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Photoinduced renormalization and electronic screening of quasi-two-dimensional Dirac states in BaNiS$_2$

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By means of time- and angle-resolved photoelectron spectroscopy, we give evidence of a remarkable reduction of the Fermi velocity of out-of-equilibrium Dirac bands in the quasi-two-dimensional semimetal BaNiS$_2$. This effect is accompanied by a nonrigid shift of the bands at the center of the Brillouin zone. Analytical and first-principles calculations indicate that this band renormalization is ascribed to a change in nonlocal electron correlations driven by a photoinduced enhancement of screening properties. Our results are explained by a scaling relation between inverse screening length and electronic temperature that is of general relevance for the description of out-of-equilibrium dynamics and screening effects in all layered Dirac materials.

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I. INTRODUCTION

The recent discovery of Dirac and Weyl fermions in crystalline solids [1–5] provides a favorable experimental environment for the study of these exotic electronic states and for the development of novel electronics applications. For instance, the interaction of Dirac fermions with ultrafast light pulses is attractive in view of novel concepts of devices combining unprecedented mobility and subpicosecond response or of enabling photoinduced topological phase transitions [6–8]. The electronic band structure of these materials is characterized by several linearly dispersing bands that originate from and are protected by the interplay between real- and momentum- space properties [9–12]. Therefore, the ability to manipulate these bands is a prerequisite for any functional application of relativistic Dirac fermions.

In the present paper, we demonstrate both experimentally and theoretically that Dirac bands can be renormalized using ultrafast light pulses in the quasi-two-dimensional (quasi-2D) Dirac semimetal BaNiS$_2$. With time-resolved angle-resolved photoemission spectroscopy (trARPES), we show that photoexcitation significantly reduces the Fermi velocities of Dirac electrons. This remarkable behavior is also accompanied by nonrigid, orbital dependent band shifts at the center of the Brillouin zone [13]. Both these effects are captured by our ab initio calculations, demonstrating that the key underlying mechanism is the dynamic reduction of the electron interaction range upon photoexcitation. With the help of an analytical model we show that the screening enhancement is related to the temperature increase of excited electrons, so that our results provide a framework for a general description of the effects of screening in quasi-2D Dirac materials.

The ultrafast renormalization of the band structure has so far been investigated only in insulating phases [14–17] or in thin films where the substrate significantly affects the screening properties [18,19]. The present study on BaNiS$_2$ demonstrates that ultrafast light pulses alter dramatically the fermiology and the transport properties of Dirac semimetals.

II. ARPES EVIDENCE OF ULTRAFAST RENORMALIZATION OF DIRAC STATES

In our experiment, BaNiS$_2$ single crystals were cleaved in ultrahigh vacuum on the $a$-b plane at the temperature of 130 K. The trARPES measurements were performed by pumping with 1.55-eV infrared pulses and by probing photoelectrons with 6.28-eV ultraviolet pulses [20] [see Fig. 1(a)]. The time resolution is 80 fs. The equilibrium Fermi surface consists of a fourfold-symmetric Dirac cone along $\Gamma$-M, and by a small electron pocket (EP) at $\Gamma$. Both structures are visible in the top panel of Fig. 1(b). The d-orbital nature of each band [13] is indicated in the figure legend. The photoexcited band structure is shown in the bottom panel for $\Delta t = 250$ fs.

In Fig. 1(c) we present the energy distribution curves (EDC’s) relative to the Dirac cone: by fitting the data with a Fermi-Dirac distribution, one can extract the electronic temperature, which yields a value $T_F \approx 400$ K at $\Delta t = 200$ fs.
FIG. 1. (a) Experimental geometry of the trARPES setup. (b) Top: Reference spectra of the bands along $\Gamma - M$ (Dirac cone at $k_f \approx -0.5 \, \text{Å}^{-1}$ and electron pocket at $\Gamma$). Bottom: Photoexcitation of the electronic states at a 250-fs delay with a pump fluence of 0.2 mJ/cm². (c) Integrated EDC’s around $k_f \approx -0.5 \, \text{Å}^{-1}$ corresponding to the Dirac states: dashed lines show their photoemission spectral weight for negative and positive time delays. Solid lines are fits giving the corresponding electronic temperature (130 K at negative delays; 400 K at 200 fs). Intensities are plotted in logarithmic scale.

We first focus on the ultrafast dynamics of each Dirac band by tuning the light polarization, as described elsewhere [13,21]. Figures 2(a) and 2(b) show the reference spectra of the $dz^2$ and $dx^2 - y^2$ bands, respectively. Figures 2(c) and 2(d) show a series of difference trARPES spectra corresponding to the difference between positive and negative delay images with a pump fluence of 0.2 mJ/cm². The difference spectra show an intensity gain (red) above the Fermi level due to the photoexcitation of electrons to the unoccupied states. As illustrated in Fig. 2(g), these states reach a maximum of occupation around 250 fs due to electron scattering and impact ionization [22]. The subsequent decay dynamics occurs with a timescale of 500 fs. We now discuss the evolution of the valence bands below the Fermi level shown in Figs. 2(c) and 2(d). These states display two distinct features: (i) an expected intensity loss (shown in blue) due to a transient depletion of the electron population and (ii) a remarkable intensity gain (shown in red), a signature of an ultrafast and time-dependent renormalization, i.e., a nonrigid shift, of the Dirac states. In order to study the dynamics of the band dispersion, $E(k,t)$, for each $\Delta t$ we analyzed the momentum distribution curves (MDC’s) of the ARPES yield, as reported in Appendix A. Figures 2(e) and 2(f) depict the band dispersion of the $dz^2$ and $dx^2 - y^2$ band, respectively, before ($-100$ fs) and after ($+250$ fs) the arrival of the pump pulse. We clearly observe an energy-dependent band shift corresponding to a reduction of Fermi velocity. This renormalization is purely photoinduced and is different from a thermal effect (see Appendix A). The renormalization is also qualitatively different from the effects of chemical substitution, even though the available data allow a comparison only with heavy Co doping [21]. In the right panel of Figs. 2(e) and 2(f), we plot the band renormalization, e.g., the change in the wave vector at the given binding energy, for different time delays. Note that the nonrigid shift of the wave vector is larger as one moves away from the Fermi level; $\Delta k$ even exceeds 0.01 Å⁻¹ for the $dx^2 - y^2$ band. In contrast, within our experimental resolution, the Fermi wave vector does not exhibit any shift and remains in its equilibrium position for all time delays. Figure 2(h) shows the dynamics of the Fermi velocity of both bands with a large reduction up to 30% for $\Delta t \sim 250$ fs. To the best of our knowledge, such

FIG. 2. ARPES reference image for the $dz^2$ (a) and $dx^2 - y^2$ (b) bands. Difference trARPES images (after photoexcitation minus before photoexcitation) for each linear band are shown in (c) and (d): red and blue in the color scale indicate gain and loss of photoemission yield, respectively. Panels (e) and (f) show the band dispersion $E(k)$ of the $dz^2$ and $dx^2 - y^2$ bands before and after arrival of the pump pulse, respectively. The photoinduced band renormalization for different time delays is depicted on the right: the curves are obtained by subtracting the $E(k)$ curves at the negative delay from the $E(k)$ curves at positive delays. In both cases, the photoinduced renormalization effect reaches a maximum around 250 fs. (g) Dynamics of the excited-state photoemission yield, integrated above the Fermi level. The decay takes place with a time constant of 500 fs. (h) Dynamics of the Fermi velocity for the linearly dispersing bands.
dramatic photoinduced renormalization of the band structure in a semimetal, far from any electronic or structural phase transition, has not been reported before. The temperature $T_0$ of hot electrons, displayed in Fig. 3(a), is deduced by analyzing the evolution of the spectral weight of the Dirac states, as described in Ref. [23].

III. THEORY OF TEMPERATURE-DEPENDENT SCREENING IN A LAYERED-DIRAC MATERIAL

Recent out-of-equilibrium band-structure calculations [24–27], applied so far only to rather simple systems, illustrate that the electronic changes induced by a light pulse are as follows; (i) change of band occupancies, (ii) band shifts, and (iii) electronic screening variations. The latter feature arises from the increase of screening on the electronic bands. The original HSE06 value ($\omega = 0.108 \alpha_0^{-1}$) is valid for the system at equilibrium (black lines). In the photoexcited regime with $0.2 \text{ mJ/cm}^2$ prior to relaxation, $\omega$ falls approximately in between $0.3 \alpha_0^{-1}$ (orange lines) and $0.6 \alpha_0^{-1}$ (red lines).

\begin{equation}
\omega_{\text{Dirac}} = \sqrt{4 \log(2) \frac{e^2 g_S g_C}{k_B^2 \gamma^2 I_d^2} k_b T_e},
\end{equation}

where $g_S$ is the spin multiplicity, $g_C$ the number of cones in each layer, $\kappa_0$ the dielectric constant of the medium, $\gamma$ the Fermi velocity, and $I_d$ the interlayer distance. Thus, by virtue of the reduced phase space of the Dirac band structure, $\omega_{\text{Dirac}} \propto \sqrt{k_B T_e}$, which strongly differs from the conventional Debye-Hückel behavior. For BaNiS$_2$, one obtains [29] the function plotted in Fig. 3(b). This relates $T_e$ of the photoexcited electrons—measured as a function of the time delay [Fig. 3(a)]—to the effective $\omega_{\text{Dirac}}$ in the material. Therefore, the screening enhancement is maximum when unoccupied states close to the Fermi level are maximally populated, then it progressively decays as the system relaxes to equilibrium. This dynamics can be mapped into equilibrium ab initio calculations, provided the screening parameters are modified accordingly. This approximation is justified in presence of a slow relaxation rate as indeed observed in the Dirac states [Fig. 2(g)]; the electrons are already thermalized before the band renormalization reaches its maximum. To implement these calculations, the Heyd, Scuseria, and Ernzerhof (HSE) functional [30–32] is particularly convenient, for it modifies the Perdew, Burke, and Ernzerhof (PBE) [33,34] exchange-correlation ($xc$) functional by the addition of the screened interaction. This interaction is treated at the Fock level ($E_{xc}^{\text{HSE}}$), such that the resulting functional reads as $E_{xc} = E_{xc}^{\text{PBE}} + \alpha (E_{xc}^{\text{HSE}} - E_{xc}^{\text{PBE}})$, where $\alpha$ is the screening parameter. In our case $\alpha = 0.07$, as determined in BaNiS$_2$ by comparison with the experimental measurements [35]. The screened interaction is written as $V_{xc}^{\text{HSE}} = \text{erfc}(\omega r)/r$, where erfc is the complementary error function. For the system at equilibrium, $\omega = 0.108$ in atomic units, i.e., the HSE regular value. In the case of a photoexcited system, $\omega$ increases and reaches its maximum when photoexcited electrons reach their maximum temperature as plotted in Fig. 3(b).

We performed ab initio fully relativistic calculations using the modified HSE functional for $\alpha = 0.07$ and $\omega = (0.108, 0.2, 0.3, 0.6)$ implemented in the QUANTUM ESPRESSO package [36,37]. Details of the calculations are explained in Appendix C. The calculated low-energy band structure along the $\Gamma-M$ direction is shown in Fig. 3(c). The full bandwidth is reduced by the screening enhancement, which contrasts the well-known tendency of the Coulomb exchange to widen the bandwidth [38]. According to our calculations, photoexcitation reduces the Fermi velocities by $\approx10\%$, in both Dirac cone branches of $d_{x^2-y^2}$ character. Experimentally, the Fermi velocity reduction is even larger and reaches $30\%$ [Fig. 2(h)] for both sides of the cone. This discrepancy is attributed to the dynamical nature of photoexcitation and to further correlation effects beyond the screened exchange, neglected in our scheme.
FIG. 4. (a) The ARPES spectrum before the arrival of the pump obtained with s-polarized light. The right panel shows the corresponding EDC’s for several wave vectors. Arrows indicate the contribution of each of the three bands to the total intensity. We follow the binding energy of the EP and band 1 for each wave vector at different time delays. In (b) we display the band dispersion of these bands before and after photoexcitation. The solid lines are guides to the eyes. The electron pocket shows a larger shift toward lower energies with respect to band 1. (c) Photoemission intensity integrated in a k interval of 0.2 Å⁻¹ around Γ₁. The arrow shows the direction of the shift for the electron pocket that corresponds to an increase of screening effects. This is reproduced by modified HSE calculations and depicted in the right panel for two different inverse screening lengths. (d) The maximum photoinduced shift of the EP and the adjacent band (band 1) are shown as a function of the pump fluence. The error bars are estimated from the experiments. The gap separating the EP and band 1 increases with excitation density. (e) Dynamics of the center of the electron pocket for various fluences.

IV. OUT-OF-EQUILIBRIUM DYNAMICS OF THE BAND STRUCTURE

In order to further verify the agreement between theory and experiment, we analyze the expected behavior of the band structure around Γ₁ upon photoexcitation [Fig. 3(c)]. In the enhanced screening regime, the electron pocket at Γ₁ disappears as the generating band is shifted above the Fermi level. This subtle effect depends on the combination of enhanced screening and spin-orbit coupling, which splits the set of three bands just below the Γ₁ point in the system at equilibrium. The bottom most band is the least affected by the screening modification and keeps almost the same position in energy.

The experimental results presented in Fig. 4 confirm that photoexcitation affects not only the velocity of the Dirac states but also the energy levels of the bands at Γ₁. Figure 4(a) shows the ARPES spectrum at Γ₁ along the Γ₁-M direction taken with s polarization. We trace the dispersion of the bands for different time delays by analyzing the EDC’s, Fig. 4(a). Three bands contribute to the photoemission intensity. We label these bands located below the Fermi level in order of decreasing binding energy as bands 2 and 1, located below E_F, and EP, the electron pocket at Γ₁. In Fig. 4(b), we display the band dispersion for the electron pocket and its neighboring band 1 for the negative delay and 250 fs. The larger shift of the electron pocket towards lower binding energies points to a nonrigid shift of the band structure at the Γ₁ point. The pocket also shows a significant renormalization with photoexcitation, as reported in Appendix D. We now analyze in detail this nonrigid shift, by focusing on the integrated EDC’s in a region of 0.2 Å⁻¹ around Γ₁, as shown in Fig. 4(c). We clearly see that the shift of the band structure at Γ₁ matches well our modified HSE calculations (shown in the right panel) that take into account the influence of screening. Figure 4(d) displays the maximum of the photoinduced energy shift occurring at the delay of 250 fs. Note that the maximum of ΔE for the electron pocket reaches about 11 meV for a fluence of 0.2 mJ/cm² and 36 meV for 0.5 mJ/cm² [Fig. 4(e)]. The dynamics of the band structure reveals that while the electron pocket is highly influenced by the ultrafast photoexcitation, band 2 hardly exhibits any appreciable shift. Clearly, our experimental data give evidence of a nonrigid and fluence-dependent energy shift of the bands at the Γ₁ point, further confirming the validity of our scaling relation for the inverse screening length and of our ab initio approach. It is worth emphasizing that our description of screening effects has general validity for all quasi-2D systems presenting linearly dispersing bands: in particular, while Eq. (1) is not applicable to free-standing monolayers, it is valid for all layered materials, as long as an electronic temperature can be defined. For instance, in pump-probe studies of graphite [39] and of ZrSiSe [40], the analysis of the band-gap renormalization dynamics indicated electron-electron interactions as the origin of the gap shrinkage. This interpretation can be easily rationalized based on our screening relation with temperature, which is thus relevant not only for trARPES experiments, but in general for all out-of-equilibrium studies of materials with linearly dispersing bands.
V. CONCLUSION

In conclusion, we observed that ultrafast light pulses dramatically modify the electronic band structure of the quasi-2D Dirac semimetal BaNiS$_2$. Notable effects are a reduction of the band velocity of the Dirac states and a nonrigid shift of the bands at the center of the Brillouin zone. These effects are theoretically explained by a dynamical change of the screening length of nonlocal interactions. While particularly important for the description of ultrafast dynamics in photoexcited materials, our scaling relation with temperature presents general relevance for the interpretation of screening effects in quasi-2D Dirac systems.

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APPENDIX A: PHOTOINDUCED VS THERMAL EFFECTS ON THE DIRAC STATES

In this Appendix, we first show the analysis of the Dirac state renormalization based on MDC’s. We then prove that this renormalization is different from a pure thermal effect and is triggered when electrons are fully decoupled from the lattice.

In order to study the chronological evolution of the band dispersion, we have sliced the ARPES yield for each time delay into MDC’s, Figs. 5(b) and 5(e). The photoemission intensity peak was fitted by a Gaussian with a linear background. The center of the Gaussian shows the wave vector for each binding energy. In this way, the band dispersion can be reconstructed for each time delay $E(k, t)$. By stacking all of the $E(k, t)$ curves a 2D map is acquired where the intensity of each point shows the wave vector position in the $E−t$ plane, Figs. 5(c) and 5(f).

We now compare the photoinduced and thermal effects on the photoemission yield of the $dz^2$ and $dx^2−y^2$ bands, Fig. 6. The ARPES intensities are obtained by $k$ integration around the linearly dispersing bands. The thermal induced changes are acquired by heating the sample starting from 130 K which is the base temperature for the time-resolved experiments. We observe that only at about 4 ps, the electronic and lattice temperatures have converged. This delay corresponds to $\Delta T \sim 20$ K which corresponds to the temperature increase produced by the heat deposited by laser pulses at 0.2 mJ/cm$^2$. These data show that the renormalization of the band structure at short delays (around 200 fs) discussed in the paper cannot be explained by a thermal effect and is purely photoinduced.

APPENDIX B: RELATION BETWEEN ELECTRONIC TEMPERATURE AND SCREENING LENGTH IN LAYERED DIRAC MATERIALS

For an electronically excited regime, we would like to infer its impact on the screening length of the effective Coulomb interaction in BaNiS$_2$. For simplicity, we have assumed that the excited electrons are thermalized, i.e., we can assign an
photoemission yield. Solid lines show the photoinduced changes in the intensity with the pump fluence of 0.2 mJ/cm² (solid lines). Dashed lines are the thermal-induced changes of the intensity while heating the sample starting from 130 K. The electronic and the lattice temperature converge around 20 K. The same set of data is shown for the $d_{x^2−y^2}$ band in panels (c) and (d).

Effective temperature to them at each time delay. This assumption is supported by our trARPES experiments. According to the experiment, when the magnitude of the renormalization effects is largest (at a ≈250-fs delay), the electrons have already been fully thermalized. We thus derive the fundamental relation between electronic temperature and effective Coulomb interaction in a layered Dirac material, such as BaNiS$_2$. Its semimetallic character is given by the linearly dispersing bands at the four Dirac points (at fixed $k_z$), which dominate the low-energy physics. Another relevant feature of BaNiS$_2$ is its layered structure, with weak band dispersion along the $k_z$ direction. Thus, the Dirac points become Dirac lines if one takes into account the $k_z$ extension of the Brillouin zone in the limit of null $k_z$ dispersion.

With the aim of incorporating these main ingredients, we start from a conical model in two dimensions, with the Dirac Hamiltonian written as

$$H = \gamma (\sigma_z k_x + \sigma_y k_y) - \mu,$$

where $k = (k_x, k_y)$ is the 2D momentum, $\sigma_i$ are the Pauli matrices, $\gamma$ is the Fermi velocity, and $\mu$ is the chemical potential. For $\mu = 0$, the density of states of this model reads

$$D(\epsilon) = \frac{g_S g_C}{2 \pi \gamma^2} \left| \epsilon \right|,$$

with $g_S$ the spin degeneracy, and $g_C$ the number of Dirac cones. This model has largely been studied in literature, mainly in the context of graphene. Here, we address a very specific question, namely, the modification of the screened Coulomb interaction driven by electronic excitations in a system made of layers, separated by a distance $l_d$, where the noninteracting part of every layer is described by the 2D Hamiltonian in Eq. (B1), and the hopping between layers is neglected. We assume the system to be in the so-called intrinsic (i.e., undoped, $\mu = 0$) regime, which is appropriate for BaNiS$_2$. The dielectric function of such a model is

$$\epsilon(q, k_z) = 1 + v^{2D}(q) S(q, k_z) \Pi(q, \beta),$$

where $(q, k_z)$ is the three-dimensional (3D) momentum, $v^{2D}(q) = 2 e^2/(k_0 q)$ is the 2D Fourier transform of the Coulomb potential with $k_0$ the dielectric constant of the medium, $\Pi(q, \beta)$ is the polarization function of the conical model Eq. (B1), $\beta = 1/(k_B T)$ is the inverse temperature $T$ with $k_B$ the Boltzmann constant, and $S$ takes into account the effect of interlayer interactions:

$$S(q, k_z) = \frac{\sinh(q l_d)}{\cosh(q l_d) - \cos(k_z l_d)}.$$

The latter equation has been derived by linear response theory in Ref. [41]. In Eq. (B3), we take the static limit of the dielectric function, while keeping the full temperature dependence $\beta$. This is at variance with previous studies which focused on its zero-temperature limit. We are most interested here in the impact of electronic excitations to the static screening of the quasi-2D Dirac systems.

It is well known [42,43] that, in the intrinsic regime, the 2D polarization function of the conical model takes the form

$$\Pi(q, \beta \to \infty) = \frac{g_S g_C}{16 \pi \gamma^2} q.$$

However, the zero-momentum limit is profoundly modified by thermal excitations, which give a finite value of $\Pi(q \to 0, \beta)$. We show in the following that this holds at any finite temperature. This result can be obtained by taking the $q \to 0$ limit of the full $\Pi(q, \beta)$ function derived in Ref. [44] through quantum field theory techniques:

$$\Pi(q, \beta) = \frac{g_S g_C}{\pi \gamma^2} \int_0^\infty \frac{d\epsilon}{\beta} \log \left[ 2 \cosh \left( \frac{q \gamma \beta \sqrt{1 - x}}{2} \right) \right].$$

The $q = 0$ finite-temperature expression can also be derived by exploiting the compressibility sum rule:

$$\Pi(0, \beta) = \frac{g_S g_C}{2 \pi \gamma^2} \frac{1}{\beta} \log(2 \cosh(q \gamma \beta \sqrt{1 - x})).$$

From both Eqs. (B6) and (B7), one easily obtains that

$$\Pi(0, \beta) = \frac{g_S g_C}{2 \pi \gamma^2} 2 \log(2) \frac{1}{\beta^2}.$$
Table I. Values used to evaluate the inverse screening length \(\omega_{\text{TF}}\) in Eq. (B11) as a function of temperature for BaNiS\(_2\). An educated guess for the dielectric constant of the medium, \(\kappa_0\), is taken from the value measured for sulfur [41], as the in-layer Ni states which make the Dirac cones in BaNiS\(_2\) are surrounded by sulfur pyramids and hybridized with sulfur \(p\) states. The function \(\omega_{\text{TF}} = \omega_{\text{TF}}(T)\) is plotted in Fig. 3(b).

| \(g_s\) | \(g_C\) | \(\kappa_0\) | \(\gamma \,(\text{Å eV})\) | \(l_d \,(\text{Å})\) |
|---|---|---|---|---|
| 2 | 4 | 3 | 2.7 | 8.908 |

To compute the dielectric function of the 3D layered system, one can plug the temperature-dependent expression for the 2D polarization function of Eq. (B8) into Eq. (B3). By using the Thomas-Fermi (TF) approach, the resulting 3D screened interaction is

\[
V_{\text{screened}}^{3D}(r) = \frac{2e^2}{\pi \kappa_0} \int_{-\pi/l_d}^{\pi/l_d} dx K_0(r2D\sqrt{x^2 + \omega_{\text{TF}}^2}) \exp(iH_d x) \tag{B9}
\]

where the 3D distance is given by \(r = \sqrt{r^2_{3D} + (l_d)^2}\), with \(r_{2D}\) the distance on the layer, and \(H_d\) the out-of-plane distance, with \(l\) an integer. \(K_0\) is the modified Bessel function of the second kind. The large-\(r\) asymptotic behavior of Eq. (B9) is given by

\[
V_{\text{screened}}^{3D}(r) \sim e^2 \kappa_0 r^{-2} e^{-r/\omega_{\text{TF}}}, \tag{B10}
\]

which has the usual Yukawa form, but this time with a TF screening parameter that reads

\[
\omega_{\text{TF}} = \sqrt{4 \log(2)} \frac{e^2 g_s g_C}{\kappa_0 \gamma^2 l_d^2 \beta}. \tag{B11}
\]

Therefore, the stronger the excitations, the shorter the screening length. Indeed, the inverse screening length has a \(\sqrt{T}\) temperature dependence. This is very different from conventional metals, where the inverse screening length goes like \(1/\sqrt{T}\), according to the Debye-Hückel behavior in the high-temperature limit. An expression similar to Eq. (B11) has been found for nodal-line semimetals in an extrinsic high-temperature regime (i.e., \(0 < \mu \ll T\)) [45]. The parameters of Eq. (B11) which are appropriate for BaNiS\(_2\) are reported in Table I.

The remarkable agreement between the band-structure modification computed from first principles and the one measured in pump-probe ARPES comes directly from the phase-space restrictions of the Dirac dispersion, which implies that in a regime of excited Dirac states the screening length is shorter than in the relaxed regime.

**APPENDIX C: AB INITIO CALCULATIONS**

The BaNiS\(_2\) geometry is taken from the experiment, as described by Grey and Steinfink [46]. The ab initio density functional theory calculations are done with the modified Heyd, Scuseria, and Ernzerhof (HSE06) functional [30–32], as explained in the paper, by using the QUANTUM ESPRESSO package [36,37]. The spin-orbit coupling is included in the Ni norm-conserving pseudopotential in a \(4s^23d^8\) atomic configuration. Norm-conserving pseudopotentials are used also to replace Ba and S atomic cores. The Ba pseudopotential includes the semicore states in valence, while S is in the \(3s^23p^4\) configuration. The total energies are converged with an 8 \(\times\) 8 \(\times\) 8 electron-momentum grid, a Methfessel-Paxton smearing of 0.01 Ry for the \(k\)-point integration, and a plane-wave cutoff of 60 Ry for the wave-function expansion. To compute the nonlocal Fock operator, we employed a downsampled 8 \(\times\) 8 \(\times\) 8 angular-momentum grid, a Methfessel-Paxton smearing of 0.01 Ry for the \(k\)-point integration, and a plane-wave cutoff of 60 Ry for the wave-function expansion. To compute the nonlocal Fock operator, we employed a downsampled 8 \(\times\) 8 \(\times\) 8 q-point grid, with a half-a-grid shift in the \(z\) direction to minimize the number of nonequivalent momenta in the \(k + q\) mesh.

**APPENDIX D: ULTRAFAST RENORMALIZATION at \(\Gamma\)**

Here, we show that the electron pocket undergoes an ultrafast renormalization with photoexcitation. By analyzing the MDC’s, we plot the band dispersion as well as the wave-vector renormalization (\(\Delta k\)) for each time delay. The results are shown in Figs. 7(b) and 7(c), respectively.
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