sum_\alpha \varepsilon_{\alpha,\nu,1}(A_\alpha + A^\dagger_\alpha)$ to $H$ above, where $A_\alpha, A^\dagger_\alpha$ are the intervalley phonon operators. We have followed our earlier procedure$^{13, 14}$ to incorporate the sizable e-p coupling effects for Bi in the two-band Hubbard-like model above within the DMFT procedure (see SI(A) for details).

**TB+DMFT Results and Transport**

We solve the two-band model above using DMFT, with multi-orbital iterated perturbation theory (IPT) as the ‘impurity’ solver. Though not numerically exact, it works very well in real multi-band cases for all temperatures and band-fillings, especially in cases where there is a sizable crystal-field splitting between the bands (here, between valence and conduction band). The bands that serve as a band-structural input to the correlation calculations are obtained by diagonalizing the one-electron part of the two-band model above, and labelled henceforth as “a” and “b” band states. IPT is a fast solver and its efficacy in a wide range of real systems is known$^{11-15}$. In contrast, though state-of-the-art continuous-time quantum Monte Carlo (CTQMC) solvers are much more “numerically exact”, they cannot access temperatures below $O(20)$ K at present$^{16}$, which makes them unsuitable for investigation of the low $T(<K)$ behavior in $\text{Bi}_2$. We choose $U_{1,1} = U_{2,2} = 0.5–0.7$ eV (henceforth we use the notation $U = U_{1,1} = U_{2,2}$ and $U' = 0.1–0.2$ eV as appropriate parameters, and while ab-initio estimates will yield more precise estimates, our present choice is physically motivated. Given widths of $O(1.0–1.5)$ eV for the valence (VB) and conduction (CB) bands, we are in the intermediate coupling limit of the two-band model. This is precisely the case where DMFT works best (for specific studies in the BCS-BEC crossover, see ref. 17).

In Fig. 2, we show the TB+DMFT local density-of-states (LDOS) as $U, U'$ are cranked up. While correlations gradually close the band gap for the VB (a’-orbital) as expected$^{14}$, they reduce the LDOS at $E_F$ for the CB (b’ orbital): these contrasting behaviors are direct consequences of distinct effects of local electronic correlations on band-insulating and metallic subsets of the non-interacting band structure. Around $U \approx W_p$, the $b$-fermion band-width, we observe eventual opening of a Mott-like gap in the LDOS, seen by the fact that $\text{Im} \Sigma_\omega(\omega)$ develops a pole structure at $E_F$ in this case. Correspondingly, the $a$-fermion states retain metallic character: this would be an interesting manifestation of orbital-selective Mott physics in semi-metals and, were such a strong coupling regime to be realized, would open up the possibility to development of novel instabilities with concomitant competing orders$^{19,20}$ as, for instance, in Fe-arsenides. However, this is not the regime applicable to Bi, and so we concentrate on the smaller $U$ regime.

Choosing $U = 0.5$ eV, $U' = 0.15$ eV, we next compute the $dc$ resistivity using the Kubo formalism in DMFT. This task is facilitated by the finding$^3$ that irreducible vertex corrections appearing in the Bethe-Salpeter equations for conduc-
tivities are negligible and can be ignored to a very good approximation. Interestingly, as shown in Fig. 3, we find that a linear-in-$T$ behavior of $\rho_{\mu}(T)$ at high $T \geq 40–50$ K smoothly crosses over to a Fermi-liquid-like $T^2$ behavior up to about 10 K and, remarkably, exhibits a further low-$T$ crossover to a good metal with $\rho_{\mu}(T) \propto T^2$. This is in very good accord with experimental trends, and mandates deeper microscopic rationalization. To cement the link between transport and excitonic liquid fluctuations, Fig. 3 shows the excitonic average, computed as $\Delta_{\omega}\rho_{\mu}(\omega) = \langle (1/\pi) - \text{Im} \Sigma_{\omega}(\omega)\rangle (G_{\mu}(\omega))$ is itself computed from two-band DMFT(IPT) as previously done for TMDs.$^{13, 14}$ We find a clear correlation between $\Delta_{\omega}\rho_{\mu}(T)$ and the T-dependence of $\rho_{\mu}(T)$ over a wide T range. This provides strong theoretical evidence linking the $dc$ resistivity to microscopic processes involving scattering of the tiny number of carriers off well-formed and quasi-local excitonic correlations: at high-$T$, the latter are incoherent, leading to a quasi-linear-in-$T$ resistivity, while increasing one-fermion coherence via suppression of incoherent excitonic fluctuations provides a rationalization for onset of enhanced metallic coherence, and the near $T^2$ behavior at intermediate $T$. 

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**Note:** The above text is a natural reading of the document, focusing on the scientific content and avoiding unnecessary formatting or less relevant information.
Remarkably, however, we also obtain the $\rho_{TT}(T) \propto T^5$ dependence at very low $T$, the latter correlated with a reduced excitonic fluctuation at low $T$. Thus, we identify a new element: the $T$-dependence of sizable and dynamical 'preformed' excitonic correlations governs the $T$-dependent resistivity in a wide $T$ window. Our proposal is distinct from: (i) the plasmaron view, where weak-coupling RPA-like analysis is employed to get $\rho_{TT}(T) \propto T^2$ from long-range interactions, and (ii) pure $e$-$p$ coupling models, which could, in principle, also yield similar behavior when $T \ll \Theta_D$, something that may also obtain in Bi. We emphasize that $e$-$p$ coupling is included on an equal footing with local excitonic correlations in our approach. We propose that a way to distinguish between these distinct scenarios could be $T$-dependent tunnelling measurements at small-to-intermediate $T$: as a function of $T$, the conductance $g(V) = dI/dV$ would, in a picture involving coupling of carriers to any bosonic mode(s), show finite-voltage (energy) peak-dip-hump features. The energies and spectral weights of such features could be compared with estimates from different bosonic channels, allowing a determination of the most important fermion-boson scattering channel. However, since there will always be a symmetry-dictated coupling of interband excitons to intervalley phonons, one also generally expects two bosonic modes at different energies to show up in $g(V)$: their relative weights provide an estimate of the relative importance of carrier-exciton vis-a-vis carrier-phonon coupling.

Figure 2. The orbitally resolved density of states from TB+DMFT at various $U$ and $U'$.

Figure 3. DC Resistivity for Bi within TB+DMFT: a $T$-dependent crossover from $\rho_{TT}(T) \propto T^2$ below $T \approx 90$ K is followed by $\rho_{TT}(T) \propto T^5$ form at very low $T$ (see inset for comparison with data points) is clear. The lower-left inset shows the $T$-dependent local interband excitonic average: onset of $\rho_{TT}(T) \propto T^5$ correlates with a decreasing excitonic average. The y-axis of both the insets are divided by $10^{-3}$.
Further support for our view arises when we compare the DMFT optical conductivity, $\sigma_{\omega}(\omega)$ in Fig. 4 as a function of $T$ with published data. Since we have kept only the two lowest bands crossing $E_F$ from the SK fit, we do not expect accord at higher energies, but can readily make a comparison for the relevant energy range (few hundred milli-eV) of interest. Specifically, up to about 100 meV, our result matches quite well with data, including (i) the plasmon edge, (ii) the detailed optical lineshape as a function of energy up to about 100 meV, (iii) sizable optical spectral weight transfer upon raising $T$. The plasmon-like features are clearly visible on both regular and log-log plots as a clear absorption onset at $\approx 15$ meV at low $T = 10$ K. Interestingly, it is also preceded by a ‘prepeak’ structure, centered at $\approx 10$ meV, in accord with observations. This prepeak feature is also washed out with increasing $T$, in accord with data. Clear spectral weight transfer up to $\approx 200$ meV is also found: given the tiny $E_F$ in Bi, this is quite a large energy scale, attesting to considerable dynamical correlations. Finally, we find an isosbestic point around 5.0 meV as a function of $T$, which is another characteristic signature of dynamical electronic correlations that could be tested in extant work.

Taken together, our results strongly support the idea that a common underlying scattering process involving tiny number of carriers and uncondensed fluctuating interband excitons may be operative in Bi. That the dominant interaction in Bi has a short-range excitonic character is well known. Our results show that it seems to be a sufficient minimal input to understand transport in Bi, and ties in our view with plasmonarons. Generally, in a correlated metal, appearance of mid-infra-red features (e.g. DMFT for one-band Hubbard type models) involves interband transitions between low-energy itinerant and high-energy localized Hubbard-band states. In our case, they arise from transitions involving carrier states scattering off the incoherent and interband excitons involving precisely the renormalized VB and CB in the two-band Hubbard-like model, since $\sigma_{\omega}(\omega)$ in DMFT is just a direct convolution of the one-particle DMFT spectral functions. In Bi, this feature occurs at a very low energy because of the tiny $E_F$. The interband electronic excitations involving carriers coupled to such exciton-like entities have been christened plasmaron in prior work. We find that this characterization is not in conflict with our excitonic fluctuation picture, since we have now shown that it can be well described by analyzing effects of predominantly local and sizable excitonic liquid-like correlations in a quasi-realistic model for Bi. Finally, since excitonic correlations can trigger changes in lattice parameters, whether development of excitonic (liquid) correlations can rationalize the small but unexplained dependence of the rhombohedral parameter $(\omega)$ on $T$ is an interesting to inquire, but out of scope of the present work.

Buoyed by the agreement with the phenomenology so far, we venture to propose a specific model for superconductivity in Bi. To do so, we need to generate an effective pairing interaction arising from excitonic and/or phonon fluctuation exchange. Given the efficacy of the excitonic liquid view above for the normal state, and that the exciton formation energy scale $(\Omega_{exc} \approx \Delta_{exc})$ is quite large, we are clearly in the non-adiabatic limit. This precludes a BCS-like instability to SC in Bi. In the normal state, the interband excitons involving the VB and CB form an isospin $T = 1/2$ degree of freedom, described by $T^+ = \sum_{\alpha} a^\dagger_{\alpha} b_{\alpha}$, $T^- = \sum_{\alpha} b_{\alpha^*} a_{\alpha}$, $T^0 = \sum_{\alpha}(n_{\alpha^*} - n_{\alpha})/2$ in particle-hole space. Precisely as in the $U < 0$ Anderson lattice Hamiltonian, one has a crossover to an incoherent exciton fluctuation dominated regime at $k_B T \approx \Omega(U')$, followed by a second crossover to a quasi-coherent exciton fluctuation dominated regime at a lower exciton-Kondo scale, $T_{Kc} ^\Omega \approx U'(\sqrt{\rho})^{1/2} e^{-1/(2\Omega(U'))}$, where $\rho = \rho(E_F)$ is the LDOS at $E_F$ and $f \approx t_{2g}^2 X_{\Omega}(0)$, with $\chi_{\Omega}(0)$ the excitonic susceptibility at $\omega = E_F(=0)$: correlated FL behavior obtains below $T_{Kc} ^\Omega$, thanks to an excitonic Kondo screening implicit in a situation where a tiny number of carriers are now coupled to fluctuating isospins $T$. It is precisely this resonant scattering that governs transport in the normal state in accord with transport data as found above.

As $T$ is lowered, inter-site, residual interactions develop in full analogy with the way inter-site spin correlations, mediated by metallic carriers, arise in the Kondo lattice. They correspond to exchange coupling between isospins...
above, mediated by the tiny number of carriers, and are naturally viewed as an exciton fluctuation-exchange process. The residual interactions read\(^3\)

\[
H^{(2)}_{\text{eff}} \approx J \sum_{\langle ij \rangle, \sigma, \sigma'} \left[ n_{i\sigma} - a_{i\sigma}^+ b_{j\sigma'} - a_{i\sigma'} b_{j\sigma} - a_{i\sigma} b_{j\sigma'}^+ - a_{i\sigma'} b_{j\sigma}^+ \right],
\]

which is also \(\int \sum_{\langle ij \rangle, \sigma, \sigma'} | \eta_{\alpha} \rangle \langle \eta_{\alpha'} | \).

Absorbing the first term in the normal state Hamiltonian, \(H^{(2)}_{\text{res}}\) in momentum space is

\[
H^{(2)}_{\text{res}} = -\sum_{kpq} \left( a_{kpq}^+ b_{kq} - a_{kpq} b_{kq}^+ \right)
\]

Use of a usual BCS-like argument to derive SC from a static Hartree-Fock decoupling of \(H^{(2)}_{\text{eff}}\) is possible\(^{23}\), but problematic. In particular, this is inappropriate in the case of \(Bi\) for the following reasons. The usual weak-coupling BCS approximation starts with the one-electron Green functions and constructs the “non-interacting” pair (particle-particle, p-p) susceptibility, composed from the lowest order bubble diagram involving the DFT one-electron propagators. Next, one uses this bare p-p susceptibility along with the bare attractive interaction (derived from integrating out appropriate bosons, e.g. phonons) to generate a ladder p-p vertex. The pole of this vertex function determines the SC transition temperature within BCS theory\(^4\). This procedure works well only as long as (i) the normal state above \(T_c\) is well-described by a weakly interacting Landau Fermi liquid, adiabatically connected to a free Fermi gas, and (ii) consequently, no quasi-bound state formation occurs in the normal state. Our results for the normal state in \(Bi\) above, showing good accord with spectral and transport responses, involve pre-formed, dynamically fluctuating excitons co-existing with a small number of carriers already in the normal state. Thus, the normal state in \(Bi\) does not satisfy condition (ii) above, making it mandatory to consider the strong coupling limit of the pairing problem beyond the BCS framework (see, e.g. Pietronero et al\(^{24}\) for the non-adiabatic limit of the strong e-p coupling case).

As it stands, the effective interaction in a fluctuating excitonic “normal” state leads to possibility of both p-p and p-h condensation, and at intermediate-to-strong “excitonic Kondo” coupling, it is known that there is strong interference between both these channels at two-particle level. Then the associated logarithmic divergences appear in both channels, best illustrated within the classic parquet approach\(^{25,26}\). We have adapted Abrikosov’s original parquet approach to our case (see SI for details). We proceed as follows: the bare vertex, \(\Gamma^{(0)}_{\pi_{pq}\pi_{rs}}(p_F, p_F, P_F, p_F) = \frac{i}{\hbar} \delta_{\pi_{pq}, \pi_{rs}} - \delta_{\pi_{pq}, \pi_{rs}}\delta_{P_F, P_F}\) will obtain drastic renormalization in the ‘excitonic Kondo’ regime. To this end, we exploit DMFT results: since the \(\pi\)-band spectral function is gapped, one can replace \(G_{\pi\pi}(\omega) \sim \omega + \epsilon_{\pi}\) at low energy (since \(\text{Im} \Sigma(0) = 0\) at low energy), while the \(\pi\)-band propagator is approximated as \(G_{\pi\pi}(k, \omega) \sim \omega - e^{i k \cdot \rho} \omega_{\pi}\) with \(e^{i k \cdot \rho} = 1 - \left(\frac{d}{d\omega}\text{Re} \Sigma(\omega)\right)_{\epsilon_{\pi}}\). With these inputs, we find that the vertex is logarithmically enhanced, giving \(\Gamma^{(0)}(\omega) = J\left[1 + \rho(\epsilon_{\pi} + \epsilon_{\pi})\right]^{-1}\). The SC transition temperature is estimated from the divergence of the renormalized vertex, and we find \(T_c = E_F e^{-1/\rho}\). Using the renormalized \(E_F \approx T_c \approx 100\) K (6.2 meV) from DMFT results below which correlated FL behavior sets in (instead of the bare \(E_F \approx 23\) meV (270 K)) in \(Bi\), \(\rho(E_F) \approx 0.1\) eV\(^{-1}\) from normal state DMFT results, and the coupling \(J \approx O(1)\) (we are not in the regime \(g_{\pi\pi} \ll U\) for \(Bi\)), we estimate the SC \(T_c \approx O(1)\) meV, quite close to the experimental finding of \(T_c \approx 0.5\) meV. Given the approximations made above, this is very reasonable. Our estimate is a huge enhancement compared to the \(T_c^{\text{BCS}} \approx O(100)\) meV found using the naive BCS formula\(^1\), and reflects the strong coupling nature of SC, where non-adiabatic effects enhance \(T_c\) via huge enhancement of the vertex. Effects akin to the above have been discussed in the non-adiabatic limit of strong electron-phonon coupling\(^{24,27,28}\), and, in \(Bi\), the fact that \(g(E_F) \approx O(0.5)\) will also imply such additional enhancement arising from electron-phonon coupling. In the adiabatic or anti-adiabatic limit, it is possible to subsume this latter effect into a renormalization of the bare vertex \(J\). However, when \(g(E_F) \approx O(1)\), one cannot “integrate out” the phonons. One must include e-p couplings explicitly in such a case and, in fact, our DMFT results do include the renormalization caused by e-p coupling as well. Such strong coupling multi-band SC will also generally give an enhanced \((dH_{c2}/dT)\) in \(Bi\), as in other documented near semi-metallic superconductors\(^30\), relative to the BCS prediction.

Finally, the form of the effective interaction also shows that the excitonic insulator (EI) phase is a subleading instability in \(Bi\). It is of interest to inquire whether evidence for such a “competing order” could obtain in \(Bi\) by modification of its electronic structure, e.g. by pressure\(^1\) or an external magnetic field. Both reduce the tiny carrier density further, jacking up the effective \(U/T\) ratio and inducing tendency to localization. Actually, pressure does induce a metal-insulator-like transition in Bismuth\(^1\). Within our strong coupling analysis, the competing order in the resultant semiconducting phase would be characterized by an order parameter \(\Delta_{\text{EO}} = \sum_{\mathbf{k}} f_\mathbf{k}(\mathbf{k}) c_{\mathbf{k}00} c_{\mathbf{k}00}\) with \(f_\mathbf{k}(\mathbf{k}) = \frac{2\sqrt{3}}{\sqrt{3}} \sin\left(\frac{\sqrt{3}}{2} k_a/2\right)\sin\left(\sqrt{3} k_a/2\right)\). This excitonic order parameter does not break inversion symmetry \((\mathbf{k} \rightarrow -\mathbf{k})\) but, remarkably, is associated with an electronic nematic order\(^27\). Very recent work\(^28\) finds a nematic electronic state with a tiny gap \(O(500)\) meV on the surface of \(Bi\) under high magnetic fields. Whether a Rashba-SOC modified electronic structure as above can induce an excitonic instability as proposed here, and whether such a state can lead to a reconstructed electronic structure having the observed anisotropy of Landau level wave-functions, is an enticing open issue. Thus, future studies under pressure can confirm or refute our prediction which, at this time seems to have limited confirmation\(^{29}\). Investigation into these aspects is left for future work.

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