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Controlling surface charge and spin density oscillations by Dirac plasmon interaction in thin topological insulators

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Thin topological insulator (TI) films support optical and acoustic plasmonic modes characterized by effective net charge or net spin density, respectively. We combine many-body and electromagnetic calculations to study how these modes can be selectively excited at films and nanodisks at infrared and THz frequencies. We first discuss the excitation of propagating plasmons in a thin film by a point dipolar source. We emphasize how changing the distance between the dipolar source and the film allows to control the relative strength of the acoustic and optical plasmons and thus to excite net-spin or net-charge waves on demand. The acoustic and optical modes in a nanodisk structure can be efficiently tuned by changing the size of the disk or by applying electrostatic gating. Furthermore, these modes can be confined to regions of dimensions much smaller than the wavelength. The control of the excitation of acoustic and optical modes indicates that thin topological insulators are a promising system to manipulate the spin and charge properties of the plasmonic response, with potential applications in fast, compact and electrically-controlled spintronic devices.

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INTRODUCTION

Plasmonic resonances supported by metallic structures[1–3] allow to localize and manipulate light using very small structures, of dimensions well below the diffraction limit. Their resonant frequency is controlled by the overall geometry (shape, material...), and cannot be easily tuned after fabrication. An alternative to plasmons in metals are Dirac plasmons [4] in two-dimensional (2D) materials such as e.g. graphene[5–9]. These structures support plasmonic resonances that can be tuned in the infrared and Terahertz frequencies by applying an external voltage (electrostatic gating), potentially reaching up to visible frequencies for systems just a few nanometers large[10]. The excitation and tunability of 2D plasmons in graphene has been demonstrated in both near and far field optical measurements[11–15].

Tunable 2D plasmons are also present in three-dimensional topological insulator systems made of materials[16] such as Bi2Se3 and Bi2Te3. These topological materials are characterized by a bulk phase that is transparent in the infrared frequency range due to the aperture of a gap and by (quasi-) 2D fermion layers (2D-FLs) at the interfaces with the surrounding medium[17–20]. As we do not consider 2D topological insulators presenting one-dimensional conduction channels[21], we simply call them topological insulators (TI) in the following.

The 2D-FLs of ∼ 10 Å width are formed by surface-state electrons having a linear energy dispersion near the surface Brillouin zone center, and can support surface plasmons at infrared and lower frequencies in a similar manner as graphene [16, 18, 19, 22–24], as have been shown experimentally [25–29]. Plasmons in TI can be tuned by electrostatic gating and present the additional advantage of carrying spin, which is locked with, and perpendicular to, the direction of propagation of the plasmon, and parallel to the surface at which they are excited. These excitations can thus be considered to be transverse spin waves[30, 31].

Notably, the 2D-FLs at the two interfaces of a thin TI film can interact via Coulomb coupling. As a consequence, two hybridized modes appear, an acoustic mode characterized by an approximately linear dispersion relationship and an optical mode emerging at larger energies and showing a square-root dependence with wave number (for low frequencies)[32, 33]. The optical (acoustic) mode is characterized by in- (out-of) phase charge waves induced at opposite surfaces of the thin TI film. Despite the similarity of these modes with those in spa-
tially separated double-layer graphene[34], the real spin–
momentum locking in TIs gives rise to a new phenomenon
predicted for thin TI films—spin-charge separation. It im-
plies that due to the opposite spin-momentum locking at
the two surfaces of the film, the optical mode corresponds
to an effective (pure) charge wave, while the acoustic
mode demonstrates pure spin character[31, 35, 36]. Thus,
thin films and other nanostructures made of a TI may
serve as a nice playground both for studying the fun-
damental properties of collective excitations in the 2D
Dirac fermions system and for engineering optoelectronic
devices with tunable spin-dependent characteristics.

For practical applications in nanotechnology, it is im-
portant to understand how these modes can be excited
and manipulated in an experiment, both for infinite
films that support propagating plasmons and for local-
ized plasmons in nanoparticles. The main objective of
this paper is thus to study the selective excitation and
control of optical and acoustic modes in TI thin films
and thin disks. We use a combination of many-body calcula-
tions to obtain a non-local conductivity \( \sigma \) characterizing
each 2D-FL (along with the plasmons dispersion of the
systems) and classical electromagnetic calculations that
take \( \sigma \) as an input and provide the optical response under
illumination by an external source. This approach also
allows to study the importance of including non-locality
when predicting the dispersion relationship of the sys-
tems.

After briefly introducing the dispersion relationship
of a thin film, we analyze the excitation of propagating
Dirac plasmons in such a system by localized point-like
dipolar sources. We study the influence of the distance
between the source and the film on the efficiency of the
excitation of the optical and acoustic modes, which al-
 lows thus to manipulate the spin and charge properties
of the excited surface wave. The last section analyzes
the excitation of localized acoustic and optical plasmonic
modes in thin disks, at frequencies that strongly depend
on the size of the structures. We are particularly inter-
ested in the tunability of the system when changing the
Fermi energy of the 2D-FLs for example by applying an
external voltage.

When not stated otherwise, atomic units are used in
the equations, i.e., \( e = \hbar = m_e = 1 \), where \( e \) is the
electronic charge, \( \hbar \) is the reduced plank constant and
\( m_e \) is the mass of the electron.

**SYSTEMS AND CALCULATION METHODS**

Fig. 1 shows the schematics of the different TI systems,
which are placed in vacuum (relative permittivity \( \varepsilon_1 = 1 \))
and illuminated at angular frequency \( \omega \) (vacuum wave-
length \( \lambda \) and vacuum wavenumber \( k \)). We first briefly
discuss the dispersion relationship of a semi-infinite sub-
strate (Fig. 1(a)) and a thin film (Fig. 1(b)), focusing
next on the near- and far-field response of the later. No-
tably, acoustic and optical propagating plasmonic modes
can be excited in thin films by localized sources, with
wavenumber \( q_{sp} \) and \( q_{op} \), respectively. We last discuss
the excitation of acoustic and optical localized plasmonic
resonances on finite thin disks (Fig. 1(c)). For the near
fields, it is convenient to differentiate between the to-
tal \( E_{tot} \) and the induced \( E_{ind} \) amplitude of the electric
fields, where only the former includes the fields from the
illumination source itself.

The semi-infinite TI substrate in Fig. 1(a) supports
a 2D-FL at the interface with vacuum, which would be
parallel to the (111) surface—the natural cleaving face of
the considered TI. The TI thin-film [Fig. 1(b)] and thin-
disk [Fig. 1(c)] has a thickness \( d=10 \text{ nm} \) and support
two coupled 2D-FLs, one at each flat interface (upper
and bottom). The film extends infinitely in the lateral
dimension, while the disks diameters considered are \( D =
40 \text{ nm} \) and \( 300 \text{ nm} \). The flat surfaces of the film and the
disk are perpendicular to the \( z \) direction and parallel to
the \( x-y \) plane (see axes in the inset). \( z=0 \) corresponds
to the central plane of the thin structures, with \((0,0,0)\)
the center of the disk.

We consider a simplified thin Ti system surrounded by
vacuum, with the objective of extracting general proper-
ties of the plasmon response under a given excitation.
The response results from the electromagnetic coupling
between the 2D-FLs at the top and bottom interfaces.
In this simple model, we choose a typical value[37–40]
for the relaxation time \( \tau = 500 \text{ fs} \) and for the Fermi ve-
locity of the electrons in the 2D-FLs of the Ti Bi2Se3.

![FIG. 1: Schematic of the TI structures considered. (a) Semi-
infinite TI substrate supporting a 2D-FL at the interface with
the surrounding vacuum, (b) an infinite TI film of thickness
\( d = 10 \text{ nm} \) (suspended in vacuum) with two interacting 2D-
FLs at the interfaces and (c) a finite TI thin disk of thickness
\( d=10 \text{ nm} \) surrounded by vacuum, with a 2D-FL at the upper
interface and another one at the bottom. The 2D systems
are characterized by an in-plane conductivity \( \sigma(q, \omega) \).
We consider Bi2Se3 as the topological insulator with relative di-
electric permittivity of the bulk \( \varepsilon_0 = 25 \). The center of the disk is at
(0,0,0), and \( z=0 \) is the horizontal middle plane of the
disk and the film in (b-c), in the coordinate system sketched
in (c). We use a Fermi energy of the 2D-FLs \( E_F = 250 \text{ meV} \),
unless specified otherwise.](image-url)
\[ v_F = 0.5 \cdot 10^6 \text{ m/s.} \] The bulk relative dielectric constant is taken as \( \varepsilon_b = 25 \), which is a reasonable approximation for Bi$_2$Se$_3$ in several regions of the THz spectrum[42–45]. We thus ignore phononic contributions[20, 25, 27, 46] that may be significant in real experiments but that can complicate the phenomena of interest for this paper, i.e. the excitation of optical and acoustic modes. In the case of the disk calculations, an idealized model has also been chosen for the lateral sides, which are treated as not presenting any 2D conductivity. Further, we do not include corrections to the electromagnetic constitutive equations[47, 48] (see Appendix B). We use a Fermi energy \( E_F \) of the 2D-FLs equal to \( E_F = 250 \text{ meV} \), unless other value is specified.

Our approach combines many-body and classical electrodynamic calculations. The many-body calculations described in Appendix A allow to calculate the dispersion relationship of the semi-infinite substrate and thin film, and to obtain the 2D in-plane conductivity \( \sigma(q, \omega) \). \( \sigma \) depends on frequency and on the horizontal (parallel) wave-vector in the 2D layer, \( q \). The \( q \)-dependence of \( \sigma(q, \omega) \) captures non-local effects[49–52] that modify the plasmonic response and that, in real-space, describe the dependence of the induced polarization at a given position on the electric fields at the surrounding region. We assume that the two 2D-FLs of the film and disk only interact electromagnetically (see Appendix A for details).

The far- and near-field optical response under a given illumination is then calculated by solving the Maxwell’s Equations, where each 2D-FL is modeled by the 2D conductivity \( \sigma(q, \omega) \). A dipole of strength \( 1 \text{ e} \cdot \text{nm} \) excites the film and disk of 40 nm diameter, while a plane-wave incoming from the top illuminates the larger disks of 300 nm diameter. We describe here only the main ideas behind the calculations, which are discussed in more detail in Appendix B. The effect of the conductivity of the 2D-FLs in this classical framework is to allow the accumulation of surface charge density \( (\sigma_c) \) at the interfaces, which affects the boundary condition[11]. The thin films are always excited by a point dipole (it is not possible to excite a plasmon in this case using far-field illumination), and we obtain the induced near-fields using a plane-wave decomposition method. By tracking the poles of the electromagnetic response (more exactly, the maximum of the transmission coefficient of the system) we also obtain a dispersion relationships that we compare with the results from the many-body model. This procedure allows to include the exact non-local \( q \)-dependence of the conductivity on the substrate and thin-film calculations. Furthermore, the local response can also be obtained by following the same procedure but using \( \sigma(q = 0, \omega) \). Last, we obtain the response of the disks under plane-wave or dipolar excitation using full-wave calculations[53]. Non-locality is introduced approximately into the calculations of the disks response by taking into account that each simulation is dominated by the response at a single \( q \), corresponding to either the acoustic \( (q_{SP}^a) \) or the optical \( (q_{SP}^o) \) mode.

**EXCITATION OF PROPAGATING DIRAC PLASMONS IN INFINITE SURFACES**

**Dispersion relationship**

We study in this section the excitation of propagating plasmons in TI systems that are infinite in the lateral direction (along the \( x \) and \( y \) directions indicated in the inset of Fig. 1). We are particularly interested on the thin films depicted in Fig. 1(b) but, for completeness, we first describe the dispersion of plasmons in a TI semi-infinite layer (sketched in Fig. 1 (a)). The main panel in Fig. 2(a) and Fig. 2(b) shows the dispersion relationship of plasmons in the substrate and thin film, respectively, for \( E_F = 250 \text{ meV} \). We first obtain it by changing the angular frequency \( \omega \) of an incoming (evanescent) plane wave and tracking which parallel wavevector \( q \) of this exciting field induces the strongest classical electromagnetic response (maximum in the transmission coefficient of the system). We consider both the full non-local \( [\sigma(q, \omega), \text{blue dots}] \) and the approximated local \( [\sigma(q = 0, \omega), \text{green solid lines}] \) conductivity. The non-local results are in very good agreement with the dispersion obtained directly from the many-body calculations[blue dots].

The results for the semi-infinite substrate in Fig. 2(a) reveal the presence of Dirac plasmons in the infrared-THz region of the spectrum that are characterized by a very large parallel wavenumber \( q_{SP} \) (low wavelength \( \lambda_{SP} = 2\pi/q_{SP} \)), compared to the corresponding value of the incident plane wave. For example, a surface plasmon excited at \( \lambda = 35 \mu \text{m} \) (\( \approx 0.0354 \text{ eV} \) has a (non-local) plasmonic wavelength \( \lambda_{SP} \approx 95 \text{ nm} \), showing a squeezing to \( \lambda/370 \). The field is thus evanescent and extremely confined in the vertical \( z \) direction, i.e. the vertical wavenumber at vacuum \( (k_z^1 = \sqrt{(2\pi/\lambda)^2 - q_{SP}^2}) \) and at the TI \( (k_z^{TI} = \sqrt{(2\pi/\lambda)^2 - q_{SP}^2}) \) are imaginary and satisfy \( k_z^1 \approx k_z^{TI} \approx q_{SP} \gg 2\pi/\lambda \). For reference, the light cone is indicated by the almost-vertical black dashed line in Fig. 2(a), which is characterized by wavenumber \( 2\pi/\lambda \).

The local and non-local dispersions are similar for low energies. In contrast, for sufficiently high energy (above \( \approx 0.03 \text{ eV} \)), both curves in Fig. 2(a) exhibit important quantitative and qualitative differences: in the case of the non-local approximation, the dispersion relationship becomes approximately linear, while for the local calculations the resonant energy is proportional to the square root of \( q_{SP} \). As a result, the nonlocal slope is \( \approx 3 \) times larger at the maximum \( q_{SP} \), shown, underlying the importance of using a non-local approach to accurately obtain the response for large energies. Further, the inset shows the strong dependence of the plasmon dispersion on the
Fermi energy $E_F$ of the 2D-FL, studied for the range $E_F = 150 \text{ meV} - 350 \text{ meV}$ in the non-local description. $E_F$ can be experimentally controlled by an external voltage bias, so that the good tunability of the system constitutes a major advantage of 2D plasmons [12, 54, 55].

Fig. 2(b) shows non-local (blue solid lines) plasmon dispersion for the $d = 10 \text{ nm}$ thin TI film. The 2D-FLs at each interface interact with each other via Coulomb coupling and two hybridized plasmon modes emerge as two different branches in the dispersion, as studied in Ref. 35. The low and high energy branches of the dispersion fall at each side of the dispersion curve for the semi-infinite substrate, (red line, corresponding to the non-local calculation). The mode associated with the lower energy branch exhibits an almost linear dependence between energy and wavenumber (linear dispersion relationship). It is usually called acoustic mode, and we will show that it is characterized by anti-symmetric charges at both interfaces (sketch in Fig. 3 (b)). The acoustic plasmon is characterized by the largest plasmon parallel wavenumber $q_{CS}'$ (smallest plasmon wavelength) and thus the strongest confinement (largest $|k_z^{1}|$). On the other hand, the larger energy branch corresponds to the optical mode and present a symmetric charge distribution (sketch in Fig. 3 (c)). The wavenumber $q_{CS}'$ for the optical branch and energies smaller than $\approx 0.075 \text{ eV}$ is significantly smaller than for the acoustic mode or for the plasmon of a single interface but remains much larger than that of a propagating plane wave in vacuum, so that the electromagnetic field of both acoustic and optical plasmonic modes are well confined to the TI film.

For energies $\gtrsim 0.075 \text{ eV}$ the acoustic and optical branches become similar to the dispersion relationship of a single interface.

As for the semi-infinite substrate, the dispersion relationship at low energies change only weakly when comparing the results obtained using the local $\sigma(q = 0, \omega)$ (green solid line) and non-local conductivity $\sigma(q, \omega)$ (blue solid line). Nonetheless, non-locality plays a significant role in the high energy response. Looking first at the acoustic mode, the wavenumber $q_{CS}'$ for the optical branch and energies smaller than $\approx 0.075 \text{ eV}$ is significantly smaller than for the acoustic mode or for the plasmon of a single interface but remains much larger than that of a propagating plane wave in vacuum, so that the electromagnetic field of both acoustic and optical plasmonic modes are well confined to the TI film.

For energies $\gtrsim 0.075 \text{ eV}$ the acoustic and optical branches become similar to the dispersion relationship of a single interface.

**Fields and surface charges induced by a point dipole**

Due to their very large wavenumber $q_{SP} \gg 2\pi/\lambda$, a plane-wave cannot excite the plasmonic modes in the TI
FIG. 3: Fields excited by a dipole illuminating a 10 nm thin TI film. (a) Maps of the real part of the $z$ component of the electric field $E_{\text{ind}}^z$ induced by a dipole emitting at $\lambda = 35 \mu m$, situated 3 nm above the film and oriented along the vertical $z$ direction. (b) Real part of the field in (a) at 3 nm [short-dash lines in (a)] away from the film minus the corresponding value for 50 nm distance [long-dash lines in (a)]. This subtracted field $E_{\text{diff}}^z$ is calculated both for fields above [solid green line in (b)] and below [dashed red line in (b)] the thin film, following Eq. 1. (c) Real part of the field calculated at 50 nm distance from the upper (solid green line) and bottom (red dashed line) surfaces, along the dashed lines in (a). The insets in (b) and (c) indicate the charge distribution corresponding to the fields of the acoustic and optical modes, respectively. The Fermi energy is $E_F = 250$ meV, the dipole strength is $1e \cdot nm$ and the field is plotted in S.I. units (Vm$^{-1}$).

We thus use a point-like dipole source, which, as discussed in Appendix B, can be decomposed as an integral over plane-waves of different horizontal wavenumber $q$, with a significant contribution for a very large range of $q$ values (very broad distribution in $q$-space). More exactly, we use an electric dipole oriented in the vertical $z$ direction and emitting at $\lambda = 35 \mu m$.

We first consider a dipole positioned 3 nanometers above the TI thin film. Fig. 3(a) shows the real part of the $z$ component of the electric field induced by the dipole, $\text{Re}(E_{\text{ind}}^z)$, in the vertical $x-z$ plane. Due to the rotational symmetry of the dipolar illumination (with respect to the $z$ axis), the induced electromagnetic fields are rotationally symmetric as well. The oscillatory behavior of $\text{Re}(E_{\text{ind}}^z)$ in the radial $x$ direction is typical of a propagating surface wave. We observe clear oscillations near (and inside) the film with periodicity $\approx 70$ nm, very close to the plasmonic wavelength of the acoustic mode $\lambda_{\text{SP}}^{ac}$ obtained from the dispersion ($\lambda_{\text{SP}}^{ac} = 2\pi/q_{\text{SP}}^{ac}$, with $q_{\text{SP}}^{ac}$ indicated by the blue open circle in Fig. 2b). These fast oscillations clearly suggest the excitation of the acoustic plasmon.

The fast decay in the vertical $z$ direction of these fast field oscillations indicate that they are strongly localized to a thin region near the TI film. The fields also decay along the direction of propagation ($x$ axis), but a large number of oscillations can be observed, revealing a large propagation distance (measured in units of the plasmonic wavelength). The decay of the plasmon in this direction is both due to the material losses associated with the relaxation time and to the $1/\sqrt{x}$ dependence of the electric field of a lossless propagating plasmon required by energy conservation (due to the cylindrical character of the plasmon wavefronts). A large propagation distance is desired for effective coupling and information transfer with distant systems.

As implied above, the $\approx 70$ nm periodicity strongly supports the excitation of the acoustic plasmon[35], characterized by fields oscillating in phase just above and below the TI (symmetric field distribution, corresponding to anti-symmetric charges as discussed below). According to the location of the maxima and minima of $\text{Re}(E_{\text{ind}}^z)$ in Fig. 3(a), however, one can appreciate that the phase of the fields is not exactly identical at both sides of the film, and that the difference becomes larger in planes further away from the film (comparing the fields at $|z|$ to those at $-|z|$ for large $|z|$). Indeed, for planes sufficiently far away from the TI, the fields oscillate approximately with opposite phase above and below the film. This asymmetric field distribution corresponds to the expectation for optical modes[35].

We attribute this behavior to the simultaneous excitation of both an optical and an acoustic propagating surface plasmon; the optical plasmon dominates the fields
induced far away from the film, while the contribution of the acoustic plasmon become very important near the interfaces. More in detail, the excitation of the acoustic mode lead to the presence of the fast oscillations near the film (set by $q_{sp}^{ac}$) characterized by symmetric field with respect to the central plane. The large $q_{sp}^{ac}$ implies that the (evanescent) fields decay very fast in the vertical $z$ direction ($k_z^1 \approx q_{sp}^{ac}$), so that it becomes difficult to appreciate these oscillations for $|z|$ larger than a few tens of nanometers. In contrast, the optical mode, characterized by asymmetric field distribution and larger plasmonic wavelength, becomes dominant at sufficiently large $|z|$ due to its comparatively smaller $|k_z^1| \approx q_{sp}^{op} < q_{sp}^{ac}$.

To confirm the simultaneous excitation of acoustic and optical modes with opposite field symmetries, we obtain $E_z$ at both sides of the film, at 3 nm and 50 nm away from it (along the two short-dash and two long-dash lines in Fig. 3(a), respectively). Since the acoustic mode decays faster than the optical plasmon in the $z$ direction, we expect to be able to discriminate the fields associated with the former by subtracting the fields excited at the larger distance to those at the smaller, i.e.,

$$E_z^{\text{diff}} = E_z^{\text{ind}}(z = \frac{d}{2} + 3 \text{ nm}) - E_z^{\text{ind}}(z = \frac{d}{2} + 50 \text{ nm})$$

$$E_z^{\text{diff}} = E_z^{\text{ind}}(z = -\frac{d}{2} - 3 \text{ nm}) - E_z^{\text{ind}}(z = -\frac{d}{2} - 50 \text{ nm}) \quad (1)$$

above and below the TI, respectively. Indeed, Fig. 3(b) demonstrates that the fast oscillations of the subtracted signal $E_z^{\text{diff}}$ above and below the film are in phase, as expected for the acoustic plasmon, and that the $\approx 70 \text{ nm}$ periodicity is in good agreement with the value $2\pi/q_{sp}^{ac}$ obtained from the acoustic branch of the dispersion relationship (as marked by the open circle in Fig. 2(b))

As we evaluate the fields further away from the film the relative weight of the optical mode increases. We plot in Fig. 3(c) the real part of $E_z^{\text{ind}}$ at a distance of 50 nm above and below the film [along the long-dash lines in Fig. 3(a)]. The fields have an opposite sign at each side of the film, i.e. opposite orientation or $\pi$ phase difference, with a periodicity of the oscillations of $\approx 1000 \text{ nm}$.

This anti-symmetry corresponds to symmetric charges (see below) and is typical of an optical mode in thin TI films [35], and the periodicity matches well with the value $q_{sp}^{op}$ obtained from the corresponding branch of the dispersion relationship (marked by the blue square in Fig. 2(b)). The fields in Fig. 3 are thus consistent with the excitation of both an acoustic and optical mode, the former strongly contributing to the fields in the vicinity of the film and the latter predominating at positions further away.

The field symmetries can be translated into charge symmetries through the fields at both sides of each 2D-FL. The surface charge density $\sigma_z$ at the upper $\sigma_z^u$ and bottom $\sigma_z^b$ interfaces (in atomic units) are

$$\sigma_z^u = \frac{\varepsilon_1}{4\pi} E_{1z}^{\text{tot}}(z = \frac{d}{2}) - \frac{\varepsilon_b}{4\pi} E_{1z}^{\text{tot}}(z = -\frac{d}{2})$$

$$\sigma_z^b = -\frac{\varepsilon_1}{4\pi} E_{1z}^{\text{tot}}(z = -\frac{d}{2}) + \frac{\varepsilon_b}{4\pi} E_{1z}^{\text{tot}}(z = -\frac{d}{2}) \quad (2)$$

where $E_{1z}^{\text{tot}}(z = \pm \frac{d}{2})$ is the total field along $z$ on the vacuum side of the interface, and $E_{1z}^{\text{tot}}(z = \pm \frac{d}{2})$ the equivalent value inside the TI, both evaluated just at the film-vacuum interfaces. The sign difference between the equations of $\sigma_z^u$ and $\sigma_z^b$ is due to the dependence of the charge density on the direction of the surface normal. The anti-symmetric $E_z$ field exhibited by the optical modes thus corresponds to a charge distribution characterized by equal sign at both interfaces, which explains the comparatively larger energy of the optical branch in the dispersion relationship as a consequence of Coulomb repulsion between equal charges at the upper and bottom 2D-FLs [inset of Fig. 3(c)]. In contrast, the acoustic
modes present charges of opposite sign at the two TI-vacuum interfaces [inset of Fig. 3(b)]. As we will see in the next section, these charge symmetries are the opposite to those of the spin, which introduces a handle to control the net charge and net spin separately. Furthermore, we also note that these charge symmetries are the opposite to those of the optical and acoustic plasmonic modes in a thin metallic film [57], where the difference between TIs and metal films appears because the latter exhibit a screening effect due to the 3-D electron gas (described by the negative value of the real part of the permittivity) that is not present in TIs.

**Controlled excitation of charge and spin waves by dipole-sample separation control**

Since the contribution to the fields induced by the dipole in Fig. 3 from the acoustic and optical modes can be dominant near and far from the TI film, respectively, reciprocity [58] suggests that the acoustic mode would be effectively excited by a dipolar source situated near the film, whereas the optical mode would be predominantly excited when the dipole is far away from the film. With this in mind, Fig. 4 shows the real part of the surface charge densities \( \sigma \) induced in the upper and bottom interfaces by a dipolar source radiating at \( \lambda = 35 \mu m \) situated at (a) 5 nm and (b) 150 nm away from the surface of the film.

When the dipole is at a distance of 5 nm above the 10 nm thin film [Fig. 4(a)] the induced charges at the upper and bottom surfaces has opposite sign and a spatial periodicity of \( \approx 70 \) nm, indicating as expected the excitation of an acoustic mode. As an aside, Fig. 3 showed that a dipole near the thin film also excite an optical mode that contributes to the near-fields outside the TI, but whose contribution to the charge distribution in Fig. 4(a) is difficult to appreciate; this contrast between field and charges is possible because \( \sigma \) depends on the (displacement) fields outside and inside the film (Eq. 2).

For a dipole placed 150 nm above the surface, [Fig. 4(b)], the charge densities oscillate in phase at both interfaces and oscillate with the larger spatial periodicity of \( \approx 1000 \) nm characteristic of the optical plasmon. The charges are thus indeed dominated by the acoustic mode for 5 nm distance, and by the optical mode for 150 nm. In consequence, it is possible to tune the relative strength of the two modes by varying the dipole-film distance. In this way, we can control not only the charge but also the spin properties, as we discuss in more detail below.

We analyze next the response in terms of net charge and net spin plasmonic waves. The two top panels in Fig. 5(a) show the real part of the induced charge density oscillations at the upper (\( \sigma^u \)) and bottom (\( \sigma^b \)) 2D-FL excited by the vertical dipole at 5 nm above the film. As just discussed, the surface charge density with \( \approx 70 \) nm period is characterized by opposite phase at the upper and lower faces, corresponding to the acoustic plasmon wave. Thus, the total effective charge density \( \sigma^c_{\text{eff}} \), defined as \( \sigma^c_{\text{eff}} = \sigma^u + \sigma^b \), is nearly zero.
Indeed, \( \text{Re}(\sigma_x^{\text{eff}})/2 \), shown in the second panel from the bottom, is only comparable to either \( \text{Re}(\sigma_x^0) \) or \( \text{Re}(\sigma_x^c) \) near the position of the dipole, where many of the evanescent components of the dipolar excitation contribute to the response, not only those associated with the excited plasmonic mode. On the other hand, a dipole at 150 nm above the film, which excites predominantly the \( \sigma_{\text{SP}} \approx 1000 \text{ nm} \) optical mode, results in a strong total effective charge, as the charge densities at both surfaces of the film oscillate in phase (Fig. 5(b)).

The peculiarity of the TI film, compared, for instance to two graphene layers coupled electrostatically, is the surface-state spin texture with well-defined helicity at the upper and bottom film surfaces. Thus, these TI films are potentially interesting for spintronic applications. The spin is parallel to the 2D-FLs, transverse to the direction of propagation of the excited plasmon and its direction depends on the surface normal. More specifically, the transverse spin polarization density at the interfaces, of amplitude \( s_T \) couples with the charge density as given by \( s_{\text{SP}}^{u,b}(q, \omega) = \mp \text{Re}(\sigma_{\text{SP}}^{u,b})(q, \omega) \) for the upper \( s_T^u \), minus sign) and bottom \( (s_T^b, \text{ plus sign}) \) surfaces[30, 33]. Thus, for the in-phase charge oscillations characteristic of the optical mode the transverse spin polarization have opposite direction at the two 2D-FLs. In contrast, the acoustic resonance is characterized by the same spin-polarization direction at both interfaces.

To provide a more precise characterization of the behavior of the effective spin, we calculate the spin density amplitude as a function of position \( \mathbf{r} \) by performing the Fourier transform of \( s_T^{u,b}(q, \omega) \). We then plot \( \sigma_{\text{SP}}^{\text{eff}}/2 = (s_T^u + s_T^b)/2 \) in the bottom panels of Fig. 5(a,b), which confirms that \( \sigma_{\text{SP}}^{\text{eff}} \) is large when the net charge density \( \sigma_{\text{SP}}^{\text{eff}} \) is small and vice versa. In short, the acoustic and optical modes can be seen as spin-like or charge-like density waves, respectively, whose relative strength is controlled by the position of the dipole; specifically, as the dipole is displaced from very short to very large distance from the TI film, we transition from exciting spin collective excitations to inducing charge density waves.

**LOCALIZED SURFACE PLASMONS IN Bi_2Se_3 NANO DISKS**

We move next from studying propagating plasmons in infinite TI surfaces to the excitation of localized Dirac plasmons in finite TI structures, such as the nanodisks of diameter \( D \) depicted in Fig. 1(c). These disks can act as optical nanoantennas enhancing and confining the field, leading for example to strong interactions with nearby objects or molecules[59–62]. As we illustrate below, the rupture of translational symmetry also allows the excitation of (optical) plasmonic modes by an incoming plane wave.

Similarly to what has been studied for metallic structures[63, 64], plasmonic resonances in disks can be understood as Fabry-Pérot-like cavity modes, for which the reflection of propagating plasmon at the edges leads to localized resonances when constructive interference occurs. The disks support many different resonances[7, 65, 66], and, for example, the difference between edge and sheet plasmons in graphene and metallic disks have been recently studied[6, 67, 68]. For simplicity, we focus on configurations where only a few of the possible modes are excited. More in detail, we choose illumination sources that induce modes where the charge densities induced at the 2D-FLs are antisymmetric with respect to the vertical \( x = 0 \) plane (not to be confused with the symmetry with respect to the middle of the film discussed in the previous section) and present a cosine-like evolution in the polar direction (in the \( xy \) plane). In a simple model treating the plasmon as a plane wave-like surface excitation of well defined wavevector \( q_{\text{SP}} \) and assuming zero reflection phase \( \phi_r \) of the plasmon at the edges of the disk of diameter \( D[69, 70] \), these resonances emerge when \( q_{\text{SP}} \) verifies

\[
q_{\text{SP}} = \frac{\pi n}{D}
\]

with \( n \geq 1 \) an odd integer (even values of \( n \) describe rotationally symmetric resonances not excited by our illumination scheme). Eq. 3 corresponds to the condition of coherent interference that occurs when the plasmon accumulates a \( 2\pi \) difference in a full round trip. Refs. [71, 72] use a related model in terms of Bessel Functions that gives a somewhat different \( q_{\text{SP}} \) but does not change the general conclusions below. Once \( q_{\text{SP}} \) is known from Eq. 3, the resonant energy is obtained via the dispersion relationship. Notice that considering a non-zero reflection phase \( \phi_r \) would modify to some extent the position of the resonances with respect to the prediction by this simplified equation.

In the following sections, we consider the thin disks sketched in Fig. 1(c) with thickness \( d = 10 \text{ nm} \) and diameter \( D = 40 \text{ nm} \) or \( D = 300 \text{ nm} \).

**Excitation of localized optical modes with plane wave illumination**

Fig. 6(a) shows the extinction coefficient (extinction cross section normalized by the disk top surface area) of the disk of 300 nm diameter, when illuminated by a plane wave of amplitude \( E_0 \) incident from the top with linear polarization along \( x \), as schematically shown in the figure. This polarization imposes anti-symmetry of the induced charges in the direction of polarization, i.e. with respect to \( x = 0 \). Due to the symmetry of the structure with respect to the horizontal plane \( z = 0 \) and the negligible phase shift associated with the propagation of the plane-wave across the small thickness of the disk, the
FIG. 6: Optical response of a thin TI disk of diameter 300 nm and thickness $d = 10$ nm that supports localized optical plasmons. The excitation is a plane wave polarized along the $x$ direction and incident from the top ($z$ axis), as shown in the schematic (top right). The coordinate axis indicates the different directions, and the center of the disk is situated at $x = 0$, $y = 0$, $z = 0$. (a) Extinction coefficient of the disk, for the Fermi energy $E_F = 250$ meV (blue solid line) and $E_F = 350$ meV (red solid line) of the 2D-FLs. (b) Maps of the imaginary part of the $z$ component of the total electric field normalized to the amplitude of the incoming wave, $\text{Im}(E_z^\text{tot})/E_0$, at the prominent lowest-energy spectral peak at $\lambda = 34.5$ $\mu$m for $E_F = 250$ meV. In the top panel, the fields correspond to the horizontal plane $z = 10$ nm, parallel to the disk and 5 nm over the top interface. The fields in the bottom panel are obtained in the $y = 0$ vertical plane passing through the center of the disk and parallel to the incident electric field. (c) as in (b), but for the second lowest-energy peak at $\lambda \approx 24.5$ $\mu$m in the $E_F = 250$ meV spectrum.

The extinction spectra (Fig. 6a) show clear peaks associated to the localized plasmon resonances. For $E_F = 250$ meV (blue solid line), the two lowest energy peaks are found at $\lambda = 34.3$ $\mu$m and $\lambda = 25.4$ $\mu$m. For comparison, the simple Eq. 3 using $n = 1.3$ predicts resonances at $\lambda = 30.7$ $\mu$m and $\lambda = 25.5$ $\mu$m, in reasonable agreement with the simulated results. The broadness of a peak at frequency $\omega_{pl}$ can be quantified with the quality factor $Q = \omega_{pl}/\Delta \omega$, where $\Delta \omega$ is the frequency difference corresponding to the full-width half maximum of the peak. The quality factor of the $\lambda = 34.3$ $\mu$m peak is $Q = 26.9$, comparable to the values found in plasmonic systems[73].

It is in good agreement with the result of the quasistatic equation[73] $Q = \omega_{pl}^2/(4 \pi n \epsilon_0 \sigma(\omega_{pl})) \approx 27$ (evaluated at the resonant frequency), which only depends on the real and imaginary part of the conductivity, not on the exact geometry.

The extinction peaks shift significantly for moderate changes of the Fermi energy $E_F$ (which modifies $\sigma(\omega_{pl}, \omega)$). The lowest energy peak, for example, shifts from $\lambda = 34.3$ $\mu$m to $\lambda = 29$ $\mu$m when $E_F$ changes from 250 meV (blue solid line) to 350 meV (red solid line). The energy shift is $\approx 4$ times the full with at half maximum $\Delta \omega$ of the $E_F = 250$ meV peak, illustrating the high tunability of plasmonic resonances in TI disks.

In order to confirm the character of the spectral peaks, we calculate the electric field distribution around the disk at the position of the two lowest energy peaks for the $E_F = 250$ meV spectrum (blue spectrum in Fig. 6a). Fig. 6(b) shows the imaginary part of the $z$ component of the amplitude of the total electric field, normalized by the amplitude of the incident plane wave $E_0$, $\text{Im}(E_z^\text{tot})/E_0$, for the $\lambda = 34.3$ $\mu$m resonance. Fig. 6(c) presents $\text{Im}(E_z^\text{tot})/E_0$ for the resonance excited at $\lambda = 25.4$ $\mu$m. The fields are calculated in the horizontal plane 5 nm above the disk (corresponding to $z = 10$ nm, top panels) and in the $y=0$ plane of incidence of the incoming illumination going through the center of the disk (bottom panels). We can see that the fields are symmetric with respect to the $z = 0$ horizontal plane. In consequence (see Eq. 2), the induced charges are antisymmetric with respect to the same plane, i.e. same sign at opposing interfaces [insets in Fig. 6(a)], as anticipated for the excitation of an optical mode. Furthermore, both the charges and fields are antisymmetric along $x$, with a cos-like symmetry in the polar direction. Last, the field distribution directly over the disk present 1 and 3 nodes along the $x$ axis for the peaks at $\lambda = 34.3$ $\mu$m and $\lambda = 25.4$ $\mu$m, respectively. Therefore, the lowest energy plasmon at $\lambda = 34.3$ $\mu$m corresponds to a dipolar optical mode, and the resonance at $\lambda = 25.4$ $\mu$m is a higher order optical mode. Our results thus confirm the excitation of tunable optical plasmonic resonances in TI disks by an incident plane wave.

**Excitation of a localized acoustic mode by a point dipole**

We consider next a disk of almost ten times smaller diameter $D = 40$ nm and same $d = 10$ nm thickness. Using this smaller diameter allows to obtain the lowest-
The mode volume, as set by the lateral dimensions of the disk and by $k_2^2$, respectively. Applying the simple equation for the Purcell Factor $P_F = 3/(4\pi^2)\lambda^3Q/V_{\text{eff}}$, we estimate an effective volume $V_{\text{eff}} \approx 120\,\text{nm}^3$. Notably, we have assumed in Fig. 6 that the system remains on the weak coupling regime, where the effect of the nanodisk on a dipolar source is well described by the Purcell Factor. However, such small $V_{\text{eff}}$ indicate that it could be possible to reach the strong coupling regime even for sources with relatively small dipolar strength[75].

Fig. 7(b) shows $\text{Im}(E_z^{\text{tot}})$ for the lowest-energy peak in the $E_F = 250\,\text{meV}$ spectra. The left and right panels represents, respectively, the fields in the horizontal plane $z = 10\,\text{nm}$ (5 nm above the top interface) and the vertical plane $y = 0$ panel (same planes as in Fig. 6). The strength of the dipole is $1\,\text{e} \cdot \text{nm}$. The main difference between the results obtained for the larger disks under plane wave illumination (Fig. 6) and for the smaller disk excited by a dipole (Fig. 7(b)) is that the fields induced in the latter case are symmetric along the vertical direction, i.e. with respect to the central $z=0$ horizontal plane. According to Eq. 2, the symmetric fields correspond to anti-symmetric charge distribution (sketched in the inset of Fig. 7(a)) confirming the excitation of acoustic modes. Furthermore, the charges (as also $E_z^{\text{in}}$) are antisymmetric with respect to $x = 0$, with the charges excited at the upper interface forming a dipole-like pattern, which reminds the results for the large disks in Fig. 6. However, as the sketch in Fig. 7 indicates, the charges exited at the bottom interface correspond to a dipole-like pattern of opposite sign than the one at the top. Thus, the overall distribution of the induced charges is quadrupolar. A quadrupole is weakly radiative (non-radiative in the ideal case) which is a different way of arguing why plane-waves
do not couple efficiently with this acoustic mode.

We have thus shown that, by choosing adequately the size and illumination of the disks, we can excite modes that have either acoustic or optical character. In a similar manner as shown in Fig. 5 for the propagating waves, these localized modes would be dominated, respectively, by the value of the effective total spin density $\sigma_{\text{eff}}^{s}$ or total charge density $\sigma_{\text{eff}}^{q}$. Modifying $E_F$ allows to tune the resonant energy to a desired value.

**SUMMARY AND DISCUSSION**

We have combined many-body theory and classical electrodynamic calculations to analyze in detail the excitation of propagating and localized plasmons supported by thin TI structures. To focus on the pure electrodynamic interactions between the plasmons at the upper and bottom interfaces, we consider a simple case where the bulk of the topological insulator is perfectly transparent. Thus, we do not include the contribution from bulk plasmons [76] or from phonons[20, 25, 27, 46]. These effects can increase the losses and modify the dispersion, possibly introducing coupling between surface and bulk modes that can lead to phenomena such as Fano resonances or strong coupling. In the case of thin disks, edge effects[10, 77] and the complex atomic structure of the TI side surfaces[78], where a 2D-FL may form, could also shift and broaden the resonances. We also do not consider the depletion layer[36]. On the other hand, we have introduced the dependence of the 2D-conductivity on the parallel wave-vector $q$ in order to show how the strong non-locality of the response affects the optical response.

Thin TI films support optical and acoustic propagating plasmon modes. Acoustic plasmons are associated with net effective spin and vanishing total charge densities, while the optical mode is characterized by net charge oscillations with insignificant effective spin density. We have studied how to excite this system, showing that the experimental condition strongly affects the contribution of these two types of modes to the total signal. Specifically, by changing the distance between a point-like dipolar source and the TI, we are able to strongly modify the relative weight of the acoustic or the optical plasmons, and thus to control the spin and charge character of the propagating surface wave. The dipolar excitation we considered could come from a 2-level transition in single photon emitters, but finding appropriate transitions at these frequencies can be challenging. A tip, a small particle or any edge might serve as alternative localized sources[79, 80]. It is also worth noting here that the approach of coupling double-2D-FL structures to control two different properties of the excited surface wave can be extended to other situations. For example, Ref. 81 discusses the separation of purely magnetic and charge plasmons in two twisted bilayer graphene structures.

Further, we show that, by controlling the geometry and illumination, localized acoustic or optical plasmons can be selectively excited in thin TI disks at well-defined wavelengths. These modes are confined to very small volumes, as desired for large coupling strengths and quantum applications [82–84]. We have focused on separately exciting either acoustic or optical modes in the thin disks, but we have also verified that it is possible to excite both types of modes in the same disk at a similar frequency, (for example by exciting $D = 95\,\text{nm}$ disks in the $\lambda = 28\,\mu\text{m}$ range with a dipolar excitation). Additionally, we have assumed that the Fermi energy $E_F$ is the same for the 2D-FLs of the thin structures, but it may be also of interest to explore the case where the applied voltage leads to two different $E_F$.

Notably, the modes supported by the thin disks are relatively narrow spectrally, and can be easily tuned by changing the geometry or the external voltage. They thus offer an attractive path to control the nett spin and charge excited in nanostructures, by selectively exciting acoustic or plasmonic modes, or both simultaneously. The possible excitation of a well-defined net spin is a key difference with respect to other similar systems, such as bilayer graphene.

In summary, we have demonstrated the large flexibility offered by thin TIs to control spin and charge properties of plasmonic resonances, making them an attractive possibility to engineer very compact and fast [27] optoelectronic and spintronic devices.

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APPENDIX A: MANY-BODY CALCULATIONS

We describe first the many-body calculation of the conductivity $\sigma(q, \omega)$ required by the classical calculations (Appendix B) and obtained for a single 2D-FL between the TI and vacuum. We express the conductivity in terms of the non-interacting response function $\chi^0(q, \omega)$ of a single 2D-FL as

$$\sigma(q, \omega) = i\omega \frac{\chi^0(q, \omega)}{q^2}. \quad (4)$$

In the relaxation-time approximation, the non-interacting response function reads[85]

$$\chi^0(q, \omega) = \frac{1 + i/\omega\tau}{1 + (i/\omega\tau)^2} \frac{\chi^0(q, \omega + i/\tau)}{\chi^0(q, 0)}. \quad (5)$$

For the TI modeled in our study, we choose a relaxation time of $\tau = 500$ fs.[37–40] Within the random phase approximation, the response function $\chi^0(q, \omega)$ has the same expression as in the case of graphene,[86, 87] but without degeneracy neither in valley nor in spin.

We describe next how to get the dispersion relationship directly from this many-body approach, for a general system with an arbitrary number of 2D-FLs that interact via Coulomb coupling. In contrast to the classical electrodynamic formalism used in much of this work, where the interaction between 2D-FLs is captured via the full Maxwell’s equations, the many-body approach ignores retardation of the fields, which should be valid for the extremely thin systems under consideration.

Each 2D-FL is surrounded by dielectric media with certain bulk dielectric constant and it is localized in the vertical $z$-direction, with the localization $|\lambda_i(z)|$ (where $i$ runs over the layers) given by

$$|\lambda_i(z)|^2 = \frac{2}{h_i} \sin^2 \left( \frac{\pi (z^0_i - z)}{h_i} \right) \theta(h_i - (z^0_i - z)) \theta(z^0_i - z). \quad (6)$$

$h_i$ is the width of the respective 2D-FL in the $z$ direction, while $z^0_i$ sets its upper plane. The width $h_i$ accounts for the extension of the TI surface state in the direction normal to the surface plane.

The non-interacting response function of the 2D-FLs as a function of wavenumber reads[88]

$$\chi^0_i(q; z, z') = \sum_i |\lambda_i(z)|^2 \chi^0_i(q; \omega)|\lambda_i(z')|^2, \quad (7)$$

As a result, the interacting response function can be found as

$$\chi^\tau_i(q; z, z') = \sum_{i,j} |\lambda_i(z)|^2 \chi^\tau_{ij}(q; \omega)|\lambda_j(z')|^2, \quad (8)$$

where

$$\chi_{ij}(q, \omega) = \chi^0_i(q, \omega) \delta_{ij} + \chi^0_i(q, \omega) W_{ij}(q, \omega) \chi^0_j(q, \omega) \quad (9)$$

with the Kronecker delta $\delta_{ij}$ and the screened interaction $W_{ij}(q)$ between $i$-th and $j$-th 2D-FLs defined as

$$W_{ij}(q, \omega) = \left[ (1 - U \chi^0_i)^{-1} U \right]_{ij}. \quad (10)$$

Here the matrix $\chi^0_i$ is considered as a diagonal matrix with the elements $\chi^0_i \delta_{ij}$. The elements of the matrix $U$ are given by

$$[U]_{ij} \equiv U_{ij} = \int dz dz' |\lambda_i(z)|^2 \phi(q; z, z') |\lambda_j(z')|^2. \quad (11)$$

The function $\phi(q; z, z')$ entering this equation is the Coulomb interaction as obtained from the Poisson equation with the $z$-dependent dielectric constant $\varepsilon(z)$ corresponding to the considered geometry. Finally, the energies of the collective excitations in the set of 2D-FLs are defined by locating the poles of $\chi^\tau$ of Eq. (8). We find these poles from the zero crossing of $\text{det}[1 - U \chi^0_i]$. In the case of the semi-infinite substrate, we place only one 2D-FL with a non-zero thickness $\sim 10$ Å such as its upper plane coincides with the TI surface. When considering the TI film, we add one more 2D-FL of the same thickness and charge density on the other side of the film, where the lower plane of this layer is superimposed with the bottom of the TI film. Thus, the 10 nm thick TI insulator layer studied in the text includes the thickness of both 2D-FLs.

APPENDIX B: CLASSICAL CALCULATIONS

We describe next in more details how the classical calculations are performed once $\sigma(q, \omega)$ is obtained from the many-body results. In the case of thin structures with two 2D-FLs, we assume[89, 90] that the layers are sufficiently far from each other to neglect inter-layer transitions, so that we can directly use the conductivity $\sigma(q, \omega)$ obtained for a single interface as described in Appendix A. For our symmetric thin structures $\sigma(q, \omega)$ is the same for both TI-vacuum interfaces. The 2D-FLs interact via electromagnetic (Coulomb) coupling.

For these calculations, the bulk TI and the surrounding vacuum are characterized by their permittivity ($\varepsilon_1 = 1$ and $\varepsilon_0 = 25$, respectively), so that we ignore any phononic mode. We also neglect the correction to the electromagnetic constitutive equations due to induced currents at the 2D-FLs [47, 48] as we expect that this correction will only introduce second-order effects to the main phenomena studied here[17, 91].

For the calculation of the response of the substrate and thin film (Fig. 1(a,b)), which are illuminated by a point
to the optical mode. The bulk dielectric constant of the TI is characterized by the parallel and acoustic plasmon. A second peak appearing in the spectra in (a) at lower peaks in (a) associated with the acoustic plasmon. For the same incident wavelengths as in (a). The dashed vertical lengths. (b) Imaginary part of the non-local conductivity characterizing a 10 nm thin TI layer, for several incident wavelengths as in (a). The dashed vertical lines in (a) and (b) are visual guides corresponding to the peaks in (a) associated with the acoustic plasmon. A second peak appearing in the spectra in (a) at lower $q$ corresponds to the optical mode. The bulk dielectric constant of the TI is $\varepsilon_b=25$ and the Fermi energy $E_F=250$ meV

\[ r_{\text{TI}}(q, \omega) = \frac{\varepsilon_{\text{TI}}k_z^1 - \varepsilon_1 k_z^\text{TI}}{\varepsilon_{\text{TI}}k_z^1 + \varepsilon_1 k_z^\text{TI}} + \frac{4\pi\varepsilon_1 k_z^\text{TI}}{\omega} \]

where $k_z^1 = \sqrt{k_z^2 - q^2}$ and $k_z^\text{TI} = \sqrt{k_z^2 - q^2}$ are the normal components of the plane-wave at both sides of the interface. This calculation directly takes non-locality into account by adopting the corresponding conductivity $\sigma(q, \omega)$ for each plane-wave of parallel wavenumber $q$.

Besides giving the fields at all positions, the plane-wave decomposition allows to obtain the dispersion-relationship from the position of the resonances, which we then can compare with the result from the many-body calculations. More exactly, we calculate the dispersion relationship from the classical approach by calculating for each frequency of interest the parallel plasmon wavenumber $q_{\text{SP}}$ that results in a larger transmission $t_p(q, \omega)$ of the full system. With this approach it is also possible to calculate the dispersion for the often-used local approximation, by simply using the conductivity $\sigma(q=0, \omega)$ for all plane-waves.

In the case of the full-wave simulations of the disks[53], it is possible to include a 2D-layer at both the upper and bottom interfaces. However, the conductivity $\sigma$ only depends on $\omega$. We are able to introduce the effect of non-locality in an approximate manner because, for the situations under study, the optical response should be mostly determined by the response at one particular plasmonic parallel wavenumber $q_{\text{SP}}$. $q_{\text{SP}}$ can correspond to the parallel wavenumber of either the optical ($q_{\text{op}}^\text{SP}$) or acoustic ($q_{\text{ac}}^\text{SP}$) modes (obtained from the dispersion relationship of the thin film), so that the calculations use $\sigma(q_{\text{op}}^\text{SP}, \omega)$ or $\sigma(q_{\text{ac}}^\text{SP}, \omega)$. For simplicity, we only consider the 2D-layer at the flat top and bottom interfaces, and not at the lateral side surfaces. The latter correspond to the so-called side faces (other than the (111) surface) of Bi$_2$Se$_3$ (a strong TI), which have surface state bands characterized by tilted anisotropic Dirac cones with notably reduced Fermi velocities and by spin-orbital texture than entirely differs from that of the (111) surface[93, 94]. As a consequence, the disk lateral sides would be described by a different conductivity and a possibly shorter relaxation time. To take into account the specific characteristics of the surface states on strong TI side faces, a proper approximation to the non-interacting response function should be developed first. However, we expect that it may introduce a significant shift on the peak position but should not affect the main physics described in this paper.

We examine next in more detail this approach to introduce non-locality into full-wave simulations. According to the discussion in the text, the localized acoustic and optical modes in the disks can be understood in terms of the excitation of propagating plasmons, which get re-
lected at the edges and lead to Fabry-Pérot-like resonances. Thus, to better understand which wavenumbers dominate the response of the disk, we examine the behavior in $q$-space of the thin films.

With this purpose, we plot in Fig. 8(a) the $q$-dependence of the absolute value of the transmission coefficient for the 10 nm thin TI disks, for different illumination wavelengths. Each spectrum displays two peaks, corresponding to the acoustic mode at the largest $q$, and the optical at the lowest $q$. If the following two conditions are verified, we should be able to use in the full-wave calculations the conductivity at either $q_{SP}^{ac}$ or $q_{SP}^{op}$: i) the response should be dominated by the $q$-components of one of the peaks, and ii) $\sigma(q)$ should not change strongly over the width of the peak.

To fulfill the first condition, we either use a plane-wave to illuminate the structure, which does not excite the acoustic modes (large disks simulations), or we choose a frequency region where optical modes are not resonantly excited (calculations of the smaller disks).

To study in more detail the second condition, Fig. 8(b) shows the behavior of the imaginary part of $\sigma(q, \omega)$ for the same excitation wavelengths as in Fig. 8(a) (the real part gives similar conclusions). In general, the spectra varies relatively slowly with the parallel wavenumber, except for the presence of a clear resonant feature at a $q$ that depends on the excitation frequency and that is related to the edge of the electron-hole continuum (smoothed in the presence of non-zero losses). For excitation wavelengths $\lambda < 50 \mu m$, the relatively flat region of the conductivity extend over the $q$-values corresponding to both the acoustic and optical peaks in 8(a). Thus, we expect our results in Fig. 6, 7 to be a reasonable approximation of the non-local response.

[1] W. L. Barnes, A. Dereux, and T. W. Ebbesen, Nature 424, 824 (2003).
[2] J. Pitarke, V. Silkin, E. Chulkov, and P. Echenique, Reports on progress in physics 70, 1 (2006).
[3] M. Pelton, J. Aizpurua, and G. Bryant, Laser & Photonics Reviews 2, 136 (2008).
[4] M. Orłita and M. Potemski, Semicond. Sci. Technol. 25, 063001 (2010).
[5] L. Ju, Baisong, J. Horng, C. Girit, M. Martin, Z. Hao, H. A. Bechtel, X. Liang, A. Zettl, Y. R. Shen, et al., Nat. Nanotechnol. 6, 630 (2011).
[6] A. Nikitin, P. Alonso-González, S. Vélez, S. Mastel, A. Centeno, A. Pesquera, A. Zurrutuza, F. Casanova, L. Hueso, F. Koppens, et al., Nat. Photon. 10, 239 (2016).
[7] Z. Fang, Y. Wang, A. E. Schletter, Z. Liu, P. M. Ajayan, F. J. García de Abajo, P. Nordlander, X. Zhu, and N. J. Halas, Nano lett. 14, 299 (2013).
[8] H. Yan, X. Li, B. Chandra, G. Tulevski, Y. Wu, M. Freitag, W. Zhu, P. Avouris, and F. Xia, Nature nanotechnol. 7, 330 (2012).
[9] S. A. Mikhailov, Phys. Rev. B 84, 045432 (2011).
[10] F. J. García de Abajo, ACS Photonics 1, 135 (2014).
[11] F. H. L. Koppens, D. E. Chang, and F. J. García de Abajo, Nano Lett. 11, 3370 (2011).
[12] J. Chen, M. Badioli, P. Alonso-Gonzalez, S. F. H. Thongrattanasiri, J. Osmond, M. Spasenovic, A. Centeno, A. Pesquera, P. Godignon, A. Z. Elorza, et al., Nature 487, 77 (2012).
[13] M. Jablan, H. Buljan, and M. Soljacic, Phys. Rev. B 80, 245435 (2009).
[14] Z. Fei, A. S. Rodin, G. O. Andreev, W. Bao, A. S. McLeod, M. Wagner, Zhang, L. M., Z. Zhao, M. Thiennes, et al., Nature 487, 82 (2012).
[15] A. Manjavacas and F. J. García de Abajo, Nat. Commun. 5, 3548 (2013).
[16] H. Zhang and S. C. Zhang, Nature Phys. 5, 438 (2009).
[17] R. Schütky, C. Erterl, A. Trügler, and U. Hohenester, Phys. Rev. B 88, 195311 (2013).
[18] D. Hsieh and M. Z. Hasan, Nature 460, 1101 (2009).
[19] Y. P. Lai, I. T. Lin, K. H. Wu, and J. M. Liu, Nanomater. Nanotechnol. 4, 1 (2014).
[20] S. Sim, H. Jang, N. Koirala, M. Brahelek, J. Moon, J. H. Sung, J. Park, S. Cha, S. Oh, M.-H. Jo, et al., Nat. Commun. 6 (2015).
[21] M. Z. Hasan and C. L. Kane, Rev. Mod. Phys. 82, 3045 (2010).
[22] C. L. Kane and E. J. Mele, Phys. Rev. Lett. 95, 226801 (2005).
[23] C. Wu, B. A. Bernevig, and S. C. Zhang, Phys. Rev. Lett. 96, 106401 (2006).
[24] H. Z. Lu, W. Y. Shan, W. Y. Han, S. M. Ruan, and S. Q. Shen, Phys. Rev. B 81, 115407 (2010).
[25] P. Di Pietro, M. Tortolani, O. Limaj, A. Di Gasparr, V. Gliberti, Giorgianni, Brahelek, N. Bansal, N. Koirala, S. Oh, et al., Nat. Nanotechnol. 8, 556 (2013).
[26] J. Y. Ou, J. K. So, G. Adamo, A. Sulaev, L. Wang, and N. I. Zheludev, Nat. Commun. 5, 5139 (2014).
[27] M. Autore, H. Engelkamp, F. DApuzzo, A. D. Gasparr, P. D. Pietro, I. L. Vecchio, M. Brahelek, N. Koirala, S. Oh, and S. Lupi, ACS Photonics 2, 1231 (2015).
[28] Y. Deshko, L. Kruis-Eilbaum, V. Menon, A. Khanikaev, and J. Trevino, Opt. Express 24, 7398 (2016).
[29] J. Guozhi, W. Peng, Z. Yanbang, and C. Kai, Sci. Rep. 6, 25884 (2016).
[30] S. Raghur, S. B. Chung, X. L. Qi, and S. C. Zhang, Phys. Rev. Lett. 104, 116401 (2010).
[31] T. Stauben, J. Phys. Condens. Matter 26, 123201 (2014).
[32] R. E. V. Profumo, R. Asgari, M. Polini, and A. H. MacDonald, Phys. Rev. B 85, 085443 (2012).
[33] T. Stauben and G. Gómez-Santos, New Journal of Physics 14, 105018 (2012).
[34] E. H. Hwang and S. Das Sarma, Phys. Rev. B 80, 205405 (2009).
[35] T. Stauben, G. Gómez-Santos, and L. Brey, Phys. Rev. B 88, 205427 (2013).
[36] T. Stauben, G. Gómez-Santos, and L. Brey, ACS Photonics 4, 2978 (2017).
[37] D.-X. Qu, Y. S. Hor, J. Xiong, R. J. Cava, and N. P. Ong, Science 329, 821 (2010).
[38] J. Qi, X. Chen, W. Yu, P. Cadden-Zimansky, D. Smirnov, N. H. Tolik, I. Miotkowski, H. Cao, Y. P. Chen, Y. Wu, et al., Appl. Phys. Lett. 97, 182102 (2010).
[39] Y. D. Glinka, S. Babakariy, T. A. Johnson, A. D. Bristow, M. B. Holcomb, and D. Lederman, Appl. Phys. Lett. 103,
