Failed excitonic quantum phase transition in Ta$_2$Ni(Se$_{1-x}$S$_x$)$_5$

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We study the electronic phase diagram of the excitonic insulator candidates Ta$_2$Ni(Se$_{1-x}$S$_x$)$_5$ ($x = 0, \ldots, 1$) using polarization resolved Raman spectroscopy. Critical excitonic fluctuations are observed that diminish with $x$ and ultimately shift to high energies, characteristic of a quantum phase transition. Nonetheless, a symmetry-breaking transition at finite temperatures is detected for all $x$, exposing a cooperating lattice instability that takes over for large $x$. Our study reveals a failed excitonic quantum phase transition, masked by a preemptive structural order.

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Introduction. One of the fascinating manifestations of interactions between electrons in solids is the emergence of electronic orders. The fluctuations close to the respective quantum critical points are also believed to be the drivers of a wealth of yet unexplained behaviors, including strange metallicity and high-$T_c$ superconductivity [1–4]. In many cases (such as, e.g., nematic [5,6] or density-wave [7,8] orders) electronic order breaks the symmetries of the crystalline lattice and the corresponding transitions can be, symmetry-wise, identical to structural ones. This raises the question of the role of the interplay between electronic and lattice degrees of freedom in the ordering. Even in cases where the lattice only weakly responds to the transition [9,10], the critical temperatures [11,12] and quantum critical properties [13] can be strongly modified. Moreover, in a number of cases the origin of the order is still under debate [14–16], as the lattice may develop an instability of its own.

The electronic-lattice dichotomy has recently come to the fore in studies of Ta$_2$NiSe$_5$ [17–19]—one of the few candidate materials for the excitonic insulator (EI) phase [16,20–24]. The EI results from a proliferation of excitons driven by Coulomb attraction between electrons and holes in a semiconductor or a semimetal [25–29]. Ta$_2$NiSe$_5$ exhibits a phase transition at $T_c = 328$ K; while the pronounced changes in electronic structure [30,32] and optical [30] properties are consistent with the ones expected for an EI, they allow an alternative interpretation in terms of a purely structural phase transition [31–34]. Indeed, the EI state in Ta$_2$NiSe$_5$ is expected to break mirror symmetries of the lattice due to the distinct symmetries of the electron and hole states forming the exciton [35], similar to a structural transition [31]. Intriguingly, substitution of Se with S has been shown to suppress $T_c$ in transport experiments to zero [19], suggesting a possible quantum phase transition (QPT) at $x = x_c$ in Ta$_2$Ni(Se$_{1-x}$S$_x$)$_5$.

Increasing $x$ enhances the band gap in the electronic structure [36], which is known to suppress the EI [27,28], consistent with an EI QPT. On the other hand, the lattice degrees of freedom also evolve with $x$ making it imperative to separately assess the roles of electronic and lattice degrees of freedom throughout the phase diagram of Ta$_2$Ni(Se$_{1-x}$S$_x$)$_5$.

Experimentally, this is a challenging task. As transitions caused by the lattice and electronic degrees of freedom break the same symmetries [31,35], their signatures may appear identical in thermodynamic (e.g., specific heat [19]) and symmetry-sensitive (x-ray diffraction [17]) probes, as well as in the single-electron spectra [18,32]. Probing the collective dynamics out of equilibrium could provide more information [33,34,37], but raises the question of whether the nonequilibrium state preserves the interaction between the structural and electronic modes intact [38]. Finally, due to the even-parity nature of the critical mode [39], the dipole selection rules forbid direct observation of the order parameter response in optical absorption [30]. A promising technique to address the near-equilibrium collective dynamics is polarization-resolved Raman scattering [39–41] that also allows us to detect symmetry breaking independently [42–44]. Furthermore, analysis of the Raman data [38,39] enables us to deduce the individual contributions of the lattice and electronic modes to the transition, making this technique unique in its scope.

In this Letter we use polarization-resolved Raman scattering to study the dynamics of electronic excitations throughout the phase diagram of Ta$_2$Ni(Se$_{1-x}$S$_x$)$_5$. We reveal the presence of low-energy excitonic modes that soften on cooling towards $T_c(x)$. This softening indicates that in the absence of lattice effects, a purely excitonic transition would have taken place at $T_c(x)$, which we deduce to be smaller than $T_c(x)$. On increasing sulfur content $x$, $T_c(x)$ is suppressed to negative values and for $x = 1$ low-energy excitons are no longer observed, as expected for an excitonic insulator quantum phase transition. However, the actual $T_c(x)$ remains finite for all $x$, implying the presence of a cooperating lattice instability, obscuring the suppression of the excitonic order. The study thus reveals a “failed” excitonic quantum phase transition in...
FIG. 1. Overview of the polarization-resolved Raman response $\chi''(\omega, T)$ in $\text{TaNi(Se}_{1-x}\text{S}_x)\text{S}_5$. (a)–(d) Response in $aa$ polarization geometry corresponding to fully symmetric ($A_g$) excitations above (red) and below (blue) $T_c$ for $x = 0–1$. Shading highlights the bare electronic contribution to the response. (e) and (f) Same for $ac$ geometry, probing the excitations with the symmetry of the order parameter ($B_{2g}$) for $T > T_c$ (red). Unlike $aa$ geometry, phonons show an extremely anisotropic Fano line shape (hatching), indicating their strong interaction with the electronic continuum (red shading). For $T < T_c$ (blue) excitations observed in $aa$ geometry above $T_c$ appear (arrow) due to symmetry breaking. (i)–(l) Temperature dependence of $\chi''(\omega, T)$; an enhancement at low energies near $T_c$ (arrow) is observed for $x = 0, 0.25, 0.67$. For $x = 1$ no low-energy response is present. (m)–(p) Details of $\chi''(\omega, T)$ for regions marked by dashed lines in (i)–(k). (m)–(o) Fano line shape of low-energy phonons due to interaction with electronic continuum. (p) High-energy peak due to an uncondensed exciton in $\text{TaNiS}_5$.

$\text{TaNi(Se}_{1-x}\text{S}_x)\text{S}_5$ masked by a preemptive structural order that takes over as the electronic instability is suppressed.

**Experiment.** We performed Raman scattering experiments on single crystals $\text{TaNi(Se}_{1-x}\text{S}_x)\text{S}_5$ with varying Se/S content, grown using the chemical vapor transport (CVT) method [38,45]. The measurements were performed in a quasi-backscattering geometry on samples cleaved to expose the $ac$ crystallographic plane with the 647 nm line from a Kr$^+$ ion laser excitation, details presented in Ref. [38]. The selection rules in the high-temperature orthorhombic (point group $D_{2h}$) phase imply that $ac$ polarization geometry probes excitations with $B_{2g}$ symmetry (same as that of the order parameter), while $aa$ geometry probes the fully symmetric $A_g$ ones [38–41]. Below $T_c$, the point group symmetry is reduced to $C_{2h}$ and the two irreducible representations merge, such that excitations from $ac$ geometry above $T_c$ may appear in $aa$ geometry and vice versa. Their appearance allows us to determine $T_c$ from the Raman spectra.

**Data overview.** Summarized temperature dependence of the Raman susceptibility $\chi''(\omega, T)$ is presented in Fig. 1. The samples with $x = 0, 0.25, 0.67$ show qualitatively similar spectra. At low energies, phonon peaks are observed on top of a smooth background, which we attribute to electronic excitations. On cooling, a pronounced redistribution of electronic intensity in a wide range of energies is observed, leading to a formation of a gaplike suppression followed by a high-energy feature, Figs. 1(a)–1(c). This feature at 380 meV for $\text{TaNiSe}_5$ has been attributed to the coherence factors at the gap edge of an Eli [39]. In $ac$ geometry, a pronounced enhancement at low energies is evident close to $T_c$, consistent with critical mode softening near a second-order phase transition, Figs. 1(i)–1(k). In the same temperature region, the line shapes of the low-energy phonons show strongly asymmetric Fano form [Figs. 1(m)–1(o)]—a known signature of interaction with an electronic excitation continuum [38,39,46]. This indicates the presence of low-energy symmetry-breaking electronic
excitations that soften close to $T_c$. At low temperatures, the asymmetry disappears [Figs. 1(e)–1(g)], a behavior consistent with a gap opening in an EJ. On increasing sulfur content $x$, the temperature where the strongest low-energy enhancement is observed progressively lowers [Figs. 1(i)–1(k)], and the $A_x$-symmetry feature at about 380 meV moves slightly to lower energy and becomes less pronounced.

The signatures for $\text{Ta}_3\text{NiS}_5$ ($x = 1$) are rather different: in the $ac$ geometry, low-energy electronic excitations are absent at all temperatures, indicating the presence of a direct gap. This implies that between $x = 0.67$ and $x = 1$ the electronic structure undergoes a Lifschitz transition from a semimetallic to an insulating one. The intensity in the $aa$ geometry at low energies is also pronouncedly smaller than for the other samples and no broad high-energy peak is observed at low temperatures. On the contrary, a sharp feature at about 0.3 eV emerges in $ac$ geometry on cooling below 100 K.

**Symmetry-breaking transition.** We address first the presence of a phase transition by studying the appearance of new modes in the broken-symmetry phase, as outlined above. In Fig. 2(a) we show the temperature dependence of such a “leakage” phonon intensity marked by arrow in Figs. 1(e)–1(h). One can see the appearance of the leaked intensity below $T_c$ in the pure Se case, as well as the decrease of $T_c$ with $S$ doping. Leakages of other modes appear below the same temperature $T_c$ [38,45]. At low $x$ the obtained values of $T_c$ agree with the ones deduced from transport and specific heat measurements [19,38], Fig. 3. However, in contrast to the transport data reported in Ref. [19], we find that the symmetry-breaking transition persists for all compositions, although the leakage intensity is strongly suppressed with higher $x$. The latter suggests that the phase transition signatures in thermodynamic and transport measurements may become too weak to be observed at large $x$, especially since the system becomes more insulating with $x$. The structural signatures, e.g., the deviation of the monoclinic angle $\beta$ from 90°, should be also strongly suppressed, being already weak at $x = 0$ [17].

The phase transition for $x = 1$, where no low-energy softening is observed [Fig. 1(l)], indicates a different transition mechanism. Below we analyze our data to elucidate the origin of the transition as a function of $x$.

**Electronic contribution to the phase transition.** We investigate first the soft-mode behavior observed for $x \leq 0.67$ [Figs. 1(i)–1(k)]. In particular, we analyze the asymmetric line shapes of the low-energy part of $\chi^\prime\prime(\omega, T)$ around $T_c$ [Figs. 1(m)–1(o)] using an extended Fano model [38,39]. The model assumes three phononic oscillators (which is the number of $B_{2g}$ modes in the orthorhombic phase) interacting with a continuum of excitonic origin. The latter is expected to arise from the excitonic fluctuations in a semimetal, overdamped due to the allowed decay into particle-hole pairs. Close to the transition, the dynamics of the excitonic mode is governed by the time-dependent Landau equations [47–50]. Together with the standard oscillator dynamics of the phonons the system is described by

$$\{\partial_t + \Omega_x(T)\} \varphi + \sum_{i=1}^{3} \bar{v}_i \eta_i = 0,$$

$$\{\partial_t^2 + 2\gamma_x(T)\partial_t + \omega_p^2(T)\} \eta_i + \bar{v}_i \varphi = 0,$$

where $\eta_{i=1,2,3}$ and $\varphi$ are the collective coordinates (order parameters) of the optical phonons and excitons, respectively. $\Omega_x(T)$ is the characteristic energy of the excitonic fluctuations, $\omega_p(T)$ and $\gamma(T)$ are the phonon frequencies and scattering rates, and a bilinear exciton phonon-coupling $\bar{v}_i$ is assumed. The linear response of the system Eq. (1) determines the Raman susceptibility. The resulting model [45] is a generalization of the standard Fano model [46] for Raman scattering in metals to the case of three phonons and the continuum response determined from the Landau equation, Eq. (1). The purely excitonic part of the response has then the form of a broad continuum $\chi^\prime\prime_{\text{cont}}(\omega, T) \propto \omega^{-1}$, in contrast to the Lorentzian phonon peaks. The interaction between the phonons and the excitonic continuum leads to an asymmetric broadening of the peaks [38], allowing us to capture the observed line shapes in great detail [38,39,45].

We now discuss the parameters deduced from the Fano model fits. The phonon frequencies $\omega_p(T)$ do not soften near $T_c$ [38], ruling out a zone-center phonon instability [31]. On the other hand, $\Omega_x(T)$ [Fig. 2(b), solid lines] consistently
softens above $T_c$ for all semimetallic samples. The linear temperature dependence $\Omega_e(T) \sim T - T_{ex}$ implies that a purely electronic transition would have taken place at $T_{ex} < T_c$ for $x = 0, 0.25$ (Fig. 3, blue symbols). The strongly negative $T_{ex}$ for $x = 0.67$ indicates that the exciton softening alone would not have led to a transition at this sulfur concentration.

The suppression of the excitonic instability with $x$ is even more evident in Ta$_2$NiSe$_5$ [Figs. 1(d), 1(h) and 1(l)], where the low-energy electronic response is altogether absent due to a direct band gap [36]. Instead, we observe a sharp $B_{2g}$ symmetry mode at 0.3 eV [Fig. 1(p)], consistent with an in-gap exciton. It is followed by a weaker feature at 0.325 eV and an intensity “tail” at higher energies up to around 0.4 eV. The natural interpretation of the second peak is the second state of the Rydberg series (i.e., 2S exciton), while the high-energy tail can be attributed, in analogy with optical absorption spectroscopy, to the Rydberg states of higher order and interband transitions [51,52] with possible contributions from phonon-assisted exciton transitions [53,54]. A leakage of the exciton features is also observed in $aa$ geometry due to symmetry breaking, Fig. 1(d) [45]. On heating, all the features broaden and eventually smear out above 100 K. The increase of the linewidth of the excitonic features can be attributed to the interaction with acoustic and optical phonons [55].

**Ferroelastic transition in Ta$_2$NiSe$_5$.** The presented observations show that the excitonic instability on its own cannot explain the occurrence of a transition for samples with large $x$, calling for a more careful consideration of the lattice effects. The most vivid is the case of Ta$_2$NiS$_5$, where the excitonic response is confined to high energies [Fig. 1(p)]. Three $B_{2g}$ optical phonon modes [Fig. 1(l)], on the other hand, exhibit some (around 15% maximum) softening on cooling. However, their energies never soften below 6.5 meV, nor exhibit anomaly at the transition temperature 120 K. We note that the number of $B_{2g}$ modes is restricted to three by the space group of the orthorhombic Ta$_2$NiS$_5$, implying the absence of any other $B_{2g}$ optical modes beyond those shown in Fig. 1(l). Consequently, an instability of zone center phonons in Ta$_2$NiS$_5$ [56] is ruled out by the data.

The only remaining option for the transition origin in Ta$_2$NiS$_5$ is an instability of the acoustic modes, i.e., ferroelasticity [57,58], driven by softening of the $B_{2g}$ shear modulus $C_{ac}(T)$. Indeed, the acoustic modes are not observed directly in Raman due to their extremely low energies and weak coupling to light [38,59]. However, we will show now that the effects of the ferroelastic instability can be observed at $x < 1$ via its coupling to the low-energy excitons.

**Electronic-structural phase diagram.** We will now demonstrate that the entire phase diagram of Ta$_2$Ni(S$_1-x$Se$_x$)$_5$ can be understood by including the interaction between excitonic and lattice modes. As has been noted above, the bare excitonic transition temperature $T_{ex}$ (Fig. 3, blue line) is significantly lower than the actual $T_c$. However, even the coupling of excitons with the otherwise inert optical phonons can affect the transition temperature. For a coupled excitonic-optical phonon system, the transition temperature $T_{comb}(x)$ corresponds to the appearance of a zero-energy solution of Eq. (1) deduced from [45]:

$$\Omega_e[x, T_{comb}(x)] - \sum_i \frac{\tilde{v}_i^2}{\omega_{pi}^2[x, T_{comb}(x)]} = 0,$$

where all the parameters of this equation are deduced from the Fano analysis of the Raman data, Figs. 1(m)–1(o), following Ref. [38]. The resulting temperature $T_{comb}(x)$ is shown in Fig. 3 (green line). While higher than $T_{ex}(x)$, there is still discrepancy with $T_c$: for example, $T_{comb}(x = 0.67)$ is negative, while the actual $T_c(x = 0.67)$ is 170 K.

We now include the effects of coupling of the excitonic order parameter $\varphi$ to the $B_{2g}$ strain $\varepsilon_{ac}$ (acoustic modes). A linear coupling between the two is allowed by symmetry [11,12,39] and leads to a further modified equation for the transition temperature [39,45]:

$$\Omega_e[x, T_{comb}(x)] - \sum_i \frac{\tilde{v}_i^2}{\omega_{pi}^2[x, T_{comb}(x)]} - \lambda^2 / 2 C_{ac}^{-1}(T_{comb}(x)) = 0,$$

where $\lambda$ and $C_{ac}(T)$ is the strain-exciton coupling constant and the $B_{2g}$ is shear modulus, respectively. To capture the ferroelastic instability at $x = 1$, we assume a Curie-Weiss behavior of the shear modulus $C_{ac}^{-1}(T) = C_{ac}^{-1}(0) - \frac{a}{T - T_{0}}$. Using $\lambda^2 C_{ac}^{-1}(0)$ and $\lambda^2 a$ as fitting parameters, the observed $T_c(x)$ can be captured very accurately, see the black line in Fig. 3. Importantly, the effects of the ferroelastic softening become noticeable well before $x = 1$. The purple dashed line $T_{str}(x)$ in Fig. 3 shows the transition temperature $T_{str}(x)$ obtained ignoring ferroelastic softening [i.e., taking $C_{ac}^{-1}(T) = C_{ac}^{-1}(0)$ in Eq. (3)]. The result deviates strongly from actual $T_c(x)$ already at $x = 0.67$. Continuing the trend further suggests a complete suppression of ordering at
$x_\varepsilon \approx 0.8$ in the absence of ferroelasticity. At the same time, for low $x$, $T_{\text{crit}}^{\text{fer}}(x)$ and $T_{\text{comp}}^{\text{fer}}(x)$ are almost indistinguishable, suggesting that ferroelasticity does not play a role in that case.

This picture bears important consequences for the physics of Ta$_2$Ni$(\text{Se}_{1-x}\text{S}_x)$. At low $x$, the transition is driven, to a good approximation, only by the excitonic softening. On increasing $x$, the lattice softening becomes more important, and for $x = 1$ the transition is purely ferroelastic. In the absence of ferroelasticity, an electronic QPT would have occurred at $x_\varepsilon \approx 0.8$. We remind that the excitonic and lattice orders break the same symmetries in Ta$_2$Ni$(\text{Se}_{1-x}\text{S}_x)$. Consequently, no change in symmetry occurs as a function of $x$ at low temperatures and a true QPT at $T = 0$ is avoided (unlike the case of superconductivity emerging near quantum critical points). However, this does not preclude critical electronic fluctuations, associated with the “failed” QPT at $x_\varepsilon \approx 0.8$ to be observed at sufficiently high temperatures (higher than the bare lattice transition temperature of 120 K) [13].

The presence of quantum critical fluctuations due to a failed excitonic QPT lends a natural explanation to the signatures of strong correlations observed in Ta$_2$NiSe$_5$. In particular, a filling-in, rather then closing of the gap in the excitonic band structure, the EI QPT has been predicted to lead to non-Fermi liquid behavior [63], mass enhancement [64], or emergence of superconductivity [65]. Interestingly, a superconducting dome near the end point of the monoclinic phase has been recently reported in Ta$_2$NiSe$_5$ under pressure [66].

**Conclusions.** In this work we used polarized Raman scattering to study the phase diagram of the excitonic insulator candidates Ta$_2$Ni$(\text{Se}_{1-x}\text{S}_x)$, disentangling the roles of structural and electronic ordering. We revealed a failed excitonic insulator quantum phase transition at $x_\varepsilon \approx 0.8$, avoided due to a lattice instability below 120 K. At low sulfur content $x$ we observed a soft excitonic mode driving the transition, while at large $x$ this mode ultimately transforms into a high-energy exciton, unable to drive the transition. We further exclude the instability of optical phonons and demonstrate that a ferroelastic instability yields an explanation of the observed symmetry-breaking transition. While the excitonic quantum phase transition is avoided due to the low-temperature lattice instability, the associated critical fluctuations can still be present at high temperatures [13], explaining the correlation effects in Ta$_2$NiSe$_5$. Furthermore, selective control of the structural and excitonic instability by strain [62] or pressure [19] can turn Ta$_2$Ni$(\text{Se}_{1-x}\text{S}_x)_{\varepsilon}$ into a platform to study the excitonic QPT at low temperatures as well as quantum critical ferroelasticity [67].

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