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Circular Dichroism and Superdiffusive Transport at the Surface of BiTeI

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We investigate the electronic states of BiTeI after the optical pumping with circularly polarized photons. Our data show that photoexcited electrons reach an internal thermalization within 300 fs from the arrival of the pump pulse. Instead, the dichroic contrast generated by the circularly polarized light relaxes on a timescale shorter than 80 fs. This result implies that orbital and spin polarization created by the circular pump pulse rapidly decays via many-body interaction. The remnant dichroic contrast at longer delay times is due to the helicity dependence of superdiffusive transport. We ascribe it to the lack of inversion symmetry in an electronic system far from equilibrium conditions.

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The application of surface and interface states in spintronic devices motivates an increasing interest in semiconductors with sizable spin-orbit interaction. The ternary compound BiTeI is a valuable example of non centrosymmetric material with surface states supporting large spin polarization [1–5]. A macroscopic spin injection in these surface states may be obtained by optically pumping the system with circularly polarized photons. Such a technique has been widely employed to optically orient the spins of bulk semiconductors [6] and GaAs/AlAs quantum wells [7]. Here the optical transition selection rules of the Bloch bands are inherited from their parent atomic orbits. Alternatively, the spin polarization can be entangled to valleys degrees of freedom by intercellular currents [8]. It has been shown in monolayer MoS₂ that this spin polarization comes along with the contrasted circular dichroism in different regions of the Brillouin zone [9, 10]. As for the aforementioned systems, also BiTeI holds a large spin-orbit coupling and lacks inversion symmetry. These properties lead to a finite spin orientation upon photoexcitation with circular polarized beams [11]. Despite it, the surface of BiTeI does not have a band gap but metallic electronic states [1, 3, 4]. Therefore, fast scattering channels can relax the orbital and spin polarization on a femtosecond timescale. On the other hand, superdiffusive transport from the surface towards the bulk leave a remnant trace of the initial polarization for a longer time.

This letter aims to a better understanding of photoexcited states and the electronic transport phenomena at the surface of materials with strong spin-orbit interaction and without inversion symmetry. The ternary compound BiTeI is a benchmark example where to investigate the electronic evolution after an optical perturbation. We probe the electronic states in the transient regime by means of time resolved photoelectron spectroscopy. Our experimental technique has the unique capability of detecting the population of electronic states upon photoexcitation by pump pulses with different helicities. By these means, we observed an internal thermalization of the photoexcited electrons within 300 fs from the arrival of the pump pulse. Interestingly, the dichroic contrast acquired 80 fs after photoexcitation is already different from the one expected from the point group symmetry of crystal. We deduce that the initial orbital and spin orientation decay on a timescale much faster than the electronic thermalization. The remnant dichroic contrast arises from the helicity dependence of the superdiffusive transport of the electrons that are highly out of equilibrium. This transport phenomenon occurs only in the period when non-thermalized electrons do not respect the detailed balanced conditions and it is strictly linked to photogalvanic effects [12].

Photoelectron spectra with photon energy of 28 eV have been collected at the Cassiopée beamline of Soleil Synchrotron. Time resolved photoelectron spectroscopy experiments have been instead performed with the FemtoARPES setup, using a Ti:Sapphire laser that generates 35 fs pulses centered at 790 nm with repetition rate of 250 kHz. Part of the beam is employed to generate the fourth harmonic by a cascade of frequency mixing in BBO crystals (β-BaB₂O₄) [13]. The 197.5 nm probe and the 790 nm pump are focused on the sample with a spot diameter of 100 µm and 200 µm, respectively. Their
FIG. 1: A): Calculated band structure of the Te terminated BiTeI surface for parallel wavevector along Γ-K and Γ-M. B): Photoelectron intensity map acquired along the Γ-K direction with photon energy of 28 eV.

A cross-correlation in a BBO crystal has a full width at half maximum (FWHM) of 80 fs. The bandwidth of the 197.5 nm beam (6.3 eV) limits the overall energy resolution of TRPES spectra to 60 meV. All the time resolved measurements have been performed at the base temperature of 130 K base pressure of 10^{-10} mbar. The samples are high quality single crystals of BiTeI cleaved in situ just before the data acquisition. If not differently specified, the employed pump fluence is 30 µJ/cm^2.

BiTeI is a polar compound in which Bi, Te, and I form stacking layers with the structure of an honeycomb lattice. The system admits a three fold rotation around the c axis and lacks inversion symmetry. Due to the large spin-orbit interaction and small bandgap the electronic states hold a large Rashba-like splitting [5, 14]. Spin resolved ARPES measurements of the surface states report an in-plane polarization with helical spin texture [1, 4]. Bulk bands hold a similar structure but generate closed Fermi surfaces around the A symmetry point of the 3D Brillouin zone [4]. Although the cleavage of BiTeI, may expose different terminations [3], our data are indicative of a surface with topmost Te atoms and negative band bending. We show in Fig. 1A, the calculated band structure of the Te terminated surface. The two dimensional electronic pockets generated by the surface states extend to higher binding energy and partially overlap with the bulk states. Figure 1B displays a photoelectron intensity map acquired along Γ-K with photon energy of 28 eV. The two dispersing branches are surface states split by the spin-orbit interaction. The bulk electronic states are not visible at this photon energy because of the extreme surface sensitivity of the photoemission spectroscopy. Instead, the surface and bulk states attain comparable photoemission intensity at photon energy below 7 eV [1]. The intensity map of Fig. 2A has been obtained with photons of 6.3 eV and it is compatible with previous ARPES experiments based on laser sources [1]. Even if individual bands cannot be discriminated, the size and shape of the electrons pockets match well the calculations of Fig. 1A.

Next, we focus on the dynamics of the electrons in non-equilibrium conditions. Figure 2B-D shows intensity maps at different delay times after the arrival of the linearly polarized pump beam. Upon photoexcitation, the electronic states above the Fermi level become partially filled. The largest part of the primary excitations takes place in bulk-like states, populating two dispersing states up to 0.8 eV (see Fig. 2B). The temporal evolution reported by the intensity maps of Fig. 2A-D indicates that the Fermi liquid thermalizes by electron-electron interaction in ∼300 fs. During this quasi-adiabatic transient, the impact ionization channels increase the spectral intensity in electronic states near to the Fermi level. Afterwards, the hot electrons can be described by a Fermi-Dirac function with effective temperature and chemical potential. We trace the temporal evolution of BiTeI by monitoring the photoelectron intensity at energy 0.15 eV above the Fermi level (see Fig. 2E). This signal displays a step-like increase followed by a sub-picosecond rise and a steadily reduction. The cooling of hot electrons by phonon emission lasts ∼10 ps, until local equilibrium is established. Notice that similar timescales have been observed also in elemental bismuth [15] and topological insulators [16]. Three reasons explain this slow dissipation of electronic energy. First, the lack of large Fermi
surface hinders the carriers recombination in the bulk [4]. Second, the small quantum energy of the phonon modes results in scattering processes that are weakly inelastic [17]. Third, the weak band dispersion along the surface normal hinders an efficient diffusion of absorbed energy into the bulk [15].

Next, we describe the dichroic properties of BiTeI both in equilibrium and after photoexcitation. Figure 3A shows the geometry of the FemtoARPES setup for the dichroic measurements. The electrons are always detected in the incidence plane of the incoming light (namely xy). Both pump and probe beams define an angle of 45° with respect to the surface normal. We are going to compare the results of two complementary experiments: i) circular-probe measurements monitor the dichroic contrast obtained by switching the circular polarization of the probe photons and damping the pump beam, ii) circular-pump measurements provide instead the dichroic contrast collected by switching the circular-polarization of the pump-beam and polarizing the probe beam in the xy plane. Before any measurement, we have verified that the intensity of different polarizations are equal within an uncertainty of 1%.

Figure 3B shows the circular-probe signal along Γ-K as a function of energy and electronic wavevector. The contrast is maximal for electronic states lying ∼0.1 eV below the Fermi level and has a nearly odd parity with respect to the in-plane wavevector. We disclose the symmetry of this signal by integrating the photoelectron intensity in the [-0.2,0] eV interval and varying the orientation of the sample with respect to the xy plane. As shown in Fig. 3C, the resulting contrast changes sign on the Γ-M direction and is maximal along Γ-K. The observed dichroism is a very general property of C3v surfaces, arising from the inherent lack of the inversion symmetry in a photoemission process [18]. When the electronic states hold large spin-orbit splitting, the circular-probe dichroism can be also linked to the spin polarization [19]. As a consequence, dichroic maps have been readily employed to gain information about spin textures in topological insulators [20–22]. In our case, this comparison could be compelling only if the individual bands were clearly resolved. Nonetheless, the general trend in Fig. 3C may be compatible with an out-of-plane buckling of the spins in the electronic pockets [19, 23].

Figure 4A shows a circular-pump map acquired in the Γ-K direction at delay time of 80 fs. An almost identical map has been measured along Γ-M or other directions of the surface plane. The dichroic signal of Fig. 4A is nearly even with respect to k∥ and changes sign at the Fermi level. This result strongly deviates from the scenario expected in an independent particle picture. In-
Indeed, the atomic structure of the BiTeI surface has strong similarity with an honeycomb lattice without inversion symmetry. If many-body interactions could be dismissed, the absorption difference between left- and right-handed light would display a strong wavevector dependence. In particular, the dichroic signal should be maximal along \(G - K\) and with nearly-odd parity with respect to \(k_{||}\) [9].

Based on these general considerations, we conclude that the circular-pump map of 4A should look similar to the circular-probe map of 3B. In this case the photoexcited system would also acquire a macroscopic spin and orbital polarization [7, 8, 11]. Heuristic arguments suggest that the spin polarization should be roughly given by ratio between the average spin-orbit splitting \(\Delta_{SO}\) and the pump photon energy \(\hbar \omega\) [11]. By estimating \(\Delta_{SO} \approx 0.2\) eV and knowing \(\hbar \omega \approx 1.56\) eV, the resulting polarization should be \(\Delta_{SO}/\hbar \omega \approx 10\%\). Clearly, the presence of interactions inevitably relaxes the population imbalance generated upon the absorption of circularly polarized light.

Nonetheless, a nearly-odd circular-pump map could still be expected at early delay times. An interesting example has been reported in monolayer molybdenum disulphide, where the photoexcited carriers get trapped in valleys at the \(K\)-point of the Brillouin zone. Apparently, the inter-valley scattering of monolayer MoS\(_2\) is so weak that the circular-pump dichroism leads to a sizable polarization of the photoluminescence [9, 10]. The reason of such weak inter-valley scattering is the formation of excitons in the spin-polarized valleys [10]. As a consequence an electron-hole pair would scatter from \(K\) to \(K'\) valley have to overcome an energy barrier that roughly corresponds to the ionization potential of the exciton.

A different instance takes place in BiTeI, since no energy gaps and valleys can stabilize excitons. In contrast, the photoexcited electrons thermalize within \(\approx 300\) fs near to the zone center. In principle, a detectable orbital and spin polarization may still be expected during this thermalization time. However, the circular-pump map of Fig. 4A indicates that the measured dichroism attains an even-like symmetry already at delay times of 80 fs. This finding suggests that electron-electron scattering erases the orbital and spin polarization within the duration of the pump pulse. The remnant dichroic signal of \(\approx 15\%\) is the long living trace of the helicity dependent excitation set in by the pump pulse. Figure 4B shows the difference between maps of spatial inversion symmetry. Indeed, we have experimentally verified that no circular-pump contrast can be detected in bulk centrosymmetric systems as Bismuth and Bi\(_2\)Te\(_3\). We conclude that the different transport of energy density is due to chiral scattering events of an electronic distribution that does not respect detailed balanced conditions. In this sense, the observed dichroism shares common aspects with photogalvanic effects and circular photocurrents [12].

In conclusion, we have characterized the dynamics of electronic states photoexcited BiTeI. The thermalization of the electronic system occurs within 300 fs from the arrival of the pump beam. Upon excitation with circularly polarized pulses we observe a circular dichroism with a symmetry that differs from the ones expected from primary optical excitations. We conclude that spin and orbital polarization relax on a timescale faster than 80 fs. The expected temporal evolution of this dichroic contrast could resemble the relaxation of transient currents in image potential states [28]. Future experiments with shorter pulse duration may be capable to capture this dynamics. On the other hand, the remnant dichroism already indicates that the energy density at the surface of the sample depends on the helicity of the pump pulse. We ascribe this effect to superdiffusive transport occurring while the systems is out-of-equilibrium. The experimental evidence of this transport phenomenon may be of interest in related subjects as ultrafast demagnetization [26, 27].

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