Optical excitations in electron microscopy

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(Dated: March 10, 2009)

This review discusses how low-energy, valence excitations created by swift electrons can render information on the optical response of structured materials with unmatched spatial resolution. Electron microscopes are capable of focusing electron beams on sub-nanometer spots and probing the target response either by analyzing electron energy losses or by detecting emitted radiation. Theoretical frameworks suited to calculate the probability of energy loss and light emission (cathodoluminescence) are revisited and compared with experimental results. More precisely, a quantum-mechanical description of the interaction between the electrons and the sample is discussed, followed by a powerful classical dielectric approach that can in practice be applied to more complex systems. We assess the conditions under which classical and quantum-mechanical formulations are equivalent. The excitation of collective modes such as plasmons is studied in bulk materials, planar surfaces, and nanoparticles. Light emission induced by the electrons is shown to constitute an excellent probe of plasmons, combining sub-nanometer resolution in the position of the electron beam with nanometer resolution in the emitted wavelength. Both electron energy-loss and cathodoluminescence spectrosopies performed in a scanning mode of operation yield snap shots of plasmon modes in nanostructures with fine spatial detail as compared to other existing imaging techniques, thus providing an ideal tool for nanophotonics studies.

PACS numbers: 68.37.Lp,79.20.Uv,78.60.Hk,73.20.Mf

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I. INTRODUCTION

Investigating minute physical processes has been essential for advancing knowledge and generating the right questions all the way from the beginning of modern science up to recent developments in nanotechnology. Far-field optical microscopes have contributed to this end, but they are limited by diffraction to a spatial resolution of about half the light wavelength in practice. Actually, the best resolution (< 1 Å) is currently achieved using electrons rather than light (Nellist et al., 2004; Ruska, 1987), for which the de Broglie wavelength (de Broglie, 1925) is well below 0.1 Å at energies above 20 keV.

Besides acquiring static images of the nanoworld, we are interested in finding out about the excitations that small objects can sustain, which inform us on their dynamical evolution and are relevant for encoding and manipulating information and for exploring a myriad of applications in fields such as molecular biology. Thus, our ultimate goal is to perform spectroscopy at the shortest possible length scale with the highest energy resolution. With this focus in mind, we can classify the available experimental techniques as shown in Fig. 1, clearly indicating that electron-based spectroscopies offer the best choice for optimizing spatial resolution.

Two routes have been devised so far for dealing with the diffraction limit: \(^1\) (1) to reduce the wavelength of the probe and (2) to employ near-field detection techniques. (1) Moving to shorter photon wavelengths is difficult due to the lack of versatile lenses and mirrors beyond the ultraviolet. Nonetheless, recent advances in x-ray microscopy (XRM) have allowed 15 nm imaging resolution (Chao et al., 2005) at energies above 250 eV. Alternatively, shifting from photons to electrons of the same energy encompasses a significant decrease in wavelength and improved spatial resolution. \(^2\) (2) Exploiting the near field is another option, particularly when relying on evanescent components, the fast decay of which provides an extra handle for enhancing resolution using localized probes. For instance, near-field scanning optical microscopy (NSOM, based on a subwavelength tip at the end of a fiber that brings, collects, or scatters light) and tip-enhanced NSOM can push spatial resolution down to tens of nanometers (Betzig et al., 1992; Hartschuh, 2008). Even better detail in the sub-Å domain is routinely achieved using the formerly developed scanning tunneling microscope (STM), on which scanning tunneling spectroscopy (STS) yields 0.1 eV energy resolution (Hörmandinger, 1994). However, STS only works with metals, and its extreme spatial accuracy (~0.01 Å in the vertical direction) requires close proximity between the tip and the sample surface, so that single-electron excitations dominate the spectra and mask collective modes relevant to optics, such as plasmons.

Electron microscopes are thus the best option for resolving both localized and extended excitations with subnanometer spatial detail and < 0.1 eV energy resolution in any type of material (Lazar et al., 2006). These instruments are sufficiently versatile to be surface-sensitive and to simultaneously procure information on bulk properties. Their performance has considerably improved in recent years due to extraordinary advances in energetic electron sources and optics. A number of studies in different fields benefit from the unprecedented spatial resolution of electron microscopy, which no other technique can currently match.

Electron microscopy has different flavors depending on the type of signal that we measure [inelastic losses, cathodoluminescence (CL) light emission, secondary electrons (SEs), etc.]. Several widely available types of setups are schematically represented in Fig. 2. The transmission electron microscope (TEM) provides by far the

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\(^1\) The existence of a diffraction limit has been questioned by recent investigations of super-oscillating functions (Zheludev, 2008).

\(^2\) Heavier particles such as protons have been argued to provide good spatial resolution compared to electrons (Demkov and Meyer, 2004). Actually, a helium ion microscope has been recently released (Ward et al., 2006) and achieved 2.4 Å resolution (Carl Zeiss 2008 announcement).
most powerful combination of spectral and energy resolution, mainly via analysis of loss events. This type of microscope can operate like a conventional optical microscope, projecting a bright image of an object on a phosphor screen or a CCD camera to picture the magnified sample. Energy-filtered TEM (EFTEM) allows constructing images out of those electrons that have donated a determined amount of energy to the specimen, which is useful to visualize selected losses [for instance, core excitations identifying the chemical environment of atomic species (Batson, 1993)]. Excellent spatial resolution is obtained using TEMs operated in scanning mode (STEM), in which the beam is focused and scanned over the sampled area, either to form images out of large-momentum transfers recorded by annular dark-field electron detectors or to collect loss spectra at specific locations.

Most TEMs use energetic electrons in the range of 50-400 keV and require very thin specimens (≤100 nm) that are relatively transparent to these electrons. Sample preparation is thus an important issue for achieving optimum spatial resolution.

In contrast, scanning electron microscopes (SEMs) can work with any sample that is covered by a thin metal layer (∼1 − 2 nm), forming images by collecting SEs upon bombardment with a primary beam of ≥2 nm in diameter. Furthermore, one can perform spectroscopy on SEMs through wavelength-resolved CL emission. Although CL spectroscopy can also be collected on a TEM (Yamamoto et al., 1996a), it is most commonly available in SEMs equipped with a light collector system (e.g., an ellipsoidal mirror) and a photon spectrometer.

Low-energy electron microscopy (LEEM) (Rocca, 1995) relies on the use of 1-100 eV electrons and can achieve < 50 nm spatial resolution accompanied by sub-electronvolt energy resolution. As an example of application, LEEM has been recently used to observe acoustic plasmons (Diaconescu et al., 2007). In contrast, photoemission electron microscopy (PEEM) (Bauer, 1994) is particularly useful to map the light intensity in illuminated nanostructures, as recently demonstrated for lithographically patterned Ag nanoparticle arrays (Aeschlimann et al., 2007; Cinchetti et al., 2005). The femtosecond dynamics of plasmon excitations has been also resolved using multiphoton PEEM with pump-probe illumination (Kubo et al., 2005, 2007).

Finally, there are other configurations that make use of electrons to perform spectroscopy. For instance, reflection electron energy-loss spectroscopy (REELS) has been recently used to determine optical properties of noble metals (Went et al., 2008; Werner, 2006; Werner et al., 2007) and graphite (Calliari et al., 2008) after careful data analysis.

![FIG. 2 (Color online) Schematic representation of various types of electron microscopes equipped to perform spectroscopy. (a) Scanning transmission electron microscopes (STEMs) allow analyzing the energy distribution of electrons transmitted through a thin specimen (∼100 nm thick), on which the incident beam is focused down to a < 0.1 nm spot. Large-momentum transfers are collected through annular dark-field detectors (ADFs). Cathodoluminescence (CL) light detection is also possible on a STEM. (b) Scanning electron microscopes (SEMs) collect secondary electrons (SEs) to form images or CL emission to perform spectroscopy, with spatial resolution down to ~2 nm. (c) Photoemission electron microscopes (PEEMs) can render spectroscopic information on the photo-excitation process and electron-hole (e-h) dynamics with ≤50 nm resolution.]

A. Spectroscopy using electron microscopes

A swift electron impinging on a microscope specimen gives rise to secondary electron emission (SEE) and CL that can be energy-analyzed, thus yielding information on the excitation spectrum of the sample. Electron energy-loss spectroscopy (EELS) performed in STEMs adds up to this suite of probes. The main advantage of electron microscopes is that these types of spectroscopies can be performed with truly nanometer spatial resolution.

Unlike freely propagating light, the bare field of an electron is evanescent, as we show in Sec. II.A. This is advantageous to investigate localized excitations, involving wave-vector components of the electromagnetic field that lie outside the light cone. For example, electron beams were instrumental in discovering and characterizing collective excitations of conduction electrons in metals (Chen and Silcox, 1975b; Powell and Swan, 1959; Raether, 1980; Ritchie, 1957; Rutherford, 1948; Watanabe, 1956), known as plasmons because they are quasi-particles associated to oscillations of the electron gas plasma. Specifically, bulk plasmons in a homogenous metal are conspicuous in EELS, since they are electro-
static and longitudinal in nature, and for this reason they couple very efficiently to moving charges.

Similarly, plasmons can be confined at the surface of a metal, and they are actually the source of interesting phenomena and applications that configure the field of plasmonics (see Sec. VI). Surface plasmons (SPs) are versatile entities: they can be trapped in metal particles that are much smaller than the wavelength (Myroshnychenko et al., 2008b), and they can hybridize with light extending over larger metallic structures (Coyle et al., 2001). More precisely, surface plasmon polaritons (SPPs), which are a subset of SPs capable of propagating on planar surfaces (Barnes et al., 2003; Ozbay, 2006; Zia et al., 2006) or along 1D waveguides (Bozhevolnyi et al., 2006), hold great promise to become the natural link between current nanoelectronics and future integrated nanophotonics, operating at frequencies that are $\sim 10^5$ times higher than microchip clocks.

As illustrated in Fig. 3, the passage of a fast electron can excite localized plasmons in metallic nanoparticles, but it can also launch SPPs in planar metallic surfaces (Bashevoy et al., 2006; Heitmann, 1977; van Wijngaarden et al., 2006) or in metal nanowires (Vesseur et al., 2007).

Quite different from SPPs, localized plasmons can decay radiatively, thus contributing to CL, although all sorts of plasmons can partially decay via inelastic channels that involve electronic excitations, including electron-hole (e-h) pair creation and SEE if the electron is near the surface and its energy above the vacuum level. We discuss this matter more thoroughly in Sec. IV.A.

The electron can directly excite e-h pairs too, the decay of which gives rise to the emission of CL and SEs. This produces in general a complex cascade of SEs, which a transport equation approach is suited to model reliably (Rössler and Brauer, 1991). The cascade includes energetic electrons that generate further SEs and CL, adding complexity to this scenario. However an element of simplicity comes from the separation between coherent and incoherent processes, from the point of view of the emitted light, as we explain in more detail in Sec. IV. In particular, the coherent CL signal is dominant in metals.

We are interested in low-energy excitations, which involve electrons and holes in the valence and conduction bands, as well as collective modes (e.g., localized plasmons and SPPs). This is by far the most intense part of the loss spectrum (see Fig. 4). Its analysis yields information on the materials optical response with the kind of spatial resolution that is currently dreamed of in the context of nanophotonics. It is our purpose to review historical developments, to summarize recent advances in this area, to present an overview of theoretical methods, and to point out some opportunities opened by electron microscopy in order to expand and complement nanophotonics studies in a way that can be particularly beneficial for emerging areas such as plasmonics.

II. INTERACTION OF SWIFT ELECTRONS WITH MATTER

We first consider some fundamental aspects of the interaction between fast electrons a matter, as well as theoretical approaches suitable to simulate EELS.

A. An evanescent source of light in matter

The electromagnetic field that accompanies a point charge moving in vacuum can be regarded as an evanescent source of radiation which permits exploring regions of momentum-energy space that lie outside the light cone. This has interesting consequences: fast electrons generate SPPs when passing near a metal surface, as we show in Sec. III.C, but they can also excite non-dipolar modes in small particles (Chu et al., 2009), which would be dif-
Direct solution of Maxwell’s equations yields a straight-line trajectory with constant velocity vector \( \mathbf{v} \) and stronger interaction in low-index dielectrics), and a term proportional to the velocity vector in the numerator, which adds transversal components to the field.

It is helpful to examine first the electric field produced by a swift charged particle moving inside a homogeneous medium. The field can be conveniently decomposed into different frequency contributions using the Fourier transform

\[
E(r, t) = \int \frac{d\omega}{2\pi} E(r, \omega) e^{-i\omega t},
\]

and also into momentum components with \( \exp(i\mathbf{q} \cdot \mathbf{r}) \) spatial dependence. We assume that the electron describes a straight-line trajectory with constant velocity vector \( \mathbf{v} \), crossing the origin at time \( t = 0 \). This is a reasonable assumption for the swift electrons and relatively thin specimens typically examined with TEMs. Then, the electron charge density becomes \(-2\pi\epsilon\delta(\omega - q \cdot \mathbf{v})\) in \( \mathbf{q} - \omega \) space. Direct solution of Maxwell’s equations yields

\[
E(r, \omega) = \frac{ie}{\pi} \int d^3q \frac{q/\epsilon - kv/c}{q^2 - k^2\epsilon} e^{i\mathbf{q} \cdot \mathbf{r}} \delta(\omega - q \cdot \mathbf{v}),
\]

where \( k = \omega/c \) is the light wavenumber in free space and \( \epsilon \) is the dielectric function of the homogeneous medium. It should be noted that \( \epsilon \) can depend on both \( \mathbf{q} \) and \( \omega \). The consequences of the wavevector dependence are discussed later in Sec. III.B, but they are ignored in the remainder of this section.

Equation (2) contains some interesting elements. Retardation effects show up both through \( k^2\epsilon \) in the denominator, which modifies the strength of the field (producing, for example, weaker interaction in metals and stronger interaction in low-index dielectrics), and through a term proportional to the velocity vector in the numerator, which adds transversal components to the field.

The delta function inside the integral of Eq. (2) expresses energy conservation for transfers of frequency \( \omega \) and wavevector \( \mathbf{q} \) from the electron to the material. Neglecting relativistic corrections, the electron energy is reduced from \( m_e v^2/2 \) to \( m_e v - hq^2/2m_e \) during the transfer, so that the energy difference \( h\omega \) leads to

\[
\omega = q \cdot \mathbf{v} - hq^2/2m_e \approx q \cdot \mathbf{v}.
\]

The right-most expression is the non-recoil approximation, which works extremely well under the usual condition \( hq \ll m_e v \). This approximation remains valid if the energy transfer is computed using relativistic expressions. The kinematically allowed transfers span a solid area for all possible relative orientations between \( \mathbf{v} \) and \( \mathbf{q} \), as shown by the shaded region in Fig. 5 with upper boundary \( \omega = qv \).

The zeros of the denominator in the integrand of Eq. (2) signal the dispersion relation of light in the medium, \( q = k\sqrt{\epsilon} \). This has been represented in Fig. 5 for a polaritonic material described by the dielectric function of Eq. (A3) (thick solid curves). The figure shows two different frequency domains separated by the condition \( v^2\epsilon = c^2 \) (point C in Fig. 5 is a graphical solution of this equation). At lower frequencies with respect to C, the electron does not couple to excitations in the medium and the spectral components of the electric field decay exponentially away from the trajectory, as explicitly shown by solving

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4 Electrons with typical TEM energies above 80 keV have velocities \( v > 0.5c \). When they undergo valence losses \( h\omega < 50\text{eV} \), the momentum transfer is \( hq \approx h\omega/v < 0.0004 m_e v \) along the direction of the trajectory.
announced exponential decay with \( \gamma \) where \( \gamma = 0.7c \) (\( \gamma \approx 1.4 \) and kinetic energy \( \approx 200 \text{ keV} \)) along the positive \( z \) axis. The only non-vanishing components \( (E_R, E_z, \text{ and } H_\phi) \) decay exponentially at large distance \( R \) from the trajectory. The inset shows the orientation of these components relative to the electron velocity vector. The small \( R \) limit is dominated by the \( 1/R \) divergence of \( E_R \) and \( H_\phi \).

the integral in Eq. (2). We find (Jackson, 1999)

\[
E(r, \omega) = \frac{2e\omega}{v^2 \gamma_\epsilon} g(r),
\]

where

\[
g(r) = e^{i\omega z/v} \left[ \frac{i}{\gamma_\epsilon} K_0 \left( \frac{\omega R}{v\gamma_\epsilon} \right) \hat{z} - K_1 \left( \frac{\omega R}{v\gamma_\epsilon} \right) \hat{r} \right].
\]

\( \gamma_\epsilon = 1/\sqrt{1 - e^2/\epsilon^2} \) is the Lorentz contraction factor\(^5\) and the notation \( r = (R, z) \) with \( R = (x, y) \) has been employed (see inset of Fig. 6), with the velocity taken along \( \hat{z} \). Similarly, the magnetic field reduces to

\[
H(r, \omega) = \frac{-2e\omega}{v\gamma_\epsilon} K_1 \left( \frac{\omega R}{v\gamma_\epsilon} \right) e^{i\omega z/v} \hat{\phi},
\]

where \( \hat{\phi} \) is the azimuthal unit vector (see Fig. 6). The announced exponential decay with \( R \) of both \( E(r, \omega) \) and \( H(r, \omega) \) arises from the asymptotic behavior of the modified Bessel functions \( K_m \) for large arguments (Abramowitz and Stegun, 1972). The non-vanishing components of these fields are represented in Fig. 6. The electromagnetic field extends up to distances of the order of \( \sim v\gamma c/\omega \), the Bohr cutoff. Notice however that one cannot assign this value to a characteristic Coulomb de-localization distance, since the field diverges at the origin as \( \sim 1/R \), so that large interaction contrast is expected across small distances in the region close to the trajectory.

When the electron moves faster than light in the medium, under the condition

\[
v > c/\sqrt{\epsilon},
\]

there is overlap between the photon dispersion relation and the shaded region in Fig. 5, so that the electron can emit Cherenkov radiation (CR) (see Sec. IV.D). The field displays oscillatory behavior and decays as \( 1/\sqrt{R} \) away from the trajectory [this stems from the modified Bessel functions \( K_0 \) and \( K_1 \) for imaginary argument in Eq. (5)].

It should be stressed that Eqs. (4)-(6) and Fig. 6 refer to each monochromatic component of the electromagnetic field, evolving with time as \( \exp(-i\omega t) \) [see Eq. (1)]. An electron moving in vacuum can be actually regarded as an evanescent source of super-continuum light with the spectral dependence shown in Fig. 6.

B. Classical dielectric formalism

The pioneering work of Fermi (1940) on the stopping of fast charged particles in dielectric materials opened up the application of classical electrodynamics to describe the interaction of swift electrons with matter. Following this useful tradition, we now extend the dielectric formalism of the previous section to cope with inhomogeneous samples and discuss in particular the spectral loss probability, which is relevant to EELS experiments.

The energy loss suffered by a fast electron moving with constant velocity \( \mathbf{v} \) along a straight line trajectory \( \mathbf{r} = \mathbf{r}_e(t) \) can be related to the force exerted by the induced electric field \( E_{\text{ind}} \) acting back on the electron as (Ritchie, 1957)

\[
\Delta E = e \int dt \mathbf{v} \cdot E_{\text{ind}}[\mathbf{r}_e(t), t] = \int_0^\infty \hbar \omega d\omega \Gamma_{\text{EELS}}(\omega),
\]

where the \(-e\) electron charge has been included (i.e., \( \Delta E > 0 \)) and

\[
\Gamma_{\text{EELS}}(\omega) = \frac{e}{\pi \hbar \omega} \int dt \{ e^{-i\omega t} \mathbf{v} \cdot E_{\text{ind}}[\mathbf{r}_e(t), \omega] \}
\]

is the so-called loss probability, which is given per unit of transferred frequency \( \omega \). The problem of calculating the loss probability reduces then to solving the electric field set up by the electron. A large deal of work has been devoted to obtaining the electric field for many geometries, including planar surfaces, isolated spheres, neighboring

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\(^5\) Interestingly, the Lorentz factor \( \gamma_\epsilon \) involves the velocity of light in the material, \( c/\sqrt{\epsilon} \), which is in turn frequency-dependent.

\(^6\) The square roots are chosen to yield positive real parts in this work. Notice that \( \text{Im}(\epsilon) \) is always positive in the retarded response formalism followed here, and it becomes a positive infinitesimal in non-lossy dielectrics.
spheres, circular cylinders, wedges, and more complex shapes, using both analytical and fully-numerical methods, either within the non-retarded approximation, based upon solutions of Poisson’s equation, or with full inclusion of retardation effects by solving Maxwell’s equations (see Sec. III and references therein). Next, we outline the general features of this formalism.

1. Non-retarded approximation

In the non-retarded approximation, we neglect the delay experienced by the electromagnetic signal that mediates the electron-sample interaction. Then, the electric field admits the form $\mathbf{E}(\mathbf{r}, \omega) = -\nabla \phi(\mathbf{r}, \omega)$, and we can disregard $\mathbf{H}$ in the absence of magnetic response.

It is useful to express the electric potential $\phi$ in terms of the screened interaction $W(\mathbf{r}, \mathbf{r'}, \omega)$, defined as the potential created at $\mathbf{r}$ by a unit point charge located at $\mathbf{r'}$ [an implicit $\exp(-i\omega t)$ time dependence is understood]. This quantity has to be combined with the charge density corresponding to the moving electron. Considering a non-coiled straight line trajectory, and assuming without loss of generality that the velocity vector is directed along the positive $z$ axis [i.e., the trajectory is $\mathbf{r}_e(t) = \mathbf{r}_0 + vt$ and $\mathbf{v} = e\mathbf{z}$], the electron charge density in frequency space $\omega$ reduces to

$$\rho(\mathbf{r}, \omega) = \frac{e}{v} \int dt \ e^{i\omega t} \delta(\mathbf{r} - \mathbf{r}_0 - \mathbf{v}t)$$

$$= \frac{e}{v} \delta(\mathbf{R} - \mathbf{R}_0) \ e^{i\omega(z-z_0)/v}, \quad (9)$$

where the notation shown in the inset of Fig. 6 has been used. From here, the potential reads

$$\phi(\mathbf{r}, \omega) = \frac{e}{v} \int dz' W(\mathbf{r}, \mathbf{R}_0, z', \omega) \ e^{i\omega(z'-z_0)/v}.$$ 

Finally, plugging these expressions into Eq. (8), the non-retarded (NR) loss probability is found to be

$$\Gamma_{\text{EELS}}^{\text{NR}}(\mathbf{R}_0, \omega) = \frac{e^2}{\pi \hbar^2} \int dz dz' \cos \left[ \frac{\omega(z-zA}{v} \right]$$

$$\times \ \text{Im} \left\{ -W(\mathbf{R}_0, z, \mathbf{R}_0, z', \omega) \right\}, \quad (10)$$

where the dependence of $\Gamma_{\text{EELS}}^{\text{NR}}$ on the electron impact parameter $\mathbf{R}_0$ is explicitly shown.\(^7\)\(^8\)

The loss probability can be thus derived from the knowledge of the frequency-dependent screened interaction $W$. A local description of the sampled materials in terms of a frequency- and space-dependent dielectric function $\epsilon(\mathbf{r}, \omega)$ provides often reliable results. Detailed expressions of $W$ are offered in Appendix B for planar, spherical, and cylindrical geometries. Nevertheless, Eq. (10) is valid beyond the local response approximation, and subtle effects such as local-field corrections can be incorporated via more realistic quantum-mechanical representations of the screened interaction.

2. Retardation effects

In high-voltage TEMs, retardation may become important because the speed of the charged projectiles is a sizeable fraction of the speed of light. For instance, the Lorentz contraction factor at 200 keV takes already a value significantly different from one, $\gamma = 1.4$. Retardation has two noticeable consequences for spectroscopy: (1) it increases the range of interaction of the electron probe in directions normal to the trajectory, as clearly illustrated by Eq. (5) and Fig. 6; and (2) it produces redshifts in excitation mode energies (Myroshnychenko et al., 2008b). The latter are discussed below in further detail, but we can anticipate that this effect turns out to be important when the excitations extend over specimen distances that cannot be neglected in front of the corresponding light wavelength. A proper description of these effects requires calculating the electric field of Eq. (8) from Maxwell’s equations in the presence of the moving electron and the sample under consideration.

The electromagnetic response of a structured material is fully captured in its electric Green tensor. In particular, the electric field produced by an external current density $\mathbf{j}(\mathbf{r}, \omega)$ in an inhomogeneous medium of permittivity $\epsilon(\mathbf{r}, \omega)$ can be written in frequency space $\omega$ as

$$\mathbf{E}(\mathbf{r}, \omega) = -4\pi i \omega \int d\mathbf{r}' \ G(\mathbf{r}, \mathbf{r}', \omega) \cdot \mathbf{j}(\mathbf{r}', \omega)$$

(11)

in terms of $G$, the electric Green tensor of Maxwell’s equations in Gaussian units, satisfying

$$\nabla \times \nabla \times G(\mathbf{r}, \mathbf{r}', \omega) - k^2 \epsilon(\mathbf{r}, \omega) G(\mathbf{r}, \mathbf{r}', \omega)$$

$$= \frac{-1}{\epsilon^2} \delta(\mathbf{r} - \mathbf{r}')$$

(12)

and vanishing far away from the sources ($|\mathbf{r} - \mathbf{r}'| \to \infty$ limit).

For the electron charge density of Eq. (9), the external current density reduces to $\mathbf{j} = v\rho$, which upon insertion into Eq. (11), and this in turn into Eq. (8), allows us to write the loss probability as

$$\Gamma_{\text{EELS}}(\mathbf{R}_0, \omega) = \frac{4e^2}{\hbar} \int dz dz' \cos \left[ \frac{\omega(z-zA}{v} \right]$$

$$\times \ \text{Im} \left\{ -G_{zz}[\mathbf{R}_0, z, \mathbf{R}_0, z', \omega] \right\}, \quad (13)$$

\(^7\) It should be noted that the reciprocity theorem $[W(\mathbf{r}, \mathbf{r'}, \omega) = W(\mathbf{r'}, \mathbf{r}, \omega)]$ and the fact that the bare Coulomb interaction is a real function have been utilized in the derivation of Eq. (10).

\(^8\) The induced field in Eq. (8) can be safely replaced by the total field, because the bare field of the moving charge does not produce stopping. We have accordingly dropped the superscript ind in Eq. (10).
where \( G_{zz} = \hat{z} \cdot G \cdot \hat{z} \).\(^9\) Interestingly, this expression works for any sign of \( v \), and therefore the loss probability is independent on whether the electron moves towards positive or negative \( z \)'s.

\[ \frac{d\Gamma_{\text{NR}}}{dt} = 2\pi e^2 \sum_{f,n} \left| \int d^3rd^3r' \frac{\psi_f^*(r)\psi_i(r)}{|r - r'|} \delta (\varepsilon_f - \varepsilon_i + \omega_n - \omega_0) \right|^2 \]

\(^9\) In the retarded case, the reciprocity theorem states that \( G(r, r', \omega) = G^T(r', r, \omega) \), a fact that we have used to recast exponential factors involving \( \omega/v > k \), so that it cannot contain wavevectors \( \omega/v > k \). This guarantees that the integral in Eq. (13) yields zero for an electron moving in vacuum, and that we are allowed to utilize the total rather than the induced field to obtain this equation.

C. Quantum approach

The quantum nature of both the electron probe and the excitations sustained by the targeted materials permeate many aspects of the electron-sample interaction. However, Ritchie and Howie (1988) showed that a quantum mechanical description of EELS yields the same results as a semiclassical formalism if all the inelastic signal is collected, and therefore readers that are more interested in applications and in a general overview of the subject can overlook this section, unless they feel motivated by Sec. V.B.2.

We work for simplicity in the non-retarded limit, in which the validity of Eq. (10) is demonstrated next under very common experimental conditions (Ritchie and Howie, 1988).

1. Quantum description of the target

The last century has witnessed important developments in the field of interaction of fast charges with solids, particularly in the community of electronic and atomic collisions (see, for instance, Palmer and Rous, 1992, Ziegler, 1999, Winter, 2002, and references therein), from which our theoretical understanding of electron microscopy has benefited considerably. In that context, ion and electron stopping was of great importance to understand dynamical screening in solids, with interesting developments such as the description of plasmon excitations using second quantization schemes in planar (Lucas and Sunjić, 1971) and spherical surfaces (Ashley and Ferrell, 1976). Moreover, a non-retarded, fully quantum-mechanical expression for the loss probability was derived using a self-energy formalism (Echenique et al., 1987, 1990). A quantum treatment of the target has been shown to be necessary for a correct assessment of delocalization in the excitation of core levels (Allen et al., 2003; Oxley and Allen, 1998). We here obtain a general expression for the loss probability, starting from the more widely used Fermi golden rule, and assess the conditions under which it agrees with the semi-classical formalism presented above, as previously discussed by Ritchie and Howie (1988).

During the interaction of a fast electron with a target, the latter can undergo transitions from its ground state \( |0 \rangle \) of energy \( \hbar \omega_0 \) to excited states \( |n \rangle \) of energy \( \hbar \omega_n \), while the incoming electron of energy \( \hbar \varepsilon_f \) and wave function \( \psi_f(r) \) acquires components \( \psi_f(r) \) of lower energy \( \hbar \varepsilon_f \). Since the interaction with very energetic electrons is generally small, the transition rate is well described within first-order perturbation theory (Fermi’s golden rule):

\[ \frac{d\Gamma_{\text{NR}}}{dt} = \frac{2\pi e^2}{\hbar} \sum_{f,n} \left| \int d^3rd^3r' \frac{\psi_f^*(r)\psi_i(r)}{|r - r'|} \delta (\varepsilon_f - \varepsilon_i + \omega_n - \omega_0) \right|^2 \]

where we have used the target-probe Coulomb interaction and \( \hat{\rho} \) is the target electron-density operator (see Fig. 7). We can now recast this expression into a more convenient form by relating the target matrix elements to the linear-response susceptibility (Pines and Nozières, 1966)

\[ \text{Im} \left\{ \chi(r, r', \omega) \right\} = -\frac{\pi}{\hbar} \sum_n \langle 0|\hat{\rho}(r)|n\rangle \langle n|\hat{\rho}(r')|0\rangle \delta (\omega_n - \omega_0 - \omega), \]

valid for \( \omega > 0 \), and this in turn to the screened interaction

\[ W_{\text{ind}}(r, r', \omega) = \int d^3r_1d^3r_2 \frac{\chi(r_1, r_2, \omega)}{|r - r_1||r' - r_2|}. \]

Then, the sum of Eq. (14) can be separated into specific values of the frequency transfer \( \omega = \varepsilon_i - \varepsilon_f \) as

\[ \frac{d\Gamma_{\text{NR}}}{dt} = \int_0^\infty d\omega \frac{d\Gamma_{\text{NR}}(\omega)}{dt}. \]
where
\[
\frac{dI^{\text{NR}}(\omega)}{dt} = \frac{2e^2}{\hbar} \sum_f \int d^3r d^3r' \psi_f(\mathbf{r}) \psi_i^*(\mathbf{r}) \psi_f^*(\mathbf{r}') \psi_i(\mathbf{r}') \\
\times \text{Im} \left\{ -W(\mathbf{r}, \mathbf{r}', \omega) \right\} \delta(\varepsilon_f - \varepsilon_i + \omega). \tag{15}
\]
This expression is general for incident electrons of well-defined energy, and we remind the reader that \( W \) contains all quantum-mechanical details of the sample response, although expressions obtained from dielectric theory such as those offered in Appendix B yield reliable results in most situations encountered in practice.

2. Quantum effects in the fast electrons

Several authors have analyzed the formation of images in electron microscopes from a quantum-mechanical viewpoint, considering the influence of instrumental parameters, as for example the beam aperture and the collection angle (Batson, 1985; Kohl, 1983; Muller and Silcox, 1995; Ritchie and Howie, 1988). Particular emphasis has been made on taming delocalization, which is relevant to devise ways of improving the spatial resolution of the inelastic signal (Allen et al., 2003; Oxley and Allen, 1998). We are not going to enter into details of the instruments here, but it is instructive to consider effects related to partial detection of the transmitted electrons.

We can regard each electron in a TEM as consisting of a coherent superposition of plane waves, yielding for instance a narrow focus close to the sample in STEM mode. A very accurate approximation can be adopted if we keep in mind that the angular aperture of the beam has remained in the range of a few milliradians since the early days of electron microscopy (Ruska, 1987). The components of the incident electron wavevector perpendicular to the beam direction \( z \) are thus \( p_{i\perp} \lesssim 10^{-2} p_i \), where \( p_i \) is the total wavevector. The parallel components become \( p_{i\parallel} = (p_i^2 - p_{i\perp}^2)^{1/2} \approx p_i \), where we are neglecting \( p_{i\perp}^2 / 2p_i \lesssim 10^{-4} p_i \) in front of \( p_i \). We conclude that the incident charge can be reliably described near the sample by the wave function
\[
\psi_i(\mathbf{r}) = \frac{1}{L^{1/2}} e^{i\mathbf{p}_i \cdot \mathbf{r}} \psi_{i\perp}(\mathbf{R}), \tag{16}
\]
where \( L \) is the quantization length of the trajectory and the \( \mathbf{R} \) dependence reflects the variation perpendicular to \( z \). This should be valid for typical sample thickness \( \Delta z \sim 50 \text{ nm} \), along which the lateral divergence of the beam can be quantified as \( \Delta R \lesssim 0.5 \text{ nm} \), a small value compared to typical dimensions of common photonic and plasmonic structures.

The transmitted electrons can be again described by plane waves of wavevector \( \mathbf{p}_f \). Then, the frequency transfer reduces to \( \omega \approx \mathbf{q} \cdot \mathbf{v} \) under the approximation of Eq. (3), where \( \mathbf{q} = \mathbf{p}_f - \mathbf{p}_i \) is the wavevector transfer and \( \mathbf{v} = (\hbar/m_e) p_i \hat{\mathbf{z}} \) is the incident electron velocity.

We are now prepared to recast Eq. (15) into a more practical formula. Using delta-function normalization for the final states, multiplying the transition rate by the interaction time \( L/v \), and making use of the above considerations, we find
\[
\Gamma^{\text{NR}}(\omega) = \int d^2\mathbf{p}_{f\perp} \frac{dI^{\text{NR}}(\omega)}{d\mathbf{p}_{f\perp}},
\]
where
\[
\frac{dI^{\text{NR}}(\omega)}{d\mathbf{p}_{f\perp}} = \frac{e^2}{4\pi^3\hbar v^2} \int d^3\mathbf{r} d^3\mathbf{r'} \psi_{i\perp}^*(\mathbf{R}) \psi_{f\perp}(\mathbf{R'}) \times e^{i\mathbf{p}_{f\perp} \cdot (\mathbf{R} - \mathbf{R'})} e^{i\omega(z' - z)/v} \text{Im}\left\{ -W(\mathbf{r}, \mathbf{r}', \omega) \right\}
\]
is the loss probability resolved in \( \mathbf{p}_{f\perp} \) (the lateral wavevector of the transmitted electron \( \perp \mathbf{v} \)), or equivalently, the outgoing direction. This formula indicates that STEMs can be used to retrieve the full nonlocal dependence of \( \text{Im}\left\{ -W(\mathbf{r}, \mathbf{r}', \omega) \right\} \), but we defer a detailed discussion of this point to Sec. V.B.2, in which we explore vicinage effects.

In practice, electron beams are polychromatic. However, a trivial extension of the above formalism leads to the intuitive result that the loss probability is then the average of Eq. (17) over the incident-beam spectrum, thus precluding interference between components of different primary energy.

Equation (17) leads to a powerful result, established by Ritchie and Howie (1988), regarding the validity of the classical dielectric formalism employed in Sec. II.B.1. The unrestricted integral over all possible values of \( \mathbf{p}_{f\perp} \) yields \( \delta(\mathbf{R} - \mathbf{R'}) \), so that the loss probability reduces to
\[
\Gamma^{\text{EELS}}_{\text{EELS}}(\omega) = \int d^2\mathbf{R} |\psi_{i\perp}(\mathbf{R})|^2 \Gamma^{\text{NR}}_{\text{EELS}}(\omega),
\]
where \( \Gamma^{\text{NR}}_{\text{EELS}}(\omega) \) is given by Eq. (10). In other words, the EELS probability is well described by Poisson’s equation if all the inelastic signal is collected (i.e., using a wide acceptance angle in the spectrometer), but it needs to be averaged over electron impact parameters weighed by the spot intensity \( |\psi_{i\perp}(\mathbf{R})|^2 \).

III. ELECTRON ENERGY-LOSS SPECTROSCOPY

Hillier and Baker (1944) were the first to propose and demonstrate EELS in TEMs, although earlier pioneering experiments reported energy losses of transmitted electrons in thin films (Leithäuser, 1904). This technique has become standard in the electron microscopy community and is capable of providing information on electronic band structures and plasmons in the low-energy loss region, as well as atomically-resolved chemical-identity encoded in core losses (Browning et al., 1993). During its prolific existence, valence EELS has contributed to fields as varied as biochemistry (for example, in the study of excitations sustained by nucleic acid bases, reported by
Crewe et al., 1971), interplanetary science (for instance, in the explanation of a 5 eV strong absorption feature in cosmic dust found by Bradley et al., 2005), and microelectronics (in particular, in the investigation of the resistivity of CMOS elements performed by Pokrant et al., 2006).

Since the early days of EELS, transmission electron microscopes have undergone a tremendous series of improvements that permit currently achieving \( \sim 0.1 \) eV energy resolution for a sub-nanometer-size electron beam. This opens up new vistas in the low-energy loss region, such as addressing the optical properties of photonic structures with unprecedented spatial detail.

The excitation spectrum of a typical material is illustrated in Fig. 4 for bulk Ag. The EELS intensity for a given energy loss \( \hbar \omega \) directly reflects the strength of specimen modes corresponding to that energy (Egerton, 1996). The loss spectrum has been traditionally separated into valence- and core-loss regions, with the division between them arbitrarily established at \( \sim 50 \) eV. With the customary use of thin samples, the main feature in an EELS spectrum is the zero-loss peak (ZLP) of unscattered electrons, in which unresolved, very-low-energy excitations (e.g., phonons) are buried. The intensity of valence features is over two orders of magnitude lower than the ZLP in the 20 nm Ag film considered in Fig. 4, while higher-energy core excitations are even less probable. This presents a typical scenario in which the inelastic signal is superimposed into the tail of the ZLP. Removal of the ZLP is thus important, and various sophisticated procedures have been devised that produce very reliable results (Lazar et al., 2006; Nelayah et al., 2007a).

Valence losses are generally more intense than core losses and allow collecting any given amount of inelastic signal with less electron irradiation, therefore producing minimum damage to the sample. However, the interpretation of low-energy-loss images is less direct because it involves the excitation of delocalized modes (Erni et al., 2008; Howie, 2003). In this respect, it is useful to rely on theoretical simulations, which are now advanced to the point of dealing with virtually any sample geometry in a predictive way.

The books of Raether (1980) and Egerton (1996) are classic references of EELS that extensively cover these subjects and provide detailed considerations on the observation and principles of TEMs. Several review papers present shorter summaries (Brydson, 2001) and collections of experimental spectra (Ahn, 2004). Comprehensive reports on valence EELS (Wang, 1996) and its theoretical analysis (Rivacoba et al., 2000) are also available. The present article intends to supplement this field by providing a more extended overview of low-energy losses, their characteristics and theoretical understanding, and several examples of novel application to nanophotonics, for which they might become particularly useful.

### A. Space, momentum, and energy resolution

Nowadays, some commercial TEMs incorporate the latest achievements in space and energy resolution. The latter has been tremendously improved over the last decade by the development of aberration correctors in the electron optics, particularly the spherical aberration (Uhlmann and Haider, 1998). This has reduced the size of the beam spots from the 1-2 nanometers commonly achieved without such correctors down to sub-Å dimensions (Krivamek et al., 1999), which allows for instance imaging closely spaced atoms in crystal samples using core losses (Batson et al., 2002; Nellist et al., 2004; Varela et al., 2004, 2007), and even identifying details of chemical bonding (Muller et al., 2008). However, the interpretation of atomically-resolved images is sometimes difficult and requires to rely on simulations (Oxley and Pennycook, 2008; Wang et al., 2008a). The search for improved spatial resolution through aberration-correction is actually a work in progress, for example through theoretical analysis for improved incoherent imaging (Intaraprasok et al., 2008) and through new sub-Angstrom beam characterization techniques (Dwyer et al., 2008).

The factors that limit the spatial resolution of a microscope are typically known as delocalization effects (Egerton, 2003, 2007; Erni and Browning, 2005). There are several of them, of very different nature, that can be grouped into three distinct categories: (1) instrumental or lens delocalization, which as noted above can be reduced below 1 Å; (2) Coulomb delocalization associated to the finite range of the field that accompanies a fast electron (see Fig. 6 and discussion in Sec. II.A); and (3) the extended nature of the excitations that are probed, varying from macroscopic distances in the case of low-energy SPPs and CR losses, to nanometers in particle plasmons, and less than 1 Å for core losses. For example, sub-atomic resolution is currently limited by the size of the probe rather than by the extension of the ionizing interaction that is employed to resolve core levels (Allen et al., 2003).

Additionally, spectral resolution has dramatically improved with the arrival of new spectrometers yielding \( \sim 50 \) meV accuracy (Brink et al., 2003), and electron monochromators (Su et al., 2003), which roughly consist in filtering out incident electrons outside a narrow energy window at the expense of reducing the beam current. An energy resolution of 0.1 eV has been achieved. Measuring excitations down to less than \( \hbar \omega = 0.5 \) eV is now possible thanks to the limited extension of the monochromatized ZLP (Lazar et al., 2006; Terauchi et al., 1999). This spectral resolution should be sufficient for studying most collective excitations supported by metallic systems, in which the intrinsic width produced by absorption is generally larger than a tenth of an electronvolt.

Deconvolution techniques, utilized to eliminate the ZLP and enhance spectral resolution, have progressed tremendously in recent years (Lazar et al., 2006; van Benthem et al., 2001). Furthermore, dynamical insta-
B. Bulk losses and determination of bulk dielectric functions

Although the volume of homogeneous materials offers little chances for performing microscopy, bulk losses are a splendid source of information on optical properties (Pallik, 1985; Raether, 1980) and they configure an area of little chances for performing microscopy, bulk losses are a function that limit the performance of electron microscopes, as recently reviewed by Egerton (2007). However, many factors are remaining uncorrected, and they constitute a correction of order \(q^2\) in virtue of Eq. (3).

The momentum dependence of the transversal permittivity can be safely neglected in this discussion, since it constitutes a correction of order \(v_F/c\) to the response, where \(v_F\) is the Fermi velocity (e.g., \(v_F = 1.39 \times 10^6\) cm s\(^{-1}\) in gold). Besides, the Lindhard formula (Lindhard, 1954) for the response of a free-electron gas (see explicit expression in Sec. A1) provides a fair description of the longitudinal dielectric function in good metals such as Al, although local-field corrections can be also relevant (Vast et al., 2002).

In this context, the response of gold has increasing technological importance, but detailed experimental analysis of the momentum-dependent optical constants is still missing. The local dielectric function determined by Johnson and Christy (1972) from ellipsometry measurements is routinely employed in plasmonic studies, and it

function \(\Im\{-1/\epsilon\}\) (Egerton, 1996; French et al., 1998; Pfüger and Fink, 1991; Shiles et al., 1980; Zhang et al., 2008). However, retardation corrections like CR losses (see Sec. IV.D) can be very damaging in the determination of electronic band gaps and they must be carefully removed from the measured spectra in some cases (Erni and Browning, 2008; Jouffrey et al., 2004; Stöger-Pollach et al., 2006; Stöger-Pollach and Schattschneider, 2007).
can be well described by the Drude-like formula of Eq. (A7) in the visible and NIR part of the spectrum (see Table II). In a recent report (García de Abajo, 2008), the author has proposed a nonlocal $\epsilon_{\text{lon}}$ for gold constructed as

$$
\epsilon_{\text{lon}}(q, \omega) = \epsilon^{\exp}(\omega) - \epsilon^{D}(\omega) + \epsilon^{M}(q, \omega),
$$

(21)

where $\epsilon^{D}$ and $\epsilon^{M}$ are the Drude and Mermin dielectric functions given by Eqs. (A1) and (A4), respectively, and corresponding to the values of $\omega_p$ and $\eta$ listed in Table II, whereas $\epsilon^{\exp}$ is the measured local permittivity. Further experimental effort is still required to contrast this model.

The spectrum in bulk metals is dominated by plasmon losses, first observed by Ruthermann (1948) and Lang (1948), and later identified as collective oscillations of the conduction-electron gas by Pines and Bohm (1952). These modes are signaled by the vanishing of $\epsilon_{\text{lon}}$, as deduced from the non-retarded (NR) limit of Eq. (20),

$$
\frac{d\Gamma_{\text{bulk}}(\omega)}{dq_{\perp}} = \frac{e^2 L}{\pi^2 \hbar^2 q^2} \text{Im} \left\{ \frac{-1}{\epsilon_{\text{lon}}(q, \omega)} \right\}.
$$

(22)

The dispersion relation of bulk plasmons has been extensively studied in the past using EELS (Batson et al., 1976; Chen et al., 1980, 1976; Raether, 1980; Watanabe, 1956) and is shown in Fig. 8(a) for Al, as measured by Batson and Silcox (1983) (symbols). When broad collection angles are considered, one can reliably approximate $\epsilon_{\text{lon}}(q, \omega)$ by a plasmon-pole analytical expression (Ritchie, 1957), reproduced in Eq. (A6). The plasmon dispersion relation derived from this formula provides a reasonable interpolation between the measured plasmon and the Bethe ridge $\omega = \hbar q^2/2m_e$, shown in Fig. 8(a) as a solid curve.

Spatially-resolved bulk-plasmon spectroscopy has been used to identify the presence of metals in a specimen [e.g., Na in either Na$_2$O or silicate glasses (Jiang et al., 2008)] and to demonstrate the effect of quantum confinement in carbon nanostructures by tracing local variations in the plasmon energy (Stolojan et al., 2006), which are related to changes in the conduction-electron density through Eq. (A2).

Finally, magnetic circular dichroism has been measured in the response of iron at x-ray energies (Schattschneider et al., 2006), suggesting that further work could eventually explore optical activity at visible and NIR frequencies via EELS, for instance in chiral (Rogacheva et al., 2006) and left-handed (Boltasseva and Shalaev, 2008) metamaterials.

C. Planar surfaces

Surfaces and interfaces can host trapped modes that modify the local band structure and the optical response with respect to bulk materials. For instance, we find electronic surface states confined by directional gaps in noble-metal interfaces (Smith, 1985), but also specific collective modes of conduction electrons (e.g., surface plasmons). As we show later on, these features can be characterized by EELS performed in STEMs with high degree of spatial resolution.

A particularly instructive situation is presented when the beam is directed parallel to a planar interface separating two different media (Fig. 9, left). This configuration has been extensively studied both experimentally (Lecante et al., 1977; Moreau et al., 1997; Powell, 1968; Walls and Howie, 1989) and theoretically (Echenique and Pendery, 1975; García-Molina et al., 1985a; Lucas and Šunjić, 1971). The analysis of this geometry in the non-retarded limit can be readily made from the knowledge of the screened interaction $W(\mathbf{r}, \mathbf{r'}, \omega)$, defined as the potential created at point $\mathbf{r}$ by a charge oscillating with...
frequency $\omega$ and placed at $\mathbf{r}'$. Expressions of $W$ for simple geometries, including that of Fig. 9, are given in Appendix B. A closed-form expression for the loss probability of the electron following the parallel trajectory of Fig. 9 (left) is then obtained from Eqs. (10) and Eqs. (B3). One finds

$$\Gamma^{\text{NR}}_{\text{planar}}(\omega) = \frac{2e^2L}{\pi\hbar^2} \left\{ \ln\left(\frac{q_0v}{\omega}\right) \text{Im}\left\{\frac{-1}{\epsilon_1}\right\} + K_0\left(\frac{2\omega b}{v}\right) \text{Im}\left\{-\frac{2}{\epsilon_1 + \epsilon_2}\right\} - \text{Im}\left\{-\frac{1}{\epsilon_1}\right\} \right\},$$

where $L$ is the length of the trajectory. The impact parameter $b$ and the dielectric functions $\epsilon_1$ and $\epsilon_2$ are defined in Fig. 9. The first term inside the curly brackets is the bulk loss probability of Eq. (19). The square brackets in Eq. (23) contain the effect of the interface, which dies off exponentially with $b$ as $K_0(\theta) \approx \exp(-\theta)\sqrt{\pi/2\theta}$ for large $\theta = 2\omega b/v$ (Abramowitz and Stegun, 1972). The first interface term describes the excitation of intrinsic boundary modes, signalled by the condition $\epsilon_1 + \epsilon_2 = 0$. It was first derived by Echenique and Pendry (1975), who obtained the expression

$$\Gamma^{\text{NR}}_{\text{planar}}(\omega) = \frac{4e^2L}{\pi\hbar^2} K_0\left(\frac{2\omega b}{v}\right) \text{Im}\left\{-\frac{1}{1 + \epsilon}\right\},$$

for an electron moving in vacuum near a medium of dielectric function $\epsilon$. The second interface term in Eq. (23) accounts for a reduction of bulk losses (i.e., the transfer of oscillator strength from volume to surface modes). This is the so-called begrenzungseffekt.

Equation (23) has been successfully applied to explain surface losses suffered by electrons passing near MgO cubes (Cowley, 1982a,b; Marks, 1982; Walls and Howie, 1989; Wang and Cowley, 1988) in the so-called aloof configuration (see below), and also to EELS in a Si/SiO$_2$ interface and in GaAs surfaces (Howie and Milne, 1985).

Inclusion of retardation effects is critical to account for CR losses and to correctly assess the weight of interface effects, as demonstrated by Moreau et al. (1997) for the Si/SiO$_2$ interface, and later by Couillard et al. (2007) for multilayer structures containing HfO. These studies have been recently extended by Yurtseyer et al. (2008) and Couillard et al. (2008), who have measured EELS in slabs and layered stacks formed by Si and SiO$_2$ using electron beams parallel to the interfaces and have demonstrated the important role of retardation corrections for a quantitative comparison with theory.

The retarded counterpart of Eq. (23) has been reported by several authors (García-Molina et al., 1985a; Otto, 1967). A particularly simple derivation consists in expanding the integrand of Eq. (2) into $p$ (TM) and $s$ (TE) electromagnetic plane-wave components, which are later reflected at the interface to act back on the electron (Forstmann et al., 1991). One finds

$$\Gamma_{\text{EELS}}(\omega) = \Gamma_{\text{bulk}}(\omega) + \Gamma_{\text{ref}}(\omega),$$

where $\Gamma_{\text{bulk}}$ is the bulk loss inside the medium 1 in which the electron is moving, given by Eq. (18), and

$$\Gamma_{\text{ref}}(\omega) = \frac{2e^2L}{\pi\hbar^2} \int_0^\infty dq_{y} \frac{q_y^2}{q_{y}} \left[ q_y^2 - 1 \right]$$

is the loss due to reflection of the electron field at the interface. Here, $q_{\parallel} = \sqrt{\omega^2/v^2 + q_y^2}$, $q_{\perp} = \sqrt{k^2 - q_{\parallel}^2}$, and

$$r_p = \frac{\epsilon_2 q_{\perp} - \epsilon_1 q_{\perp}}{\epsilon_2 q_{\perp} + \epsilon_1 q_{\perp}},$$

$$r_s = \frac{q_{\perp} - q_{\parallel}}{q_{\perp} + q_{\parallel}},$$

are Fresnel reflection coefficients for $p$ and $s$ waves, respectively. In the non-retarded limit, one has $r_s = 0$ and $r_p = (\epsilon_2 - \epsilon_1)/(\epsilon_2 + \epsilon_1)$, from which Eq. (23) is easily recovered. The advantage of this approach, based upon optical reflection coefficients, is that it can handle more complicated surfaces, like the periodically corrugated boundary of a confined photonic crystal (García de Abajo and Blanco, 2003; Pendry and Martín-Moreno, 1994). For large impact parameters, Eq. (25) provides a retardation correction to Eq. (24), consisting in substituting $2\omega b/v\gamma$ for $2\omega b/v$ in the argument of the $K_0$ Bessel function; the apparent impact parameter at very large velocity is contracted to $b/\gamma$.

1. Excitation of surface plasmons and surface-plasmon polaritons

Surface modes, signalled by the condition $\epsilon_1 + \epsilon_2 = 0$, are characteristic of the interface between a dielectric having $\text{Re}\{\epsilon_1\} > 0$ and a metal satisfying $\text{Re}\{\epsilon_2\} < 0$.}

![Fig. 9 (Color online) Planar interface separating two media of dielectric functions $\epsilon_1$ and $\epsilon_2$, respectively. An electron is shown moving parallel to the boundary inside medium 1 (left), with impact parameter $b$ relative to the interface. Another electron is represented crossing the interface at normal incidence (right).](image-url)
These excitations were first identified and understood by Ritchie (1957) to explain anomalous valence losses that were previously observed at positions differing from bulk plasmons in the spectra of fast electrons transmitted through thin foils (see, for instance, feature D in Fig. 2 of Watanabe, 1956). Ritchie’s description in terms of a classical hydrodynamic plasma to represent the conduction electron band led Stern and Ferrell (1960) to baptize these excitations as surface plasmons, which were later confirmed in electron energy-loss experiments (Powell and Swan, 1959) (see Fig. 10). Bulk and surface modes were observed as $h\omega_p \approx 15\,\text{eV}$ and $h\omega_s \approx 10.6\,\text{eV}$ energy losses in electrons reflected from an AI surface. [Incidentally, the spectrum of Fig. 10 contains multiple plasmon losses (i.e., two or more plasmons being excited by the same electron). The coherent aspects of double plasmon excitation have deserved some attention in the past (Schattschneider et al., 1987).]

Aluminum is a prototypical example of nearly-free electron gas for which the dielectric function is well described by the Drude formula of Eq. (A1). The SP of the bare surface, satisfying the condition $\epsilon + 1 = 0$, has then a frequency $\omega_s = \omega_p/\sqrt{2}$, in excellent agreement with experimental observations.

Surface plasmons were also observed for other materials in subsequent reflection experiments performed on melted metals (Powell, 1968), and even the evolution of these modes through a solid-liquid phase transition was reported (Powell, 1965). Other pioneering electron transmission experiments managed to demonstrate the observation of SPs as a characterization technique (Raether, 1967) and to determine the dependence of SP energies on the thickness of thin films (Boersch et al., 1966a). Evidence of surface and bulk plasmon excitation has been also observed in the spectra of photoelectrons and Auger electrons emitted from metals (Osterwalder et al., 1990; Simonsen et al., 1997; van Attekum and Trooster, 1978).

Similar to the bulk plasmons of Fig. 8(a), SPs are affected by nonlocal effects and exhibit significant dispersion, which was already reported in early EELS studies (Krane and Raether, 1976). This has stimulated a rich literature intended to achieve a reasonable first-principles description of the dynamical response of crystal surfaces (see Pitarke et al., 2007, and references therein). Nonlocal effects involve comparatively short-range interactions extending a distance $\sim 1/q$ for typical wavevector transfers $q$ in the $\AA^{-1}$ domain (see Fig. 8(a)). However, we are mainly concerned here with longer-range optical excitations (e.g., SPPs), in which space dispersion can be generally overlooked, except when sharp metallic structures or narrow gaps between metals are involved (García de Abajo, 2008), as shown in Sec. V.E for electrons passing near metal-sphere dimers. Nonlocal effects can be also significant at small separations $\lesssim 1\,\text{nm}$ between the passing electron and metal surfaces (Zabala and Echenique, 1990).

An interesting scenario is presented by the aloof configuration, in which the electron trajectory does not even intersect the sample. This leads to distant interaction, capable of providing surface-specific information, free of close-encounter events between the fast probe and the target atoms. The exponential decay of surface losses with $2\omega_b/\nu^2$ indicates that low-energy transfers are favored in the aloof configuration. This has the additional benefit of minimizing sample damage.

The aloof configuration was pioneered by Lecante et al. (1977), who forced the electrons to describe parabolic trajectories, deflected from the metal surface by a bias potential. Subsequent studies managed to aim a STEM aloof beam parallel to the planar surfaces of MgO cubes (Cowley, 1982a,b; Marks, 1982). Loss spectra were later recorded after transmission through perforated metallic channels (Warmack et al., 1984). The aloof configuration has been recently revisited and compared with near-field optical microscopy (Cohen et al., 2003, 1998, 1999; Echenique et al., 1999; Itskovsky et al., 2008).

Retardation adds another source of plasmon dispersion. Since SP modes are resonances in the surface response, they must involve a divergence in the reflectivity for incident evanescent waves with the right values of $(q, \omega)$. More precisely, we obtain the plasmon dispersion relation from the vanishing of the denominator of $r_p$ [Eq. (26)], leading to

$$q_{\parallel}^{\text{SP}} = \sqrt{\frac{\epsilon_1 \epsilon_2}{\epsilon_1 + \epsilon_2}}$$

under the condition that $\text{Re}\{\epsilon_1\}$ and $\text{Re}\{\epsilon_2\}$ have opposite signs. The dispersion relation of Eq. (27) is represented in Fig. 11 (left) for a silver-vacuum interface. When the retarded nature of SPs becomes important, they are usually referred to as SPPs, because they acquire a polaritonic character reflecting a mixture of collective electron motion and propagating electromagnetic fields. This tends to happen at low wavevectors (i.e., at long wavelengths according to Fig. 11), for which SPPs increasingly approach the light line ($\omega = q_0 \epsilon$). It is important to stress that this polaritonic regime occurs for momentum transfers that are well below the region exhibiting nonlocal effects (cf. horizontal scales in Figs. 8(a)}
and 11]. This is shown in Fig. 8(b), where the hybridization of transversal bulk plasmons with light takes place at values of \( q \) that are two orders of magnitude smaller than the Fermi wavevector \( k_F \). More details on SPPs, including a comprehensive literature survey, can be found in the book of Raether (1988).

The electric and magnetic fields associated to a SPP plane wave moving along \( x \) have similar strength in Gaussian units. For instance, in the vacuum region outside a semi-infinite metal of permittivity \( \varepsilon \), for \( q \ll \varepsilon \), we have

\[
E^{\text{SP}}(r, \omega) = [(\hat{x}q^{\text{SP}}_x - \hat{z}q^{\text{SP}}_z)/k] \exp[i(\hat{q}^{\text{SP}}_x x + q^{\text{SP}}_z z)],
\]

\[
H^{\text{SP}}(r, \omega) = \hat{y} \exp[i(\hat{q}^{\text{SP}}_x x + q^{\text{SP}}_z z)],
\]

where \( q^{\text{SP}}_z = \sqrt{k^2 - (\hat{q}^{\text{SP}}_x)^2} = -k/\sqrt{\varepsilon} + 1 \). In particular, the electric field satisfies \( \nabla \cdot E^{\text{SP}} = 0 \), and therefore it is transversal.\(^{10}\) Close to the plasmon cutoff, the SPP momentum becomes increasingly large. This marks the transition towards the non-retarded regime, in which magnetic and electric fields are decoupled, so that we can approximate \( \nabla \times E^{\text{SP}} = 0 \) and define a potential \( \phi = \exp[i(\hat{x}x - \hat{z}z)q^{\text{SP}}_z] \) to write \( E^{\text{SP}} = -\nabla \phi \). The electric field appears to be longitudinal, and the plasmons are then known as SPs rather than SPPs, signalled by the condition \( \varepsilon = -1 \).

An aloof electron like that of Fig. 9 (left) couples to SPPs of momentum component along the velocity vector given by \( h\omega/v \), according to Eq. (3). Consequently, the electron only excites parallel momenta above that value. This condition is represented in Fig. 11 (left) for two different electron energies. The corresponding electric fields are shown in the same figure (right) within a plane that contains the velocity vector and the surface normal. The electron is passing at a distance of 10 nm above the surface. We can see several characteristic features in these plots: (1) the field shows a wake pattern characterized by oscillations of wavelength \( \approx 2\pi/(q^\parallel) \), with \( q^\parallel = \omega/v \) determined by the intersection points of Fig. 11 (left); (2) the electron that moves faster excites plasmons of lower energy, thus giving rise to oscillations of longer spatial period; (3) at variance with the continuity of the parallel electric field, the normal component changes sign across the surface in order to preserve the continuity of the normal displacement, because the metal permittivity is negative; (4) the normal field component takes large values compared to the parallel one, and this effect is more pronounced for faster electrons, which involve lower \( \omega \)'s.

In a different direction, the early 1990s witnessed remarkable experiments of energy losses in coincidence with SEE, initially collected in amorphous carbon samples (Müllejans and Bleloch, 1992; Pijper and Kruit, 1991). Angle-resolved measurements were also carried out in this context to conclude that the SEE yield is larger in more localized excitations, involving larger-momentum transfers (Drucker and Scheinfein, 1993). Comparative studies for amorphous carbon and silicon suggested that bulk plasmons do not play a central role in SEE (Drucker et al., 1993; Scheinfein et al., 1993), possibly due to the noted begrenzungseffekt, which limits the strength of volume plasmons in favor of SPs within the region accessible to the escape depth of SEs. Finally, coincidence experiments were conducted using the aloof beam geometry in diamond and MgO to compare EELS and SEE rates and to provide direct measurements of the probability that a surface excitation gives rise to a SE, which was found to be below \( \sim 5\% \) for low-energy losses (Müllejans et al., 1993).

2. Guided modes in thin films

The SPPs on both sides of a thin film interact to produce two hybridized modes. One of them can travel longer distances along the film due to exclusion of the electric field from the metal (Sarid, 1981). This mode is currently being applied to propagate electromagnetic signals in plasmonic devices (Barnes et al., 2003; Berini et al., 2007). The dispersion relation of coupled SPPs in thin films was obtained from the EELS signal of transmitted electrons over 30 years ago by a series of outstanding experiments conducted by Silcox and co-workers in Al (Pettit et al., 1975; Vincent and Silcox, 1973) and Si films (Chen et al., 1975). An example of these studies is given in Fig. 12 for a partially oxidized thin Al film. Both plasmon branches follow the light line at low energies, and they converge to the non-retarded limit of the Al/Al\(_2\)O\(_3\) interface for large momentum transfer. This limit corresponds to \( E_{\text{Al}} + 4 = 1 \), or equivalently, \( h\omega_s \approx 6.7 \) eV when Eq. (A1) is used to model Al, since the permittivity of Al\(_2\)O\(_3\) in this energy window is \( \approx 4 \). Large \( q^\parallel \)'s involve fast plasmon oscillations along the film, so that alternating induced charges weakens the interaction between both film sides and we recover the limit of the single interface separating two semi-infinite media. For small \( q^\parallel \), the interaction between plasmons in both sides of the film gives rise to two SPP branches.

Guided modes in graphite (Chen and Silcox, 1975a) and aluminum-oxide (Chen and Silcox, 1975b) films were also characterized using the same technique. In this case, the film behaved as a dielectric, and the electron coupled to optical Fabry-Perot resonances set up by successive total internal reflections at the film boundaries. The experiments were in excellent agreement with theory (Chen and Silcox, 1979).

In a more recent development, a complex multilayer structure containing HfO, which is of interest in microelectronics, has been characterized by Couillard et al.

\(^{10}\) Although the main \( E^{\text{SP}} \)-component of long-wavelength SPPs is along the surface normal, there is also a finite electric field along the propagation direction, but the normal components of both the wavevector and \( E^{\text{SP}} \) have a \( \pi/2 \) phase difference relative to the parallel components, leading to \( \nabla \cdot E^{\text{SP}} = 0 \).
FIG. 11 (Color online) Surface plasmon dispersion relation in Ag (left) and excitation of SPPs by swift electrons (right). The SPPs saturate at Ritchie’s non-retarded frequency for large parallel wavevector and follow the light line $\omega = q_v c$ in the low-energy limit. An electron moving with velocity $v$ parallel to the surface preferentially excites plasmons of frequency and wavevector related by $\omega \lesssim q_v v$. This condition is indicated by solid dots in the dispersion relation for the two electron energies considered on the right. Faster electrons generate plasmons of lower $q_v$ and longer wavelength. The impact parameter is 10 nm in the field plots. The dielectric function of Ag is taken from Palik (1985).

FIG. 12 (Color online) Experimental determination of the SPP dispersion relation in a thin film (adapted from Pettit et al., 1975). Symbols: measured data obtained from energy and angle distributions of fast electrons traversing a partially oxidized Al thin film. Continuous curves: SPP branches calculated from optical data.

The agreement with theory is excellent after retardation effects are incorporated to correctly account for interface plasmons and Cherenkov guided modes. A similar study has been recently presented by Yurtsever et al. (2008) in slabs and stacks of Si and SiO$_2$.

Additionally, a radiative plasmon branch inside the light cone, giving rise to losses near $\omega_p$, was predicted by Ferrell (1958) for thin metal films, and the resulting light emission was detected later by Steinmann (1960) in Ag. We discuss this issue in more detail in Sec. IV.F.1.

D. Curved geometries

The ability of TEMs to probe local response properties has been extensively exploited to study non-planar sample geometries. Particular attention has been paid to spherical and cylindrical shapes, carbon nanotubes, composites, and other more complicated structures. The intense valence losses can provide spectral images, the interpretation of which is not as direct as when core losses are employed instead. Therefore, we need to strongly rely on theory, which is actually in a very advanced state, capable of dealing with virtually any system. For example, small particles like fullerenes or carbon nanotubes are reliably described by atomistic models, although first-principles theory is also possible (Marinopoulos et al., 2003). For larger structures (e.g., nanoparticles) we have a suite of methods based upon classical dielectric theory, which gives excellent agreement with experiment in metallic objects of dimensions above $\sim 10$ nm. In this section, we explore some of these structures possessing spherical or cylindrical geometry.

1. Cylinders

Early work on cylindrical nanocavities relied on an electron beam to drill holes in AlF$_3$ (Macanhay et al., 1989), which were investigated by EELS (Scheinfein et al., 1985). The synthesis of such types of holes has considerably improved over the last few years in two different directions. (1) Bottom-up approach: chemical methods have been developed that are capable of spontaneously forming arrays of self-organized nanoholes in materials like alumina (Masuda and Fukuda, 1995); arrays of cylinders are also formed in eutectics (Pawlak et al., 2008). (2) Top-down approach: advanced electron-beam and focused-ion-beam (FIB) lithographies have been extensively employed to produce hole arrays for nanophotonics studies (Genet and Ebbesen, 2007). These and similar geometries have been investigated through TEM spectroscopic analysis (Degiron et al., 2004; García de...
one obtains the loss probability

cylinder, respectively, and
scripts 1 and 2 refer to the media outside and inside the
where the notation of Fig. 13 has been adopted, the sub-
ents of wavevector \( q \) is appropriate for the variation
along the direction of translational invariance, chosen along \( \hat{z} \). This decomposition is employed in Sec. B.2 to
write the non-retarded screened interaction, from which the cylinder modes are derived using the condition \( \Delta_m = 0 \) [see Eq. (B4)]. A relatively simple expression can be
obtained for these modes including retardation (Ashley and Emerson, 1974):

\[
\begin{align*}
x_1^2 x_2^2 \left[ \epsilon_2 x_1 I'_m(x_2)/I_m(x_2) - \epsilon_1 x_2 K'_m(x_1)/K_m(x_1) \right] \\
\times \left[ x_1 I'_m(x_2)/I_m(x_2) - x_2 K'_m(x_1)/K_m(x_1) \right]
\end{align*}
\]

\[
= m^2 (\epsilon_1 - \epsilon_2)^2 \left( q_z a^2 \right)^2,
\]

(28)

where the notation of Fig. 13 has been adopted, the sub-
scripts 1 and 2 refer to the media outside and inside the
cylinder, respectively, and \( x_j = (\omega/v) \sqrt{1 - \epsilon_j v^2/c^2} \).

The interaction of the electron with a cylinder has been
analytically described using both non-retarded (Tu et al., 2006; Zabala et al., 1989) and fully-retarded (Walsh, 1991; Zabala et al., 1989) dielectric theory. One can easily derive the
loss probability using the screened interaction non-retarded limit [see Eq. (B4)]. For electrons moving
parallel to a cylinder, one just needs to insert \( W \) inside
Eq. (10). The momentum transfer along the cylinder axis, is fully determined by the condition \( q_z = \omega/v \), and one obtains the loss probability

\[
\Gamma_{\text{ref,cyl}}(\omega) = \frac{e^2 L}{\pi \hbar v^2} \sum_{m=-\infty}^{\infty} \text{Im} \left\{ -W_m(b, b, q_z, \omega) \right\},
\]

where \( b \) is the distance from the electron beam to the
cylinder axis, and different \( m \)'s contribute separately. Re-
tardation effects are however tedious to deal with analytically (Walsh, 1991; Zabala et al., 1989), as compared to the planar interface. For instance, a beam moving along
the center of the cylinder, which only couples to \( m = 0 \)
 modes due to the symmetry of the electron-cylinder com-
bined system, experiences a loss probability

\[
\Gamma_{\text{ref,cyl}}(\omega) = \frac{e^2}{2\hbar v^2} \text{Im} \left\{ \frac{v^2 - 1}{v^2} \right\}
\]

\[
\times \frac{\epsilon_2 x_1 I_0(x_1)I_1(x_2) - \epsilon_1 x_2 I_1(x_1)I_0(x_2)}{\epsilon_2 x_1 K_0(x_1)I_1(x_2) + \epsilon_1 x_2 K_1(x_1)I_0(x_2)}
\}

where we can immediately see the presence of the \( m = 0 \) mode predicted by Eq. (28) when we
arrange the denominator of the last fraction as
\( K_0(x_1)I_0(x_2)(\epsilon_2 I_0(x_2)/I_0(x_2) - \epsilon_1 x_2 K'_0(x_1)/K_0(x_1)) \).

2. Spheres

The interaction of electrons with spheres has attracted
c onsiderable attention since the publication of pioneering
experimental loss spectra in alkali-halide (Creuzburg, 1966) and Al (Fujimoto et al., 1967) particles, followed
by their subsequent theoretical interpretation (Crowell and Ritchie, 1968; Fujimoto and Komaki, 1968). Further
EELS measurements corroborated these results and fo-
cused on the role of plasmon dispersion in small spheres
(Batson, 1980; Ouyang et al., 1992) and in the effect of
the interaction between neighboring particles (Batson, 1982b, 1985; Ugarte et al., 1992). More recently, UV
surface exciton polaritons have been observed in gold
nanospheres (Chu et al., 2008) and whispering gallery
modes measured in silica beads (Hyun et al., 2008) using
EELS.

A small metallic sphere embedded in a dielectric host
exhibits plasmon modes at frequencies dictated by the
condition (Kittle, 1987)

\[
(l + 1)\epsilon_1 + l \epsilon_2 = 0,
\]

(29)

where \( l \) labels the multipolar order of the excitation, and
\( \epsilon_1 \) and \( \epsilon_2 \) are the permittivities of the host and the metal,
respectively (see Fig. 13). In particular, for an Al sphere
in which \( \epsilon_2 \) is described by the Drude dielectric function of
Eq. (A1), the frequency of the non-retarded plasmon
oscillations becomes

\[
\omega = \omega_p \sqrt{\frac{l}{(l + 1)\epsilon_1 + l}}.
\]

(30)

For example, the dipole \( (l = 1) \) of an Al sphere in vac-
uum has energy \( h\omega_p/\sqrt{3} \approx 8.7 \) eV. For very large \( l \), the
surface-charge oscillations are so dense that they cannot
resolve the particle curvature, and therefore the mode
frequency evolves to the plasmon of the planar interface,
\( \omega_s = \omega/\sqrt{\epsilon_1 + 1} \).

The remarkable works of Fujimoto and Komaki (1968)
and Crowell and Ritchie (1968) established the basis

FIG. 13 (Color online) Notation used for homogeneous spheres and cylinders or radius \( a \), with an electron passing at a distance \( b \) from the center.
for the theoretical understanding of fast-electron interaction with spherical particles. These authors described a metallic sphere using the hydrodynamical model of Appendix A.2 and derived analytical expressions for the loss probability and photon emission rate of a broad beam neglecting retardation. Their results showed spectral features at frequencies roughly described by Eq. (30).

This theoretical development was followed by studies of voids in metals (Ashley and Ferrell, 1976), and later by the derivation of an analytical formula for the loss probability of a STEM electron passing outside a sphere (Ferrell and Echenique, 1985; Ferrell et al., 1987). Namely,

$$\Gamma_{\text{sph}}^{\text{NR}}(\omega) = \frac{4e^2}{\pi\hbar v^2} \sum_{l=1}^{\infty} \sum_{m=-l}^{l} \frac{(\omega/v)^{2l}}{(l+m)!(l-m)!} K_m^2 \left( \frac{\omega b}{v} \right) \text{Im} \{ \alpha_l^{\text{NR}}(\omega) \}, \quad kb \ll 1, \quad v/c \ll 1,$$

where

$$\alpha_l^{\text{NR}}(\omega) = a^{2l+1} \frac{\ell e(\omega) - l}{\ell e(\omega) + l + 1}$$

(32)
is the non-retarded multipolar polarizability of a sphere of radius $a$ in vacuum, and $b$ is the distance from the trajectory to the sphere center, as illustrated in Fig. 13. Equation (31) can be derived upon insertion of the non-retarded screened interaction near a sphere [Eq. (B6)] into Eq. (10). This formula successfully explained previous measurements for aloof electrons (Batson, 1982a) and was soon applied to explain EELS of spheres made of tin (Achêche et al., 1986), silicon (Ugarte et al., 1992), silver (Ouyang et al., 1992), and aluminum (Stöckli et al., 1997), as well as corrugated metal surfaces (Mamola et al., 1987), and diamond particles (Fehlhaber and Bur sill, 1999, 2000; Peng et al., 2001). The effect of coatings, which can be important due to oxidation (Ferrell et al., 1987; Ugarte et al., 1992), is described by Eq. (31) if we use the polarizability of a coated sphere instead of Eq. (32).

Optical modes in nanoparticles (e.g., plasmons in metallic spheres) are well-known to be redshifted due to retardation when the particle size is not negligible in front of the wavelength (see Sec. IV.F.2): the interaction between distant regions of a particle is mediated by electromagnetic signals that propagate at the speed of light, but the signal simply takes longer to travel over larger distances as the particle dimensions increase, so that the period of the oscillation modes decreases, thus producing the noted redshift. This effect was first described by Mie (1908) in his analysis of light scattering by metallic spheres. Similar phenomena occur in particles made of polaritonic materials (Fuchs and Kliewer, 1968).

The extension of Eq. (31) to incorporate retardation also leads to an analytical formula (García de Abajo, 1999c; García de Abajo and Howie, 1998):

$$\Gamma_{\text{sph}}(\omega) = \frac{e^2}{c\hbar \omega} \sum_{l=1}^{\infty} \sum_{m=-l}^{l} K_m^2 \left( \frac{\omega b}{v} \right) \left[ C_m^M \text{Im} \{ t_m^M \} + C_m^E \text{Im} \{ t_m^E \} \right],$$

(33)

where the Lorentz factor $\gamma = 1/\sqrt{1-v^2/c^2}$ accounts for the contraction of the impact parameter $b$ as seen by the moving electron. Here, $C_m^M$ and $C_m^E$ are coupling coefficients that depend exclusively on the $v/c$ ratio, and $t_m^M$ and $t_m^E$ are magnetic and electric Mie scattering coefficients independent of beam energy and impact parameter. Analytical expressions for these coefficients are given in Appendix C for homogeneous spheres. Equation (33) results from a multipolar expansion of the electric field induced by the electron [Eq. (4)] after each multipole is scattered by the sphere to produce an outgoing wave acting back on the electron and giving rise to energy loss [see Eq. (8)].

It should be emphasized that Eq. (33) can be applied to any spherical object for external electron trajectories, provided $t_m^M$ and $t_m^E$ are known for that object. In particular, this has been recently done for left-handed spheres (Xu and Zhang, 2008) and spherical metamaterial cloaks (Xu et al., 2008).

Equation (33) predicts sizable redshifts even for relatively small Al spheres of radius $a = 10$ nm, as shown in Fig. 14(a) by comparison to spheres with $a = 5$ nm (solid curves). Spectral features associated to dipolar ($l = 1$) and quadrupolar ($l = 2$) excitations are clearly discernable in the spectra. Higher-order multipoles produce a shoulder piling up near $\omega_a$ because their separation is smaller than the absorption width. Besides, the dipole feature is broader in the bigger sphere, which is again a retardation effect, since the coupling of this mode to emitted light increases with the particle dipole strength (i.e., with particle size), therefore limiting the lifetime of the dipolar plasmon due to radiative decay (Myroshnychenko et al., 2008b). This point is further discussed in Sec. IV.F.2.

For sufficiently small particles and aloof trajectories, we can just retain dipole terms ($l = 1$), in which case Eq. (33) reduces to

$$\Gamma_{\text{EELS,dip}}(\omega) = \frac{1}{\hbar \pi} \left( \frac{2\omega}{v^2 \gamma} \right)^2 f \left( \frac{\omega b}{v \gamma} \right) \text{Im} \{ \alpha \},$$

(34)

where

$$f(\zeta) = K_2^2(\zeta) + K_0^2(\zeta)/\gamma^2.$$  

(35)

Actually, this approximation works quite well even for relatively big spheres, as shown by the dotted curves of

11 Loss and CL emission probabilities are plotted per unit of lost-energy range (i.e., in eV$^{-1}$), so that the areas under the spectral curves of Fig. 14(a) yield dimensionless probabilities per electron.
FIG. 14 (Color online) Plasmon losses and spatial resolution in Al spheres. (a) The loss probability is calculated for two different particle sizes, exhibiting several multipole plasmons that redshift as the particle radius increases. Solid curves: result of Eq. (33) using local response. Dashed curves: same equation with nonlocal scattering matrices (see Appendix C). Dotted curves: prediction of Eq. (34). The Al is described by the dielectric function of Eq. (A1) with $\bar{\hbar}\omega = 15$ eV and $\bar{\hbar}\eta = 0.5$ eV. (b) Impact-parameter dependence of the dipolar mode maximum. (c) Apparent radius of the dipole excitation in an Al sphere, modeled within local response, and defined as the impact parameter $b_{1/2}$ for which the EELS probability of a 200 keV electron is half the maximum as a function of $b$. The dipole-energy dependence on particle size (right scale) is taken into account. The $v\gamma/2\omega$ distance of exponential decay in $\Gamma_{EELS,dip}$ [Eq. (34)] is shown for comparison.

Fig. 14(a), obtained from Eq. (34) with the electric polarizability

$$\alpha = 3t_{E}^{2}/2k^{3} \quad (36)$$

[see Eq. (C2); $t_{E}^{2}$ is a Mie scattering coefficient]. The dipole mode is given correctly, although higher multipoles are obviously missed. In contrast to the particle-size-independent dipole-plasmon energy predicted by Eq. (32), we find that Eq. (36) incorporates retardation effects leading to a sizeable dipole-plasmon redshift with increasing particle diameter.

Spheres offer an excellent playground to discuss Coulomb delocalization effects (see Sec. III.A). As anticipated by Eq. (4) for the external electron field, Eq. (33) predicts an exponential decay of the EELS signal with $b$, and an attenuation distance given by $v\gamma/2\omega$. This is true when the argument of the Bessel functions is large compared to unity, a condition that is not satisfied near small spheres, in which those functions have a steeper dependence on $b$ (see Sec. II.A). This is actually observed in the $b$ dependence of the loss probability shown in Fig. 14(b) (solid curves) for the dipole peaks of the spheres considered in Fig. 14(a). The probability falls down by a factor of two at a distance of $\sim 2$ nm away from the surface of the smaller sphere, so that its apparent radius $b_{1/2} \sim 7$ nm is significantly smaller than $v\gamma/2\omega \sim 11$ nm. The sphere size dependence of $b_{1/2}$ [Fig. 14(c)] fully corroborates these arguments.

Small metallic particles exhibit further peak shifts originating in quantum confinement and spatial dispersion, the effect of which is to break down the hypothesis of local response. Nonlocal effects in the performance of bounded metals can be easily included using the specular-reflection-model approximation (García de Abajo, 2008; Ritchie and Marusak, 1966; Wagner, 1966). Then, Eq. (31) is still valid if we replace $\epsilon$ by (Rojas et al., 1988)

$$\epsilon_{l}(q,\omega) = \frac{\pi}{2a(2l+1)} \left[ \int_{0}^{\infty} dq \frac{\epsilon(q,\omega)}{\epsilon(q,\omega)} j_{l}^{2}(qa) \right]^{-1} \quad (37)$$

in the polarizability of Eq. (32), where $\epsilon(q,\omega)$ is the nonlocal bulk response discussed in Secs. III.B and A.1, and $j_{l}$ is a spherical Bessel function. This level of description is similar to the work of Fujimoto and Komaki (1968). Under the conditions of Fig. 14(a), one needs however to simultaneously include nonlocal and retardation effects, for which the above procedure is not enough.

The hydrodynamic model described in Sec. A.2 incorporates both spatial-dispersion and retardation effects. When applied to spherical particles, it still yields analytical expressions for the resulting nonlocal Mie scattering coefficients, which we introduce in Appendix C. Using these coefficients in combination with Eq. (33), we predict blue shifts in the plasmons of the Al particles of Fig. 14(a) (dashed curves), even for relatively large spheres. This type of dispersion operates on a length scale comparable to the wavelength of conduction electrons at the Fermi level, which is of the order of 1 nm for typical
metals. This is consistent with the large nonlocal effect observed for the quadrupole as compared to the dipole in the \( a = 5 \, \text{nm} \) sphere, since the \( l = 2 \) mode involves more sign changes in the induced surface charge that increase the relative importance of short distances. It should be noted that spatial dispersion has been found to be partially compensated by the smooth electron density profile at the particle surface (Ruppin, 1978).

In this regard, nonlocality is important at distances \( \sim v_F/\omega \) (Larkin and Stockman, 2005; Larkin et al., 2004), as derived from the minimum momentum transfer in the e-h pair region for a given excitation frequency \( \omega \) (see Fig. 8). This is the so-called Landau damping, which has been phenomenologically accounted for through a size-dependent local dielectric function, with the dephasing rate increased by \( \sim a/v_F \) (Batson, 1980; Hövel et al., 1993; Kreibig and Vollmer, 1995). Nonlocal effects in noble metal particles are noticeable up to a size \( 10-20 \, \text{nm} \) at optical frequencies, but it can be important in other materials for larger sizes \( \sim 100 \, \text{nm} \) (e.g., silver halide nanocrystals, studied with EELS by Oleshko, 2008).

There are only scarce data on plasmon dispersion in small spheres (see, for example, Ouyang et al., 1992), and particularly on the spatial distribution of these effects, so there is a clear need for further EELS experiments that can shed more light into this subject, which is becoming increasingly important in many nanophotonics developments that make use of metallic nanoparticles in the sub-20 nm range. We hope that these results, together with the dimer calculations of Sec. V.E, contribute to trigger future research in this direction.

3. Coupled nanoparticles

Coupling between plasmon modes in neighboring nanoparticles has attracted considerable interest in the electron microscopy community. Early energy-loss experiments focused on the interaction between neighboring spheres (Batson, 1982b, 1985), leading to hybridization of individual particle plasmons, the energies of which were successfully explained through analytical derivations. Further theoretical work addressed these modes in more detail (Ruppin, 1982; Schmeits and Dambly, 1991), and eventually the coupling of the electron to two neighboring spheres was derived in the non-retarded approximation (Zabala et al., 1997).

The interaction between more than two non-aligned particles is difficult to explore by analytical techniques. A particularly suitable method that yields highly convergent results, including retardation, consists in expanding the electromagnetic field in terms of multipoles around the particles, and then allowing these multipoles to propagate among particles, thus generating a self-consistent field (García de Abajo, 1999b). More details on this method are given in Sec. III.E.4.

Cylinder pairs have been theoretically addressed as well in the non-retarded limit by Schmeits (1989). This study was later extended to include retardation and to cope with more than two cylinders (García de Abajo and Howie, 2002; Manjavacas and García de Abajo, 2008).

A great deal of work has been devoted in recent years to the interaction between metallic nanoparticles in the context of nanophotonics, due in part to the strong enhancement of the light intensity in the inter-particle region at close proximity (Xu et al., 1999). This has direct application to biosensing using surface-enhanced Raman scattering (Talley et al., 2005). Furthermore, the plasmon modes of neighboring particles can be tuned to achieve on-demand frequencies by varying their separation, as shown in optical measurements performed on lithographically printed metal dots (Atay et al., 2004). However, there are still many issues that remain unclear in the interaction between particles, specially when the gap distance is of the order of a few nanometers. Among them, the role of Landau damping and other dispersion effects. Energy loss spectroscopy turns out again to be ideally suited for exploring these situations, in which nanometer spatial resolution is needed to resolve optical excitations. For example, the singular transition in the response of touching and non-touching particle dimers (Romero et al., 2006) has already prompted EELS measurements (Chu et al., 2009; N’Gom et al., 2008), although a comprehensive experimental study of this transition is still pending.

E. More complex shapes

The real world is generally more complex than planar, cylindrical, or spherical surfaces. There are numerous experimental studies of EELS involving more complicated geometries, like Si tips (Reed et al., 1999), metallic particles supported by dielectric (Wang and Cowley, 1987a,d) or by metal (Wang and Cowley, 1987b,d) substrates, MgO wedges (Aizpurua et al., 1999; García de Abajo and Aizpurua, 1997), MoS\(_2\) platelets (Cohen et al., 2003), 2D photonic crystals (García de Abajo et al., 2003), Ag spheroids (Little et al., 1984), Ag nanotriangles (Nelayah et al., 2007a,b), Ag nanoshells (Eggeman et al., 2007), and gold nanoparticles ranging from single nanospheres and nanorods (Bosman et al., 2007; Schaffer et al., 2009) to touching dimers and interacting nanorods (Chu et al., 2009; N’Gom et al., 2008). The corresponding theoretical efforts needed to explain these measurements involve more elaborate analytical developments as well as numerical simulations based upon either Poisson’s equation in the non-retarded approximation or Maxwell’s equations if full retardation effects are necessary. These methods use in general local-response functions and assume sharp boundaries separating different dielectric phases.
1. Analytical methods

Several analytical studies of EELS have been carried out in the non-retarded approximation for geometries that can be naturally described using coordinate systems in which the Poisson equation is separable.

a. Ellipsoids The electrostatic modes of ellipsoids have been known for a long time, including analytical expressions for the polarizability (Gans, 1912; Jones, 1945). In particular, the EELS probability has been worked out for oblate ellipsoids by Illman et al. (1988).

b. Wedges Analytical solutions have been reported for the modes of sharp wedges (Davis, 1976; Dobrzenski and Maradudin, 1972), parabolic wedges (Eguiluz and Maradudin, 1976), and hyperbolic wedges (Davis, 1976). The latter display a discrete spectrum that suffers a singular transition in the limit to a sharp edge, characterized by a continuous spectrum. The EELS probability has been calculated for parabolic wedges (García-Molina et al., 1985b) with the beam running parallel to the edge.

c. Supported particles A sphere or a hemisphere intersected by a planar surface can be described by mode matching, consisting in expanding the potential into spherical harmonics and plane waves within different regions of the structure, thus generating a set of self-consistent linear equations upon application of the boundary conditions. This method, pioneered by Zaremba (1985) to explore the van der Waals interaction of an atom with a surface feature, was later applied by Wang and Cowley (1987a) to study plasmon modes of supported particles, which they measured in Al spheres sitting on AlF₃ (Wang and Cowley, 1987a) and Al surfaces (Wang and Cowley, 1987b). These authors incorporated nonlocal effects within the hydrodynamic approximation (Wang and Cowley, 1987c). Further work on this geometry focused on calculating the loss probability (Aizpurua et al., 1996; Rivacoba et al., 1992).

Several other geometries in which Poisson’s equation is separable are still unresolved, but the list of structures that can be accessed with analytical tools is short and we need in general to rely on numerical methods for more complicated shapes. We briefly discuss two of them in this section, and a third one is presented in Sec. III.G.

2. Non-retarded boundary element method

The boundary element method (BEM) consists in expressing the fields inside each homogeneous region of a given structure in terms of interface sources that are then self-consistently calculated by imposing the continuity of the potential and the electric displacement. This has the advantage that it requires a modest numerical effort, since it involves parametrization of boundaries rather than volumes.

The BEM can be traced back to Maxwell, who utilized it to calculate capacitances (Maxwell, 1891). It was pioneered by Fuchs (1975) in the computer age to simulate modes of ionic cubes, and later by Lu and Maradudin (1990) to find plasmons propagating inside channels cut into otherwise planar surfaces. Its debut in EELS was made by Ouyang and Isaacson (1989b), who obtained loss probabilities that they applied to study a particle supported on a substrate (Ouyang and Isaacson, 1989a). Subsequent developments focused on geometries characterized by either axial symmetry or translational invariance along a certain direction, thus permitting us to access a large number of complex morphologies with moderate computational effort (García de Abajo and Aizpurua, 1997), since only curves rather than surfaces need to be parametrized in these cases. This approach gives excellent results as compared to experiment for the losses of electrons passing near Si tips (Reed et al., 1999) or close to the edge of an MgO cube with different orientations of the trajectory (Aizpurua et al., 1999).

In the non-retarded BEM, the potential inside a given homogeneous region of space is separated into two terms,

$$\phi = \phi^{\text{ext}} + \phi^{\text{ref}}$$

the first of which describes the potential produced by the external charges present in that medium, treating it as if it were infinitely extended, while the second term accounts for reflection at the boundary. Assuming an inhomogeneous structure composed of two materials with dielectric functions $\varepsilon_1$ and $\varepsilon_2$, respectively, the interface $\mathcal{S}$ can be assimilated to an induced surface charge $\sigma$, so that the reflected potential becomes

$$\phi^{\text{ref}}(r) = \int_{\mathcal{S}} ds \frac{\sigma(s)}{|r - s|}.$$  

Inserting these expressions into Poisson’s equation, one finds the following self-consistent relation for the induced boundary charge (García de Abajo and Aizpurua, 1997):

$$\Lambda \sigma(s) = \mathbf{n}_s \cdot \nabla \phi^{\text{ext}}(s) + \int ds' F(s, s') \sigma(s'), \quad (38)$$

where

$$F(s, s') = \frac{-\mathbf{n}_s \cdot (s - s')}{|s - s'|^3},$$

$$\Lambda = 2\pi(\varepsilon_2 + \varepsilon_1)/(\varepsilon_2 - \varepsilon_1),$$

and $\mathbf{n}_s$ is the interface normal directed towards medium 2 (see Fig. 15). The right-hand side of Eq. (38) is actually the opposite of the electric field along the boundary normal, with the integral describing the field produced by induced interface charges. Equation (38) admits a...
representation in terms of the eigenvectors of \( F \), with eigenvalues denoted as \( 2\pi \lambda_j \). This leads to the following general expression for the loss probability (García de Abajo and Aizpurua, 1997):

\[
\Gamma_{\text{NR}}^{\text{NR}}(\omega) = \sum_{i,j} \text{Im} \left\{ -A_{ij} \left[ \frac{2}{(1 + \lambda_j)e_1 + (1 - \lambda_j)e_2} - \frac{1}{e_i} \right] \right\},
\]

where \( j \) runs over the materials 1 and 2, and \( A_{ij} \) are coupling constants that depend on the actual sample geometry and electron trajectory but not on \( \epsilon_1 \) or \( \epsilon_2 \). The second term inside the square brackets of Eq. (39) gives a negative loss probability corresponding to the begrenzungseffekt, the weight of which increases with \( A_{ij} \) in the same proportion as the interface mode described by the first term.

3. Retarded boundary element method

We can follow a similar procedure to incorporate retardation by solving Maxwell’s equations at the expense of a 6-fold increase in computational cost (García de Abajo and Aizpurua, 1998, 2002). First of all, the electric field can be expressed in terms of vector and scalar potentials as

\[
\mathbf{E} = ik \mathbf{A} - \nabla \phi.
\]

We find it convenient to work in the Lorentz gauge, in which \( \nabla \cdot \mathbf{A} = k \epsilon \phi \). Then, the potentials become

\[
\phi(\mathbf{r}) = \phi^\text{ext}(\mathbf{r}) + \int_{S_j} ds G_j(|\mathbf{r} - \mathbf{s}|) \sigma_j(\mathbf{s})
\]

and

\[
\mathbf{A}(\mathbf{r}) = \mathbf{A}^\text{ext}(\mathbf{r}) + \int_{S_j} ds G_j(|\mathbf{r} - \mathbf{s}|) \mathbf{h}_j(\mathbf{s})
\]

inside each homogeneous region \( j \) delimited by a boundary \( S_j \), where

\[
G_j(r) = \frac{\exp\left(ikr\sqrt{\epsilon_j}\right)}{r}
\]

is the Green function of Helmholtz equation inside that medium, \( \phi^\text{ext} \) and \( \mathbf{A}^\text{ext} \) are the potentials produced by external sources in an infinite homogeneous medium described by a dielectric function \( \epsilon_j \), and \( \sigma_j \) and \( \mathbf{h}_j \) are auxiliary interface charges and currents (see Fig. 15). The boundary conditions of the electromagnetic field are then imposed and one is left with a set of self-consistent surface integral equations that determine the boundary sources. For example, the continuity of the scalar potential leads to

\[
\int ds' \left[ G_1(\mathbf{s} - \mathbf{s}')\sigma_1(\mathbf{s}') - G_2(\mathbf{s} - \mathbf{s}')\sigma_2(\mathbf{s}') \right] = \phi_2^\text{ext}(\mathbf{s}) - \phi_1^\text{ext}(\mathbf{s}),
\]

where \( \phi_j^\text{ext} \) is the external potential in medium \( j \), evaluated at positions infinitesimally close to the interface. We proceed by discretizing the integrals in these equations through a finite number of representative points distributed on the interfaces. This leads to a linear set of equations that we solve by linear algebra techniques. Further details on the method and its generalization to an arbitrary number of different media have been given by García de Abajo and Howie (2002).

The BEM is particularly advantageous to simulate axially-symmetric samples, for which different azimuthal components \( m \) having \( \exp(im\phi) \) dependence on azimuthal angle are uncoupled. Then, the sample surface can be parametrized with a contour line, thus effectively reducing the problem to one dimension. Similarly, only the 1D contour defining the geometry needs to be parametrized for geometries with translational invariance along a given direction. These two special cases have been discussed in detail by García de Abajo and Howie (2002). However, the method is general and can be applied to arbitrary boundaries, as recently illustrated by Myroshnychenko et al. (2008a).

A recent example of application of the retarded BEM is presented in Fig. 16, adapted from Nelayah et al. (2007a), in which measured loss spectra for Ag nanoprisms [Fig. 16(a)] are compared to simulations [Fig. 16(b)] when the electron beam is aimed at the corner, the edge, or the center of the triangle. Different plasmon losses are observed at each of these positions. Moreover, one can obtain snap shots of these plasmons, as shown in the images of Fig. 16(c-h), by scanning the beam over the sample area and collecting the rate of selected energy losses corresponding to the three dominant features observed in the spectra of Fig. 16(a-b). These findings have been corroborated by tip-scattering NSOM measurement (Rang et al., 2008).

The fully retarded BEM has been used to investigate the optical response of nanoparticles (Aizpurua et al., 2003; Grzelczak et al., 2007; Hohenester and Kreun, 2005; Kumar et al., 2008; Myroshnychenko et al., 2008b; Sánchez-Iglesias et al., 2006) and extended nanostructures (Blanco and García de Abajo, 2004a; Cole et al., 2007), with particular emphasis placed on localized plasmon resonances, the coupling between atoms/molecules and nanoparticles (Blanco and García de Abajo, 2004b;
4. Multiple scattering approach

For specimens containing several neighboring particles, multiple scattering techniques provide a natural way of solving Maxwell’s equations in frequency space. The field induced by each of the particles adds up to the scattering produced by the other particles in a self-consistent fashion.

In practice, we need to express the electric field around each particle \( j \) in a complete basis set, for which vector spherical harmonics (Wang et al., 1993) are a convenient choice. The following specific steps are then followed to solve the electromagnetic problem in the presence of a passing electron (García de Abajo, 1999b): (1) the external field is scattered by each of the particles, giving rise to spherical outgoing waves (single-scattering contribution); (2) every outgoing multipole centered around a given object is expanded into propagating multipoles centered around each of the remaining objects; (3) these propagating multipoles are scattered by the objects around which they are centered, giving rise to outgoing waves that add up to the ones obtained after first-order scattering of the electron field; (4) this process is repeated iteratively, giving rise to the self-consistent relation

\[
\xi_{j}^{\text{ind}} = \xi_{j}^{\text{ss}} + t_{j} \sum_{j'} H_{jj'} \xi_{j'}^{\text{ind}},
\]

where \( \xi_{j}^{\text{ss}} \) and \( \xi_{j}^{\text{ind}} \) are vectors containing the coefficients of the outgoing-wave expansion of the single-scattering contribution and the self-consistent induced field, respectively, \( H_{jj'} \) is the propagation matrix that expresses outgoing waves centered around object \( j' \) in terms of propagating spherical waves centered around object \( j \), and \( t_{j} \) is the scattering matrix of object \( j \), containing the scattering outgoing-wave coefficients after interaction of propagating spherical waves with that object. Analytical expressions of \( t_{j} \) for homogeneous spheres are given in Appendix C. The multipolar expansions are truncated at some maximum orbital angular-momentum number \( l_{\text{max}} (\gtrsim ka) \) for particles of radius \( a \), so that the above equations contain \( L = 2N(l_{\text{max}} + 1)^2 \) variables, where \( N \) is the number of particles, and the factor of 2 stands for the fact that we have electric and magnetic multipoles, similar to the situation encountered in Eq. (33) for the single sphere. Direct inversion of the full system of equations takes a prohibitive computation time proportional to \( L^3 \). Instead, we can utilize a highly convergent iterative procedure based upon Lanczos’ method (see Haydock, 1980) to solve these equations (García de Abajo, 1999a,b). These computations demand an effort proportional to \( L^2 \) and yield highly converged spectra for electrons passing near clusters of up to several hundred particles in a matter of minutes with \( l_{\text{max}} \sim 15 \).

An example of application of this method is presented in Fig. 17, taken from Cai et al. (2009). A 100 keV electron is considered to pass near the end of a linear array formed by 51 spherical gold particles [Fig. 17(a)]. The spectral distribution of the electric field \( |\mathbf{E}^{\text{ind}}(\mathbf{r}, \omega)| \),...
plotted in Fig. 17(b) along a segment parallel to the array exhibits prominent features at the localized plasmon resonances of the spheres. The time-domain picture [Fig. 17(c), obtained from Eq. (1)] shows plasmon propagation mediated by inter-particle interaction. The group velocity (dashed curve) is consistent with the slope at the intensity-maximum of the guided-mode dispersion relation derived from the space Fourier transform of $|\mathbf{E}^{\text{ind}}(r, \omega)|$ [Fig. 17(d)].

F. Composite materials

A common situation arises when the sample contains composites made of a fine-scale mixture of different dielectric grains. Finding the optical response of the resulting metamaterial is not easy, specially in cases for which the actual microscopic geometry is unknown. This is even more complicated in EELS due to the local nature of the electron probe.

A first naive approach to the problem of explaining loss spectra in a composite consists in considering the appropriately weighed bulk probabilities derived from Eq. (19) for the constituents. The result is a reasonable approximation when the grains are large enough so that interface losses are small in front of volume losses. This type of analysis has been used to map water distributions in biological specimens using valence excitations (Sun et al., 1991), which are more intense than core losses and provide similar information in this case, accompanied by lower sample damage.

However, this approach breaks down for finer mixtures, as shown by Howie and Walsh (1991) through studies of losses in Al spheres created in AlF$_3$ by the ionizing action of the electron beam. They found that the spectra were well reproduced by an expression like Eq. (39) with $\lambda_l = 1/3$ (Howie, 2003; Howie and Walsh, 1991), which results from the dipolar mode of the spheres [Eq. (29) for $l = 1$], now corrected by the bulk losses due to the begrenzungseffekt. Similar conclusions were extracted for other systems, like Ni-Al mixed oxides (Cadete Santos Aires et al., 1993).

The Maxwell-Garnett (1904, 1906) effective medium theory, which yields good results in many cases, is particularly suited to describe small spheres of a given material 2 embedded in a matrix of material 1, and works extremely well for small volume fractions of the spheres, $f \lesssim 0.3$. This theory predicts an effective dielectric function

$$
\epsilon_{\text{eff}} = \epsilon_1 \frac{2(1-f)\epsilon_1 + (1+2f)\epsilon_2}{(2+f)\epsilon_1 + (1-f)\epsilon_2},
$$

accounting for the spheres through their non-retarded dipolar response [$\alpha_{\text{NR}}^{p}$ in Eq. (B5)]. Not surprisingly, the $l = 1$ mode of Eq. (29) ($2\epsilon_1 + \epsilon_2 = 0$) is recovered from the zero of the denominator of Eq. (43) in the small $f$ limit. The Maxwell-Garnett theory has been successfully employed by McComb and Howie (1995) to explain EELS measurements in SiO$_2$ polymorphs of different density.

Corrections to the loss function of granular materials beyond the dipolar terms captured in Maxwell-Garnett theory have been explored by several authors (Fu et al., 1993; Fu and Resca, 1993). In particular, Barrera and Fuchs (1995) obtained a nonlocal effective dielectric function for a random system of spheres described their local bulk permittivity. This work was followed by the derivation of a spectral representation of an effective local response for the same system (Fuchs et al., 1996). More complicated situations have been considered, such as an afoof electron moving above ordered and disordered arrays of spheres (Mendoza et al., 1998, 1999), and self-similar structures formed by a continuous distribution of sphere sizes (Sosa et al., 2001), all of them analyzed in the non-retarded limit.

In this context, the remarkable works of McKenzie and McPhedran (1977) and McKenzie et al. (1978) provide analytical expansions for the effective local response of cubic arrangements of spheres with inclusion of an arbitrarily large number of multipoles, which directly yield the loss function $\text{Im}\{-1/\epsilon_{\text{eff}}\}$. However, a similar derivation of EELS formulas for these structures is still pending, although they could shed light into the open discussion of nonlocal effects in metamaterials.

Retardation corrections to the effective dielectric function can be obtained from the photonic bands of periodic composites in the long-wavelength limit. This approach was followed by Pendry and Martín-Moreno (1994) to explain the $\sim 8$ eV loss feature measured by Howie and Walsh (1991) in aggregates of small aluminum particles. Their simulations were accompanied by a complex series of spectral features that subsequent analysis by García de Abajo and Blanco (2003) demonstrated to be due to numerical inaccuracies. In fact, the loss spectrum for small filling fraction ($f = 0.06$) of an array of equally-spaced aluminum spheres already shows a single $\sim 8.5$ eV feature (García de Abajo and Blanco, 2003) [see Fig. 18(a), solid curves]. This peak splits into two for larger $f = 0.3$, due to multipolar interaction between neighboring spheres, quite different from the individual-plasmon excitation predicted by Maxwell-Garnett theory [Fig. 18(b), dashed curves, obtained from Eq. (43)]. Interestingly, the calculated EELS peaks in Fig. 18 are displaced to the red with respect to the maxima of the surface loss function $\text{Im}\{-1/(1 + \epsilon_{\text{eff}})\}$ as a result of retardation, similar to the effect discussed in Fig. 11. Both the EELS probability and the effective dielectric function $\epsilon_{\text{eff}}$ are calculated here using a layer Korringa-Kohn-Rostoker (KKR) method, which allows solving the multiple-scattering equation (42) for parallel layers of spheres (García de Abajo and Blanco, 2003; Stefanou et al., 2000). The aluminum is described by Eq. (A1) with $\hbar \omega_p = 15$ eV and $\hbar \eta = 1$ eV. A value of $l_{\text{max}} = 6$ is sufficient to achieve convergence. We plot for comparison the loss probability computed in the absence of multiple-scattering interaction among spheres [Fig. 18(a), broken curves], predicting a single peak that is roughly independent of $f$. 
FIG. 17 (Color online) Electron-beam-induced plasmon generation in gold nanoparticle chains (taken from Cai et al., 2009). (a) Scheme of the structure and geometry under consideration. (b) Induced electric field $|E^{\text{ind}}(x,\omega)|$ (in frequency space $\omega$) produced by the electron along the points of the \(AB\) segment [see red double arrow in (a), \(\parallel \hat{x}\)]. (c) Same as (b), in the time domain: $|E^{\text{ind}}(x,t)|$ [see Eq. (1)]. (d) Fourier transform of $E^{\text{ind}}(x,\omega)$ with respect to $x$, showing the dispersion relation of plasmons excited by the electron and propagating along the chain. The velocity of the launched pulse is found to be $\sim 0.75c$ [dashed line in (c)], in good agreement with the group velocity at the maximum of excitation probability along the guided-mode dispersion curve [dashed line in (d)].

G. Carbon molecules and low-dimensional structures: The discrete-dipole approximation

Carbon molecules and other low-dimensional structures have fascinated chemists and physicists since the discovery of C\(_{60}\) (Kroto et al., 1985) and carbon nanotubes (Iijima, 1991; Monthioux and Kuznetsov, 2006), followed by BN nanotubes (Chopra et al., 1995) and other related molecules.

Plasmon modes were first observed in solid C\(_{60}\) as 6.5 eV and 28 eV EELS features (Gensterblum et al., 1991; Saito et al., 1991), corresponding to collective oscillations of $\pi$ and $\sigma$ electrons, respectively. The optical properties of the individual molecules is not significantly altered in the solid, since their binding interaction is basically due to weak van der Waals attraction. Actually, subsequent studies in gas phase C\(_{60}\) molecules corroborated these results (Keller and Coplan, 1992).

The clear-cut distinction between loosely-bound $\pi$-like orbitals and tightly-bound $\sigma$-like orbitals persisted in the loss spectra of carbon nanotubes (Kuzuo et al., 1992), in both single-wall and multishell configurations (Stéphan et al., 2002), in multishell fullerenes (Henrard et al., 1999), in BN nanotubes (Kociak et al., 2000), and in other similar types of molecules (Kociak et al., 2001).

These results were soon related to the dielectric properties of graphite, the anisotropic permittivity of which was used as input to describe fullerenes and nanotubes via continuous dielectric theory, with different $\epsilon$ for field components parallel and perpendicular to the carbon shells (Lucas et al., 1994). The dielectric approach yielded reasonable results compared to measurements of plasmons sustained by a diverse number of geometries and chemical identities, including multishell BN nanotubes, C nanotubes, and C fullerenes (Henrard et al., 1999; Kociak et al., 2000; Stéphan et al., 2002), WS\(_2\) hollow nanoparticles (Kociak et al., 2001), and WS\(_2\) nanotubes (Taverna et al., 2002). Good agreement was obtained for nanotubes of different thicknesses, down to the single-wall level (Stéphan et al., 2002).
Also, first-principles calculations have been carried out to describe the interaction of fast electrons with carbon nanotubes (Nojeh et al., 2006). The agreement with experimental loss spectra of single-wall tubes is quite satisfactory (Marinopoulos et al., 2003). However, this approach is currently limited to relatively small structures, due to the heavy computational demand of spectroscopy ab initio methods.

The discrete nature of these molecules suggests an atomistic approach to describe their response, with each atom represented by an induced dipole of different strength in the directions out-of-plane and along the tangent plane of the atomic shells. This is the so-called discrete-dipole approximation (DDA), in which the atomic polarizability is obtained from the dielectric function of the bulk material (e.g., graphite for carbon nanotubes and fullerenes) by inverting the Clausius-Mossotti relation.\(^{12}\) A detailed comparison between the DDA and the continuous dielectric theory was carried out by Henrard and Lambin (1996), who showed that both approaches are in excellent mutual agreement.

The DDA was pioneered by Purcell and Pennybacker (1973) in the study of the optical response of dielectric grains by dividing them into fine volume elements that were described through induced dipoles. Actually, the DDA is a versatile method for dealing with arbitrary sample morphologies and is extensively employed to predict experimental loss spectra of single-wall nanotubes (Nojeh et al., 2002). This versatility is currently limited to relatively small structures, due to the heavy computational demand of spectroscopy of all dipoles \(\mathbf{p}_j\), the matrix \(G^0\) is made of the components of \(G^0_{jj'}\), and so on. The electric field corresponding to a passing electron is given by Eq. (4), that is, \(\mathbf{E}^{\text{ext}}(\mathbf{r}) = (2\epsilon \omega /v^2\gamma) \mathbf{g}_j\), where \(\mathbf{g}_j = \mathbf{g}(\mathbf{r}_j)\) [see Eq. (5)].

Using the same matrix notation as in Eq. (45), the fully-retarded loss probability in the presence of our polarizable point particles representing carbon atoms reads

\[
\Gamma_{\text{EELS}}(\omega) = \frac{1}{h\pi} \left( \frac{2\epsilon \omega}{v^2\gamma} \right)^2 \text{Im} \left\{ g^+ \cdot \frac{1}{\alpha^{-1} - G^0} \cdot g \right\}. \tag{46}
\]

Notice that this expression reduces to Eq. (34) in the single-atom case (\(G^0 = 0\)).

As an example of application of Eq. (46), Fig. 19 shows the loss probability for an electron passing near a single-wall carbon nanotube under grazing incidence conditions.

\[\text{FIG. 19 (Color online) Probing plasmons in a single-wall carbon nanotube via EELS. The experiment shows prominent \(\pi\) and \(\sigma\) plasmons that are relatively well described both by the discrete-dipole approximation (DDA) and by a dielectric approach. The experimental curve and the dielectric theory have been taken from Stéphan et al. (2002). The DDA calculation performed here for a 16 \times 16 nanotube is based upon the dielectric function of graphite, taken from Palik (1991).}\]

\[\text{Equation (44) can be solved by direct matrix inversion to yield}\]

\[
p = \frac{1}{\alpha^{-1} - G^0} \cdot \mathbf{E}^{\text{ext}}, \tag{45}\]

where \(p\) represents the vector formed by the components of all dipoles \(\mathbf{p}_j\), the matrix \(G^0\) is made of the components of \(G^0_{jj'}\), and so on. The electric field corresponding to a passing electron is given by Eq. (4), that is, \(\mathbf{E}^{\text{ext}}(\mathbf{r}) = (2\epsilon \omega /v^2\gamma) \mathbf{g}_j\), where \(\mathbf{g}_j = \mathbf{g}(\mathbf{r}_j)\) [see Eq. (5)].

\[\text{In order to apply the DDA to carbon structures, each of the atoms (or alternatively, each of the bonds in a similar formulation) is assimilated to an induced point particle of polarizability tensor \(\alpha(\omega)\). The dipoles induced on these points in response to an external electric field \(\mathbf{E}^{\text{ext}}(\mathbf{r})\) are then obtained from their self-consistent interaction, so that the dipole of the particle located at position \(\mathbf{r}_j\) is given by}\]

\[
\mathbf{p}_j = \alpha \cdot \left[ \mathbf{E}^{\text{ext}}(\mathbf{r}_j) + \sum_{j' \neq j} G^0_{jj'} \cdot \mathbf{p}_{j'} \right], \tag{44}\]

where the \(3 \times 3\) symmetric tensor \(G^0_{jj'}\) describes the coupling with the dipole at \(\mathbf{r}_{j'}\).\(^{13}\)

\[\text{12 For carbon structures, the effective polarizability per atom is obtained from the dielectric function of graphite using } \alpha^{-1} = (4\pi n_C/3) \left[ (\epsilon_i + 2)/(\epsilon_i - 1) - S_i \right], \text{ where } n_C = 113.63 \text{ nm}^{-3} \text{ is the atomic density of this material, } i = ||, \perp \text{ refers to the response to an electric field either parallel or perpendicular to the carbon planes, and the constants } S_{\perp} = -2S_{||} = 2.8216 \text{ originate in the lack of inversion symmetry of the graphite atomic lattice (Ashcroft and Mermin, 1976).}\]

\[\text{13 The electric field created by dipole } \mathbf{p}_{j'} \text{ is given by} \]

\[
G^0_{jj'} \cdot \mathbf{p}_{j'} = \frac{e^{iku}}{u^3} \times \left\{ \left[ (ku)^2 + ku - 1 \right] \mathbf{p}_{j'} - \left[ (ku)^2 + 3ku - 3 \right] \frac{(\mathbf{r} \cdot \mathbf{p}_{j'}) \mathbf{u}}{u^2} \right\},
\]

where \(u = \mathbf{r}_j - \mathbf{r}_{j'}\). This quantity is related to the Green tensor \(G\) of Eq. (12) in vacuum through \(G^0 = -4\pi\omega^2G\).
The experimental data and the results of the anisotropic dielectric theory have been taken from Stéphan et al. (2002). The position of the plasmon features predicted by the above DDA formalism compares very well to both of them. It is remarkable that dielectric theory is still producing meaningful results for such thin nanotubes, although the DDA seems to attribute a relative weight to $\pi$ and $\sigma$ plasmons in better agreement with the observed spectrum.

H. Relation to the photonic local density of states

We can ask ourselves how exactly the loss probability is related to the photonic properties of the sample and whether it is possible to retrieve from measured data for bounded media a relevant quantity similar to the dielectric function. This question has positive answer in systems with translational invariance along the direction of electron motion: the EELS probability is proportional to the photonic local density of states (LDOS) (García de Abajo and Kociak, 2008b).

The photonic LDOS is defined by analogy to its electronic counterpart in solid state physics as the combined local intensity of all normalized eigenmodes of the system (Ashcroft and Mermin, 1976), but unlike electronic wave functions, the electric field is a vectorial quantity, which suggests defining separate Cartesian projections of the LDOS along unit vectors $\hat{n}$ as (Fussell et al., 2005)

$$\rho_n(r, \omega) = -\frac{2\omega}{\pi} \text{Im} \{\hat{n} \cdot G(r, r, \omega) \cdot \hat{n}\}$$  

(47)

in terms of the Green tensor defined in Eq. (12). The uniform LDOS in free space is known from black-body theory:

$$\rho_n^0(r, \omega) = \frac{\omega^2}{3\pi^2 c^3},$$

which contains an additional factor of $1/3$ arising from the projection over a specific Cartesian vector $\hat{n}$.

If we choose the direction of translation invariance along the $\hat{z}$ axis, the Green tensor $G(r, r', \omega)$ depends on $z$ and $z'$ only via $z - z'$, and consequently we can write

$$G(r, r', \omega) = \int \frac{dq_z}{2\pi} \tilde{G}(R, R', q_z, \omega) \delta(q_z(z' - z)).$$  

(48)

Using this in Eq. (13), the loss probability can be recast as

$$\Gamma_{\text{EELS}}(R_0, \omega) = \frac{4c^2 L}{\hbar} \text{Im}\{-G_{zz}(R_0, R_0, q_z, \omega)\}$$

$$= \frac{2\pi c^2 L}{\hbar \omega} \rho_z(R_0, q_z, \omega),$$

(49)

where $q_z = \omega/\nu$ is determined by energy conservation [see Eq. (3)], $L$ is the length of the trajectory, and

$$\rho_n(R, q_z, \omega) = -\frac{2\omega}{\pi} \text{Im} \{\hat{n} \cdot \tilde{G}(R, R, q_z, \omega) \cdot \hat{n}\}$$

is the photonic density of states, local in real space along the $R$ directions perpendicular to the beam and local in wavevector space $q_z$ along the direction of motion, $\hat{z}$. This relation between the photonic LDOS and the EELS probability is further illustrated in Sec. IV.D.1 for electrons moving inside 2D photonic crystals.

For a non-lossy homogeneous dielectric of real permittivity $\epsilon$, one finds $\rho_n(R, q_z, \omega) = (\omega/2\pi c^2)(1 - q_z^2/k^2\epsilon) \theta(\nu^2\epsilon - c^2)$, which reflects the fact that only electrons moving faster than light in the material can lose energy [see Eq. (7)]. This is the Cherenkov effect that we discuss in Sec. IV.D. As expected, the vacuum density of states does not contribute to the EELS signal.

Equation (49) constitutes a rigorous relation between the EELS probability and the LDOS valid for 2D systems with translational invariance along the direction of electron motion. The LDOS enters this relation through its projection over specific components $q_z$ of light wavevector along that direction. A similar relation can be obtained for fully arbitrary geometries from Eq. (13) (García de Abajo and Kociak, 2008b):

$$\Gamma_{\text{EELS}}(R_0, \omega) = \frac{4c^2 L}{\hbar} \text{Im}\{-G_{zz}^{\text{ind}}(R_0, R_0, q_z, -q_z, \omega)\},$$

$$= \frac{2\pi c^2 L}{\hbar \omega} \rho_z^{\text{ind}}(R_0, q_z, \omega),$$

where $G_{zz}^{\text{ind}}(R, R', q_z, -q_z', \omega)$ is the Fourier transform of the induced Green tensor with respect to $z$ and $z'$, $q_z = \omega/\nu$, and we have defined, by analogy to Eq. (47), the quantity

$$\rho_n(R, q, \omega) = -\frac{2\omega}{\pi} \text{Im} \{\hat{n} \cdot G(R, R, q, -q, \omega) \cdot \hat{n}\}$$

as a generalized density of states that is local in real space along the $R$ directions and local in momentum space along the remaining $z$ direction. In 2D systems, we have $\rho_n = L\rho_n$. 

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14 There is a univocal correspondence between the spatially-dependent local dielectric function and the Green tensor, as can be clearly deduced upon inspection of Eq. (12). This correspondence works frequency by frequency. Furthermore, the photonic LDOS is directly obtained from the Green tensor following Eq. (47). The inverse relation is less direct, although one can extend the equivalent of the Hohenberg and Kohn (1964) theorem to light, thus asserting that there is a univocal correspondence between dielectric functions and LDOS, with both quantities defined within a finite frequency range above $\omega = 0$ (i.e., the correspondence is not local in frequency).

15 The definition of Eq. (47) assumes that the photon eigenmodes are well defined (that is, in the absence of absorption) (Colas des Francs et al., 2001). An alternative interpretation, which holds even in the presence of lossy materials (D’Aguanno et al., 2004), comes from the realization that $(4\pi^2\omega D^2/\hbar) \rho$ gives the spontaneous decay rate for an excitation dipole strength $D$ (Barnett and Loudoun, 1996; Fussell et al., 2004). It should be also noted that a complete definition of the LDOS must include a magnetic part (Joulain et al., 2003), which is however uncoupled to our fast electrons.
I. Electronic structure determination

The publications containing EELS studies of local electronic structures are so numerous that only a succinct review is possible here. A spectacular degree of spatial resolution is achieved by resorting to high-energy losses, for instance when mapping $sp^2$ and $sp^3$ carbon in a silicon-diamond interface with sub-nanometer detail (Muller et al., 1993), and also when imaging K atoms in doped fullerenes (Guan et al., 2005). However, valence losses can provide excellent resolution too with minimum sample damage, as illustrated by Daniels et al. (2003), who mapped carbon chemical states through characteristic bulk plasmon losses with nanometer detail. In general, comparison with \textit{ab initio} theory greatly facilitates the identification of spectral features in the measured loss spectra (Keast and Bosman, 2008; Marinopoulos et al., 2003).

The determination of band gaps in insulators is successfully addressed by valence EELS (Dorneich et al., 1998), although this requires special care to deal with the subtraction of the ZLP and to get rid of CR losses, which can be misleading in the assessment of the inelastic signal threshold (Gu et al., 2007; Stöger-Pollach, 2008; Stöger-Pollach et al., 2006; Stöger-Pollach et al., 2008; Stöger-Pollach and Schattschneider, 2007).

The loss probability has been shown to be closely related to the so-called joint density of states (JDOS) (Pflüger et al., 1984, 1985), which is the convolution of density of states of occupied and unoccupied electronic bands. The JDOS gives an intuitive measure of the interband-transition probability when more subtle effects in the excitation matrix elements and lifetime effects are neglected. The JDOS predicts a dependence of the loss probability on the energy transfer $\hbar \omega$ as $\sim (\hbar \omega - E_g^d)^{1/2}$ and $\sim (\hbar \omega - E_g^i)^{3/2}$ at the edge of direct- and indirect-gap energies, $E_g^d$ and $E_g^i$, respectively (Rafferty and Brown, 1998). A good example of this is found in GaN, as shown in Fig. 20, adapted from Lazar et al. (2003). A direct gap involves a more abrupt decay of the loss probability near the gap [Fig. 20(a)-(d)]. Interestingly, both the smooth tail of an indirect gap and the sharper edge of a higher-energy direct gap are clearly resolved in the loss spectra of GaN [Fig. 20(e)].

In this context, Van Hove singularities (Van Hove, 1953) appearing in the electronic density of states at energies corresponding to maxima, minima, and saddle points of the bands have been also resolved using valence EELS (Dorneich et al., 1998).

Finally, the spatial resolution of electron microscopes can be employed at its best to study electronic bands of nanostructures. For instance, this has been done to measure optical gaps in single-, double-, and triple-wall BN nanotubes (Arenal et al., 2005), showing that the gap is rather independent of thickness and stays near the in-plane value of hexagonal bulk BN ($\sim 5.8$ eV). In a separate development, quantum confinement effects in the gaps of individual CdSe quantum dots have been observed using space-resolved EELS (Erni and Browning, 2007).

IV. CATHODOLUMINESCENCE: GENERATION OF LIGHT BY INCOMING ELECTRONS

Cathodoluminescence (CL) was first observed in the mid-nineteenth century as light emission produced when cathode rays (electrons) struck the glass of evacuated discharge tubes. Since then, a whole industry has been generated in the search for outstanding phosphorescent materials (Ozawa, 1990), and in particular, CL has been extensively used in TV screens, and even TEMs have benefited from it by projecting sample images in phosphor screens.

Cathodoluminescence offers a method for characterizing microstructures causing minimum sample damage. It can be regarded as a contactless technique under the aloof configuration (see Sec. III.C). CL is widely utilized in mineralogy (Pennycook, 2008; Pratesi et al., 2003; Schieber et al., 2000) to resolve the composition, contamination, and defects of natural structures with sub-micron detail. It is also a common tool to study the electronic bands of insulators (Balberg and Pankove, 1971) and semiconductors (Shubina et al., 2004; Yacobi and Holt, 1986), and particularly the effect of dopants (Holt and Yacobi, 1990). Recently, CL has been used as a means to read stored information by tracing phase transitions in Ga nanoparticles (Denisyuk et al., 2008), which can be in turn controlled by electron excitation (Pochon et al., 2004). Cathodoluminescence is typically performed in adapted SEMs using 1-50 keV beams, and it is now possible to record spectra for every pixel of a given image (Galloway et al., 2003). In an separate development, low-energy electrons (150-300 eV) emitted from scanning tunneling microscope tips have been employed to produce surface CL, which benefits from the small inelastic mean free path of electrons at those energies (Ma et al., 2006).
Here, we are interested in the application of CL to investigate the optical properties of nanostructured materials. Besides e-h pair recombination, which dominates CL in semiconductors, we are interested in other mechanisms of light emission such as excitation of leaky plasmons in metal nanoparticles. Overall, CL constitutes an excellent technique to access optical properties with nanometer resolution and without spurious effects due to modifications produced by the presence of a probe in the region near the sample, as is the case in NSOM. A close relative of CL is found in light emission accompanying inelastic electron tunneling, which can be assisted by localized plasmon excitations (Persson and Baratoff, 1992), although the photon yield is very low in that case.

A. Mechanisms of light emission

We find it convenient to classify the mechanisms of CL emission according to their degree of coherence with respect to the external electric field of the bombarding electrons.

1. Coherent electron-induced radiation emission

The interaction of the electron with a material gives rise to electromagnetic far-field components that are coherent with respect to the external evanescent field accompanying the moving charge, described by Eq. (4). These far-field components produce what we call electron-induced radiation emission (EIRE). There are several mechanisms of coherent light emission, the most notorious of which are transition radiation (Sec. IV.C), Cherenkov radiation (Sec. IV.D), and diffraction radiation (Sec. IV.E). The electric field associated to each of these mechanisms is subject to the same set of Maxwell equations, so that we must observe interference effects between them.

The interference of coherently emitted light is actually due to the fact that the quantum-mechanical state of the sample does not change after the emission process takes place. All the energy released by the electron in a scattering event giving rise to EIRE is liberated in the form of radiation, as schematically shown in Fig. 21(a).

EIRE sources have a relatively delocalized origin, since they involve regions of the sample that are large from the electronic point of view, although they can be small compared to the wavelength of the emitted light. As a consequence, the coherent photon yield increases with electron energy: the faster the charge, the more extended the external field (see Sec. II.A) and the more efficient the coupling to delocalized excitations.

When heavy encounters with sample atoms take place, the sudden change in the electron trajectory gives rise to radiation. This is the so-called bremsstrahlung (Jackson, 1999), which can maintain certain degree of coherence with the sources of EIRE discussed so far. In particular, theoretical analysis has shown that bremsstrahlung produces interference with transition radiation (TR) upon electron bombardment (Ritchie et al., 1964; Shieh and Ritchie, 1970). Experimental evidence of this effect has been reported under grazing incidence conditions (Cram and Arakawa, 1967), so that deflections in the electron trajectory occur close to the surface. However, the contribution of bremsstrahlung to the total photon yield under usual TEM and SEM operation conditions is minor, unless very grazing trajectories are considered (Tomaš et al., 1972, 1974).

2. Incoherent cathodoluminescence

The incoherent part of the emission bears close relation to photoluminescence: the sample is first excited and then it decays inelastically by emitting light of a different color, displaced to the red with respect to the external illumination. The only difference lies in the source of excitation: while external light preferentially couples to strong-dipole modes, electrons can efficiently excite metastable levels because they are capable of transferring relatively large momenta to the sample (see, for example, Fig. 8) that enhances the probability of non-dipole transitions. Furthermore, the electron field can be regarded as a super-continuum source of evanescent light, suitable for creating energetic excitations (for instance, in large band-gap insulators) that would be difficult to produce using a laser instead.

Figure 21(b) shows a typical incoherent emission process, whereby the excited state produced by the electron decays into an intermediate state, giving rise to light of frequency $\omega'$ below the transferred $\omega$. The sample changes its quantum-mechanical state after the interaction with the electron and the optical emission take place, and consequently, the resulting light cannot produce interference with coherent EIRE. This contribution to CL is similar to the outcome of excited-atom decay (i.e., proportional to the radiative part of the photonic LDOS).

Incoherent emission is generally associated to the creation of e-h pairs and excitons. Actually, CL is routinely used to investigate these types of excitations (Koizumi et al., 1972, 1974).

FIG. 21 (Color online) Some of the processes taking place during energy transfer from a swift electron to a sample. (a) Excitation from the specimen ground state g to an excited state e, and subsequent radiative decay of the latter (see Sec. IV.A.1). (b) Excitation followed by incoherent radiative decay to an intermediate state i (see Sec. IV.A.2). (c) Excitation followed by non-radiative decay.
et al., 2001; Rodt et al., 2005; Sauer et al., 2000), and particularly when one is interested in resolving them in space. For instance, to study quantum dots (Grundmann et al., 1995; Rodt et al., 2005), quantum-confinement in semi-conductors (Chen et al., 2006; Yamamoto et al., 2006a), and optical effects driven by surface atomic steps (Minoda and Yamamoto, 2006).

We should expect the opposite dependence of the incoherent emission yield on electron velocity as compared to coherent EIRE. The excitations in this case are highly localized, so that the strength of the coupling to the more delocalized field of faster electrons turns out to be weaker. This effect is actually reflected in the energy dependence of the electron inelastic-mean-free-path (e-imfp) (Tanuma et al., 1994), represented in Fig. 1: this quantity passes through a minimum at \( \sim 50 \) eV, in which inelastic excitation is more probable, and exhibits a monotonic increase with energy due to weaker Coulomb interaction when the electron moves faster. Incoherent CL can take very large values in suitably designed in-space. For instance, to study quantum dots (Grundmann et al., 2002; Karali et al., 2005).

Incoherent CL can be regarded as a way to deliver a localized source of light (the inelastic excitations) with which one can investigate local photonic properties. It is possible thanks to the effect predicted by Purcell (1946): an increase in the radiative decay rate of excited nuclei coupled to a resonant circuit. The Purcell effect is quantified by a linear dependence of the light emission rate of excited atoms on the photonic LDOS (see Sec. III.H) that has been observed in numerous experiments (see Andrew and Barnes, 2001, and references therein). But of course, this decay channel must compete with non-radiative processes and their relative magnitudes depend on the dielectric environment. As a thumb rule, atomic radiative decay dominates in insulators and is negligible in metals.

A microscopic quantitative analysis of incoherent CL is made more complicated by the excitation of hot electrons, which are in turn a source of CL. Some of these electrons escape from the sample surface, giving rise to SEE, the yield of which can reach values above 20 electrons per primary electron in some insulators at \( \sim 1 \) keV incidence energy (Cazaux, 2006). This high emission is due to hot charge carriers right above the insulator gap, characterized by long mean free paths and therefore larger probability of escaping from the solid (Ortega et al., 1998).

Secondary electrons of significant energy above the Fermi level are less abundant in metals because they are rapidly quenched by strong dynamical screening. Moreover, the CL emission comes from the skin-depth sub-surface layer (\( \sim 20 \) nm in the visible and near-IR for Al and noble metals), which drastically erases any effects arising from the electron cascade that is produced by the energetic impinging probe in deeper regions.

As noted above, the incoherent CL emission must compete with other relaxation channels, like Auger de-excitation [see Fig. 21(c)], which are relatively inefficient in insulators. However, radiative inelastic decay gives rise to just a minor contribution to CL in metals because electronic relaxation channels are several orders of magnitude faster. Therefore, the CL signal arising from metallic nanostructures, such as the ones employed in plasmonics, is dominated by EIRE, the details of which we discuss in the next sections.\footnote{A word of caution should be added here: while EIRE is dominant at relatively-high electron-beam energies (for example, above 10 keV), it decays and is overcome by incoherent CL at sufficiently-low electron energies.}

\[ \text{B. Calculation of coherent light emission} \]

The intensity of light emission produced by swift electrons interacting with a sample can be calculated from the induced far field. Using the retarded BEM to solve Maxwell’s equations (Sec. III.E.3), we obtain the far field from the integrals of Eqs. (40) and (41) in the \( kr \rightarrow \infty \) limit. The radiation is commonly produced and measured in the vacuum medium \( j = 1 \), in which \( \varepsilon_1 = 1 \) and \( G_1(|r-s|) \approx \exp(\text{i}(kr - k\hat{r} \cdot s))/r \) (for \( kr \gg 1 \)), with \( \hat{r} = r/r \). Then, the induced field reduces to

\[ \mathbf{E}^\text{ind}_{kr \rightarrow \infty} \to f(\Omega) e^{ikr}/r, \]

where \( \Omega \) denotes the orientation of \( r \), and the far-field amplitude \( f = \tilde{f} - (\hat{f} \cdot \hat{r})\hat{r} \) is expressed in terms of the boundary current on the vacuum side as

\[ \tilde{f}(\Omega) = ik \int_{S_1} ds e^{-ik\hat{r} \cdot s} \mathbf{h}_1(s). \quad (50) \]

Here, we are relying on the fact that the contribution of \( -\nabla \phi \) to \( \mathbf{E}^\text{ind} \) only produces longitudinal components. However, the far field is transversal, and consequently, entirely coming from the \( k \mathbf{A} \) contribution to \( \mathbf{E} \). This gives rise to \( \tilde{f} \) [Eq. (50)], from which the longitudinal part has been subtracted to obtain \( f \).

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The EIRE intensity can be obtained from the Poynting vector integrated over emission directions. The emitted energy per incoming electron is then given by
\[
\Delta E = \frac{c}{4\pi} \int dt \int d\Omega \cdot [\mathbf{E}(r, t) \times \mathbf{H}(r, t)].
\]
Now, expressing the fields in time Fourier transform [Eq. (1)] and using the far-field limit just discussed, \( \Delta E \) can be decomposed into photon-energy components \( h\omega \) as
\[
\Delta E = \int_0^\infty h\omega \, d\omega \int d\Omega \, \Gamma_{\text{CL}}(\Omega, \omega),
\]
where
\[
\Gamma_{\text{CL}}(\Omega, \omega) = \frac{1}{4\pi^2 h\omega} |f(\Omega)|^2
\]
is the number of photons emitted per incoming electron, per unit of solid angle of emission \( \Omega \), and per unit of photon frequency range \( \omega \).

C. Transition radiation

When a fast charge impinges on a metal surface, the sudden annihilation of its image acts like an induced dipole that produces radiation, as illustrated in Fig. 22. This effect was predicted by Ginzburg and Frank (1946) and observed by Goldsmith and Jelley (1959) in the visible range for protons incident on Ag, Al, and Au surfaces. In fact, transition radiation occurs whenever a swift electron crosses the interface between different dielectric phases.

A detailed exposition of TR was given by Ter-Mikaelian (1972), and a particularly useful result was produced by Kröger (1968), who reported a closed-form expression for the EELS probability of an electron traversing a thin film under arbitrary oblique incidence, including the contributions of Cherenkov modes and TR. This formula has been valuable in devising methods to eliminate retardation effects from measured spectra and to retrieve bulk dielectric functions (Stöger-Pollach, 2008). In a related development, the theoretical analysis of the radiation produced by an electron passing by the center of a sphere showed an interesting evolution from CR (see Sec. IV.D) to TR when considering large or small spheres compared to the wavelength (Pogorzelski and Yeh, 1973).

On the experimental side, the results of Yamamoto et al. (1996a,b) for CR and TR in mica, Ag, Al, and Si films, including detailed comparison with theory, constitute an outstanding reference that illustrates the complex interplay between both types of radiation emission channels in dielectric films as compared to metals (for example, Ag and Al show relatively featureless backward emission spectra for wavelengths above the bulk plasmons, in contrast to mica or Si). In a separate work, Borziak et al. (1976) reported CL spectra for 5-500 eV electrons impinging on Ag, Au, Bi, Cu, and W surfaces, and showed a dense series of features that might originate in incoherent CL, since coherent EIRE is strongly suppressed at such low energies (see Sec. IV.A.1). More recently, relativistic beams (6.1 MeV) have been utilized to produce millimeter TR that is subsequently focussed with a parabolic mirror (Naumenko et al., 2008).

The theory of TR for an electron normally incident on a planar surface is covered by several textbooks (Jackson, 1999; Landau et al., 1984). We outline it here in a form that is suitable for obtaining plasmon launching probabilities in Sec. V.D. We consider a swift electron moving along the \( z \) axis and crossing the free surface of a metal occupying the \( z < 0 \) region (Fig. 9, right).

The electric field produced by the moving charge in each medium \( j \) (with \( j = 1 \) and 2 representing the materials above and below the interface, respectively) can be separated into the contribution of the external electron field for an infinite bulk material plus the field reflected at the surface,
\[
\mathbf{E} = \mathbf{E}_j^\text{ext} + \mathbf{E}_j^\text{ref}.
\]
In particular, the electric field inside an infinite medium \( j \) obtained from Eq. (2) after integrating the \( z \) component of \( \mathbf{q} \) reads
\[
\mathbf{E}_j^\text{ext}(r, \omega) = \int \frac{d^2 \mathbf{q}_i}{(2\pi)^2} \mathbf{E}_j^\text{ext}(\mathbf{q}_i, z, \omega) e^{i\mathbf{q}_i\cdot\mathbf{r}}
\]
with
\[
\mathbf{E}_j^\text{ext}(\mathbf{q}_i, z, \omega) = \frac{4\pi ie^{i\omega z/v} \mathbf{q} - \mathbf{v} k \hat{\epsilon}_j/c}{q^2 - k^2 \epsilon_j},
\]
where \( \mathbf{q} = (\mathbf{q}_i, \omega/v) \) and \( \hat{\phi}_q = \mathbf{z} \times \mathbf{q}_i \), so that the set \( \{\mathbf{q}_i, \hat{\phi}_q, \mathbf{z}\} \) forms a positively oriented 3D reference frame. Faraday’s law permits writing the corresponding magnetic field as
\[
\mathbf{H}_j^\text{ext}(\mathbf{q}_i, z, \omega) = \frac{4\pi ie^{i\omega z/v}}{c} \frac{\hat{\phi}_q}{q^2 - k^2 \epsilon_j}.
\]
It should be stressed that the bulk fields are evanescent away from the trajectory, unless \( \epsilon_j \) is a positive real number and \( v^2 \sqrt{\epsilon_j} > c^2 \), in which case CR can be produced.

The reflected component of the field finds its sources in the induced surface charges and currents. The charges can be in turn expressed in terms of the currents by using the continuity equation. After some algebra, we obtain
\[
\mathbf{E}_j^\text{ref}(\mathbf{q}_i, z, \omega) = \frac{4\pi ie^{i\omega z/v}}{c} \frac{\hat{\phi}_q}{q^2 - k^2 \epsilon_j}.
\]

\footnote{The electron velocity varies dramatically inside the metal due to both heavy encounters with target atoms and inelastic collisions, until it is eventually stopped. However, no relevant changes in the trajectory occur with significant probability for energetic electrons (\( \geq 50 \text{ keV} \)) within a distance of the order of the skin depth underneath the surface.}
\[-\frac{2\pi k}{q_{xz}} \times \left\{ \begin{array}{ll}
\frac{e^{iq_{z1}z}}{k_{xz}^2} \left[ h_1 - \frac{q_{v1}}{k_{xz}^2} (q_{z1}, q_{z1}) \right], & j = 1 \\
\frac{e^{-iq_{z2}z}}{k_{xz}^2} \left[ h_2 - \frac{q_{v2}}{k_{xz}^2} (q_{z2}, -q_{z2}) \right], & j = 2 \end{array} \right. \]

and

\[H^\text{ref}_j(q_{||}, z, \omega) = \frac{-2\pi}{q_{xz}} \times \left\{ \begin{array}{ll}
\frac{e^{iq_{z1}z}(q_{z1}) D \mu_j \text{sign}(z) \hat{\phi}_q}{z}, & j = 1 \\
\frac{e^{-iq_{z2}z}(q_{z2}, -q_{z2}) \times h_2}{z}, & j = 2 \end{array} \right. \]

where \(q_{xz} = \sqrt{k^2 - q_{||}^2}\). Here, \(h_j\) are the noted surface currents (defined on either side of the interface), which are determined from the continuity of the parallel components of \(E\) and \(H\). Finally, the reflected magnetic field reduces to

\[H^\text{ref}_j(q_{||}, z, \omega) = -\frac{2\pi}{q_{xz}} \times \left\{ \begin{array}{ll}
\frac{e^{iq_{z1}z}(q_{z1}) D \mu_j \text{sign}(z) \hat{\phi}_q}{z}, & j = 1 \\
\frac{e^{-iq_{z2}z}(q_{z2}, -q_{z2}) \times h_2}{z}, & j = 2 \end{array} \right. \]

where

\[
\mu_1 = \frac{-(\omega/v)\epsilon_2 + q_{z2}\epsilon_1}{q^2 - k^2\epsilon_1} - \frac{-(\omega/v)\epsilon_1 + q_{z2}\epsilon_1}{q^2 - k^2\epsilon_1},
\]

\[
\mu_2 = \frac{(\omega/v)\epsilon_2 + q_{z1}\epsilon_2}{q^2 - k^2\epsilon_2} - \frac{(\omega/v)\epsilon_1 + q_{z1}\epsilon_2}{q^2 - k^2\epsilon_2},
\]

and

\[
D = \frac{2\epsilon q_{||}}{q_{z1}\epsilon_2 + q_{z2}\epsilon_1}.
\]

Performing the integral over the azimuthal angle of \(q_{||}\), one finds

\[H^\text{ref}_j(r, \omega) = \left\{ \begin{array}{ll}
-\frac{iq_{||}}{q_{xz}} \frac{dq_{||}}{D} \mu_j e^{iq_{z1}z} |z| J_1(q_{||}R), & j = 1 \\
-\frac{iq_{||}}{q_{xz}} \frac{dq_{||}}{D} \mu_j e^{-iq_{z2}z} \times h_2, & j = 2 \end{array} \right. \]

where \(\hat{\phi}\) is the azimuthal unit vector of \(r\). The TR emission is obviously \(p\) polarized for normal incidence, with the magnetic field pointing along the azimuthal direction, parallel to the surface. This is in agreement with experimental observations, in which residual \(s\)-polarized emission is claimed to originate in surface roughness (Hatten-dorf, 1977).

The homogeneous field \(H^\text{ext}_j\) decays evanescently away from the electron trajectory (see Sec. II.A). Therefore, any TR emission must arise from \(H^\text{ref}_j\), which behaves as

\[H^\text{ref}_j(r, \omega) \rightarrow f_H(\theta, \omega) e^{ikr}/r \]

in the far field. Asymptotic analysis of Eq. (55) shows that

\[f_H(\theta, \omega) = ik\cos\theta D \mu_1 \hat{\phi}.\]

This expression must be evaluated at \(q_{||} = k\sin\theta\), where \(\theta\) is the emission angle with respect to the surface normal. Finally, proceeding in a way similar to Sec. IV.B, the TR probability is given by the angular integral

\[\Gamma_{\text{TR}}(\omega) = \frac{1}{2\pi} \int_0^{\pi/2} |f(\theta, \omega)|^2 d\theta, \]

performed over the upper hemisphere (vacuum).

Typical TR spectra are shown in Fig. 23(a) (broken curves) for 200 keV electrons incident on Ag and Au surfaces. The spectral emission intensity exhibits an approximately linear decrease with wavelength above the SP threshold. The integrated emission in the 600-1200 eV region yields \(10^{-3}\) photons per incident electron and shows a monotonic increase with probe energy [Fig. 23(b)].

D. Cherenkov radiation

The electric field set up by a swift electron in a homogeneous dielectric can exhibit far-field oscillatory behavior according to Eq. (5) whenever the charged probe moves faster than light in the medium under the condition (7). This is reflected in the fact that the argument
FIG. 23 (Color online) (a) Spectral dependence of the TR and SPP emission probabilities upon electron beam bombardment under normal incidence. A 200 keV electron impinges on a planar metal surface, giving rise to TR [dashed curves, calculated from Eq. (56)] and SPPs [solid curves, obtained from Eq. (71)]. The metal dielectric function has been taken from Johnson and Christy (1972). The probability is given per unit of light wavelength range. (b) Dependence of the TR and SPP emission probabilities on electron kinetic energy for two different free-space light wavelengths, $\lambda = 800$ nm and 1500 nm. (c) Measured (symbols) and simulated (dashed curve) SPP emission probability spectra for normally-incident 30 keV electrons. The SPP emission rate produced by an oscillating dipole placed at the surface is shown for comparison (solid curve), normalized to the spectral integral of TR and far-field dipole radiation. (d) Computed TR spectrum for 30 keV electrons normally impinging on a gold surface. (c) and (d) have been taken from Kuttge et al. (2009), to which the reader is referred for more details on the microscope setup, experimental methods, and theoretical analysis.

of the Bessel functions in Eq. (5) becomes imaginary under such conditions. Actually, these oscillations signal the emission of light, originally observed by Cherenkov (1934) and successfully explained soon after its discovery (Frank and Tamm, 1937; Tamm, 1939). The theory of the Cherenkov effect in homogeneous media was later generalized to accommodate frequency dispersion (Fermi, 1940; Tamm, 1939), and more sophisticated descriptions have been produced since then (Lalor and Wolf, 1971; Patro, 1982).

Cherenkov radiation has been extensively discussed (see Ginzburg, 1996, and references therein), but a particularly interesting demonstration was reported by Ohkuma et al. (1991), who observed the effect for bunched ultra-relativistic electrons moving faster than light in air.

The probability of CR emission in a homogeneous material was first obtained by Frank and Tamm (1937). It can be easily derived by integrating Eq. (20) over wavevector transfers, assuming real and local $\epsilon(\omega)$. One finds

$$\Gamma_{\text{Che}}(\omega) = \frac{e^2 L}{\hbar} \left[ \frac{1}{c^2} - \frac{1}{v^2 \epsilon(\omega)} \right],$$

(57)

where the dependence on emission frequency comes exclusively from the dielectric function. Equation (57) is a pillar in the design of Cherenkov detectors for high-energy physics, in which CR signals the passage of charged particles resulting from energetic subatomic collisions (Julley, 1958).

Non-trivial finite-trajectory effects have been successfully described using the field of Eq. (4) in structures of large dimensions compared to the emission wavelength, for which the contribution of reflection at the boundaries can be neglected (Čiljak et al., 2003; Ružička and Zrelova, 1993; Zrelova and Ružička, 1989). However, this approximation is not sufficient for small samples like the thin films theoretically addressed by Kröger (1968). Actually, the experiments reported by Yamamoto et al. (1996a) on mica and Si films show a complex evolution of the CR intensity with film thickness (particularly for backward emission) that is correctly explained by Kröger’s formula.

The Cherenkov effect has been theoretically investigated for a charge traversing planar interfaces (Kröger, 1968; Lucas and Kartheuser, 1970; Ter-Mikaelian, 1972; Von Festenberg and Kröger, 1968) or moving parallel to a surface (Schieber and Schächter, 1998), for layered structures (Bolton and Chen, 1995; Lastdrager et al., 2000), for an electron moving along the axis of a cylinder (Olsen and Kolbenstvedt, 1980), and for beams passing near a number of different geometries (García de Abajo et al., 2004). Other related effects have been explored, like CR due to bunched electrons moving along waveguides (Onishchenko et al., 2002), solitons producing CR in optical fibers (Akhmediev and Karlsson, 1995; Skryabin et al., 2003), a second narrower Cherenkov cone in strongly dispersive active media (Carusotto et al., 2001), THz SPPs emitted by browsing a light spot at superluminal velocities on a planar surface (Bakunov et al., 2005), and inverted Cherenkov cones in materials of negative refraction-index (Averkov and Yakovenko, 2005; Matloob and Ghaffari, 2004; Veselago, 1968).

Furthermore, exotic production of CR has been observed due to superluminal vortices in Josephson junctions (Goldobin et al., 1998), whereas subluminal CR emission has been reported assisted by phonons that supply the required extra momentum (Stevens et al., 2001).
FIG. 24 (Color online) Tutorial description of the Cherenkov effect in a photonic crystal. A charge moving in a homogeneous dielectric with velocity below the Cherenkov threshold $v < c/n$ (a) cannot transfer energy and momentum [shaded region in (b)] overlapping the light dispersion line [upper line in (b)]. However, a periodic modulation of the index of refraction (c) allows light to undergo umklapp processes [i.e., absorption or emission of lattice momentum, as shown in (d)], so that band folding over the first Brillouin zone of the resulting photonic crystal (e) produces mixing and gap openings in the photonic band structure (f). The charge can now couple to propagating optical modes.

1. Cherenkov effect in photonic crystals

An interesting situation is presented when an electron moves inside a photonic crystal, consisting of a periodic modulation of the index of refraction along certain(s) spatial direction(s). The resulting CR is then affected by the photonic band structure, which can considerably differ from the vacuum light cone, as schematically illustrated in Fig. 24 for a 1D crystal. The loss of translational invariance allows subluminal electrons to couple to photonic bands of the crystals, thus resulting in the emission of CR.

This effect has been theoretically analyzed by Luo et al. (2003) and Kremers et al. (2009) for electrons moving perpendicularly with respect to the pores of a 2D photonic crystal. These authors found a complex evolution of the electric field, which is molded by the photonic bands.

An experimental investigation of the interaction of fast electrons with a 2D photonic crystal has been reported (García de Abajo et al., 2003) and is illustrated in Fig. 25. The photonic crystal consisted of a porous alumina film, grown by electrochemical methods under the appropriate conditions to form a locally regular array of pores (Masuda and Fukuda, 1995). The precise control of electron optics in a STEM allows focusing the probes so that they pass along one of the pores of the crystal in a loaf configuration, with the beam positioned tens of nanometers away from the pore wall. The resulting loss spectra display prominent peaks around 7-9 eV, depending on the electron energy, which were attributed to radiative losses produced by coupling to photonic modes of the crystal. The spectral features are of course related to the photonic LDOS, as discussed in Sec. III.H, and they exhibit singularities similar to those explained by Van Hove (1953), but now arising from characteristic points in photonic rather than electronic bands. These results are indirect evidence of CR in photonic crystals. A direct experimental detection of CL in these systems has not been yet reported.

Incidentally, similar effects could be responsible for unusual spectral features in porous silicon at large beam-surface separation (Williams et al., 2000).

E. Diffraction radiation

We have seen that radiation can be produced when an electron crosses a dielectric interface or when it moves faster than light in a homogeneous medium. Another common source of light emission originates in the passage
beams in electron accelerators (Karataev et al., 2000). Thick curves: retarded EELS calculations using Eq. (33). Dotted curve: non-retarded EELS calculation for the larger sphere, obtained from Eq. (31). Thin solid curves: CL calculations using Eq. (58). The computed probability is given per eV of energy-loss range. The dielectric function of alumina is taken from Palik (1985). (31). Thin solid curves: CL calculations using Eq. (58). The EELS calculation for the larger sphere, obtained from Eq. (33). Dotted curve: non-retarded EELS calculations using Eq. (31). Thick curves: retarded EELS outside alumina spherical particles. Symbols: measured data taken from Abe et al. (2000). FIG. 26 (Color online) Losses in aloof trajectories due to diffraction radiation. The figure shows measured and simulated energy-loss spectra for 200 keV electrons passing just outside alumina spherical particles. Symbols: measured data taken from Abe et al. (2000). Thick curves: retarded EELS calculations using Eq. (33). Dotted curve: non-retarded EELS calculation for the larger sphere, obtained from Eq. (31). Thin solid curves: CL calculations using Eq. (58). The computed probability is given per eV of energy-loss range. The dielectric function of alumina is taken from Palik (1985).

The emission probability admits a closed-form expression similar to Eq. (33) (García de Abajo, 1999c):

\[
\Gamma_{\text{CL,sph}}(\omega) = \frac{e^2}{c^2} \sum_{l=1}^{\infty} \left[ \frac{\omega}{v_n} \right]^2 \sum_{m=-l}^{l} K_m^2 \left( \frac{\omega b}{v_n} \right) \left[ C_{lm}^M |t_{lM}^M|^2 + C_{lm}^E |t_{lE}^E|^2 \right].
\]

Equation (59) arises from the condition of constructive interference in the emission originating in contiguous spheres.

Successive experimental demonstrations of this effect were carried out using 30-300 keV electrons to produce light of wavelength lying in the \( \mu m \) region (Bachheimer, 1972; Salisbury, 1970). Also, relativistic beams have been employed for generating millimeter radiation (Blackmore et al., 2008; Doucas et al., 1992; Shibata et al., 1998), which turns out to be very intense along the forward direction (Woods et al., 1995). Furthermore, 20-40 keV dense electron beams coming from conventional electron microscopes and coupled to metallic gratings of periods in the sub-millimeter domain have been used to produce super-generation of far-infrared radiation (Temkin, 1998; Urata et al., 1998).\(^{18}\)

1. Smith-Purcell emission

The effect discovered by Smith and Purcell (1953) (S-P) is a well-known example of diffraction radiation. These authors recorded light emitted when an electron beam passes parallel and above a metallic grating. The direction and wavelength of emission followed a simple relation derived from a Huygens construction. More precisely,

\[
\Delta t = d \cos \theta/c = d/v + \lambda n/c,
\]

where \( n \) is an integer number, \( \lambda \) is the wavelength, and the rest of the parameters is defined in the following inset:

\[\text{electron} \quad \begin{array}{c}
\text{d} \\
\text{a} \\
\text{v} \\
\text{A}S\text{O}_3 \\
\text{light emission} \\
\theta \\
\text{c}\end{array}\]

\[\Delta t = d \cos \theta/c = d/v + \lambda n/c, \quad (59)\]

One can easily see from Eqs. (C1) and (C2) that the scattering matrices satisfy the relation \( |t_{lM}^M| = \text{Im} \left| t_{lM}^M \right| \) in spheres made of real dielectric function, implying that the loss probability equals the CL rate in that case, because the material is unable to absorb any energy. The CL rate calculated from Eq. (58) coincides with the loss probability in the region below 7 eV of Fig. 26 (thin and thick continuous curves, respectively), which further supports the dominant effect of the diffraction radiation mechanism in that part of the spectrum.

Incidentally, the EELS spectra of Fig. 26 show absolute maxima in the 15-20 eV range, corresponding to plasmons of the alumina spheres (for comparison, the bulk plasmon occurs at 25 eV and the planar SP at 21 eV).

18 Within linear response theory, valid for most situations considered in this review, the EELS and CL probabilities are proportional to the square of the particle charge, and they scale linearly with the beam intensity for moderate currents. However, bunching can produce nonlinear scaling when the electrons move closer to each other than the emission wavelength (Urata et al., 1998), so that they behave like a composite charge, the square of which is obviously larger than the sum of the squares. This leads to super-radiant Smith-Purcell emission.
FIG. 27 (Color online) Energy loss probability and Smith-Purcell radiation emission for an electron moving parallel to an infinite periodic string of aligned Al nanoparticles (see upper inset). The probability is normalized per sphere and per eV of energy-loss range. The loss probability for an individual sphere is also shown for comparison. Adapted from García de Abajo (1999a).

1974). It was also shown that an equivalent description results from considering the currents induced by the passage of the electron on the grating surface (Brownell et al., 1998). Moreover, a large deal of work has been devoted to studying the conditions upon which this effect could be utilized for efficient x-ray generation (Moran, 1992), for instance in free-electron lasers (FELs) (Li et al., 2007a,b; Liu et al., 2007).

The excellent spatial resolution of electron microscopes could be used to direct a beam parallel to an array of nanoparticles and produce S-P emission. This situation is considered in Fig. 27, which predicts intense supercontinuum emission in the UV domain. This type of submicron light emitters could find eventual application in integrated photonic devices. Interestingly, Fig. 27 shows the presence of an absolute threshold below which no emission is possible because Eq. (59) cannot be satisfied. In recent works, the S-P emission from photonic crystals has been extensively investigated (García de Abajo and Blanco, 2003; Ochiai and Ohtaka, 2004a,b, 2005; Pendry and Martín-Moreno, 1994) and compared with gratings (Ohtaka and Yamaguti, 2001; Yamaguti et al., 2002) using theoretical tools. Besides, experimental observations have been reported for millimeter waves emitted by electrons passing near photonic-crystal opals (Yamamoto et al., 2004). Finally, interesting effects due to finite size of the periodic structure along the direction of the beam have been noted (Ochiai and Ohtaka, 2006) and theoretically analyzed (Horiuchi et al., 2006).

F. Cathodoluminescence and plasmons

Plasmons in metallic nanoparticles have finite lifetime due to radiative decay, Landau damping (e-h pair creation), and other absorption mechanisms. The former is rather efficient when the particle has non-negligible size compared to the corresponding light wavelength, as we discuss below. Actually, the CL signal resulting from plasmon decay offers an excellent opportunity to spectrally and spatially resolve plasmons.

1. Plasmons in metallic films and gratings

An electron traversing a thin film can give rise to a peak in the CL emission at the bulk plasmon frequency $\omega_p$, as predicted by Ferrell (1958), who found that the emitted light was $p$ polarized (magnetic field parallel to the film) for normal incidence, and the emission intensity vanished at normal and grazing directions for a Drude metal. This prediction was soon confirmed by Steinmann (1960) and Brown et al. (1960) in Ag films, and later by Arakawa et al. (1964) in Ag and Mg films. A study by Ritchie and Eldridge (1962) generalized Ferrell’s results to arbitrary metals and $v \ll c$, while the comprehensive work of Cram and Arakawa (1967) showed excellent agreement between theoretical and experimental results in Ag films, compared on the same absolute-intensity scale. Additionally, the Ferrell mode was observed using optical excitation rather than electron beams (Brambring and Raether, 1965), and its presence was shown to affect the response properties of thin films near the bulk plasmon frequency (Kliewer and Fuchs, 1967). Subsequent TEM experiments managed to resolve the dispersion relation of the Ferrell mode in Al (Vincent and Silcox, 1973), basically following the transversal bulk plasmon curve of Fig. 8(b).

In contrast, SPPs in a thin film cannot directly couple to external light because their parallel momentum lies outside the light cone, according to Eq. (27), with $\epsilon_1$ and $\epsilon_2$ having opposite sign. However, SPPs can be made to radiate if they find obstacles in their propagation along an otherwise planar surface. In particular, gratings provide a source of parallel momentum that plasmons can use to couple to external light. The optical emission resulting from excitation of SPPs by electrons impinging on a grating was first observed by Teng and Stern (1967). These authors managed to correlate SPP-related peaks in the angular distribution of the emission with dips observed in the specular reflectivity of periodically corrugated surfaces.

In a remarkable development, Heitmann (1977) used CL to map the SP dispersion in sinusoidal Ag gratings of different height. The gratings assisted SPPs to couple to freely propagating light. His analysis of extrapolated results in the limit of shallow corrugation were in excellent accord with both the dispersion relation and the momentum-energy width predicted by Eq. (27). Further-
more, his results for stronger modulation of the grating profile resulted in dispersion relations further apart from the light line, so that the grating contributed to increase the binding of SPPs to the surface. A similar increase in surface-mode binding driven by periodic corrugations has been also observed in microwaves trapped by drilled metal films (Ulrich and Tacke, 1973). This behavior is accompanied by gaps in the boundaries of the Bragg zones, as first discussed by Ritchie et al. (1968) for $p$-polarized light incident on gratings, and later studied in 2D plasmon crystals (Kitson et al., 1996) capable of guiding plasmons through lines of defects (Bozhevolnyi et al., 2001).

Two recent outstanding experiments have independently supplemented Heitmann’s work by out-coupling electron-generated SPPs in planar surfaces using distant gratings (Bashevoy et al., 2006; van Wijngaarden et al., 2006), thus suggesting the possibility of using fast electrons as plasmon sources. We address this issue in more detail in Sec. V.D.

2. Plasmon mapping

Cathodoluminescence has been used to map plasmon excitations in scanning mode, by analogy to the EELS plasmon-mapping study presented in Fig. 16. The CL signal is generally much weaker than the loss intensity, which is problematic when good signal-to-noise ratio is needed to resolve neighboring spectral features. However, CL has the advantage that it can be performed in SEMs, which are more widely available than TEMs, and it works for thick samples, since there is no need to collect transmitted electrons.

Several examples of application of this technique to spatially resolve plasmons in nanostructures have been presented by Yamamoto et al. (2001a), gratings (Yamamoto et al., 2001b), holes in thin films (Degiron et al., 2004), and particle dimers and nanorods (Yamamoto et al., 2006b).

A characteristic example of these studies is shown in Fig. 28 for Ag particles. The spectra of Fig. 28(a) unveil the excitation of a broad dipole mode and a narrower quadrupole. The measurement is in excellent agreement with the theory of Eq. (58). By scanning the electron beam over the particle and collecting polarized photons emitted at the peak wavelength, one can form the image of the dipole plasmon shown in Fig. 28(b). The intensity of each pixel represents the photon rate measured when the electron is positioned at that spot of the sample. This plasmon map displays two characteristic maxima on either side of the particle along the direction of the polarizer. This study was carried out for particles of various sizes, the results of which are summarized in Fig. 28(c). The plasmon modes show a distinctive redshift with increasing particle size, in good agreement with the theory of Mie (1908), who provided the basis for understanding light scattering by spherical particles. The origin of these shifts is retardation, as we discuss in Sec. III.D.2, and they occur when the diameter of the particle is a sizable fraction of the wavelength. As an empirical rule, we can assert that retardation effects show up for radius $a \gtrsim \lambda/2\pi$.\footnote{The multipolar order $l$ intervenes in determining the relevant distance to which the wavelength has to be compared for assessing the relative importance of retardation. This must be of the order of the separation between nodes in the induced surface charge, which is $\sim a/l$, since the interaction between charges at larger distances is partially washed out by cancelation of contributions with different sign.}

These effects are accompanied by an increase in the width of the multipole plasmons with particle size, which arises as a result of stronger coupling to propagating light [see Fig. 28(d)] (Myroshnychenko et al., 2008b).

Figures 28(a)-(b) clearly demonstrate that aloof geometries (external trajectories) provide significant information on the particle plasmons. This idea is further illustrated in Fig. 29, which presents a quantitative comparison of EELS and CL probabilities for Au disks. The difference between these two probabilities corresponds to events in which the lost energy is absorbed by the particle rather than being re-emitted. These calculations have been carried out using the boundary element method described in Sec. III.E.3.

The CL spectra of Fig. 29 (dashed curves) show just a subset of the features observed in the loss spectra (solid curves), as an indication that some plasmon excitations are dark modes, in the sense that their main decay mechanism is via absorption by the particle rather than coupling to radiation (Chu et al., 2009). More precisely, the feature marked by an arrow in the loss spectrum of trajectory A is not found in the light cross section of randomly oriented disks (not shown), but it can be clearly resolved by EELS.

The spatial resolution of CL should be analogous to EELS for mapping plasmons, as shown in Fig. 14(b). This figure proves that aloof beams can yield similar emission intensities as penetrating electrons, in agreement with the images of Fig. 28(b). Finally, an additional thought on the aloof configuration: the incoherent CL signal, which we argued in Sec. IV.A.2 to be small compared to the coherent EIRE contribution reported in these figures, should be actually smaller when the electron does not traverse the metal and cannot create localized excitations, which are the main source of incoherent CL.

G. Ultrafast cathodoluminescence

In a recent development, the CL technique has evolved to allow picosecond resolution using a microscope in
FIG. 28 (Color online) Plasmon mapping of single metallic nanoparticles (adapted from Yamamoto et al., 2001a). (a) Measured and calculated CL spectra for a 122 nm Ag nanoparticle under grazing incidence, showing dipole ($l = 1$) and quadrupole ($l = 2$) plasmons. The dielectric function of Ag is taken from Johnson and Christy (1972). (b) Energy-filtered CL scan corresponding to the dipole-mode maximum of the spectra shown in (a). The collected light is passing through a polarizer with the orientation shown by the arrow. (c) Symbols: measured dispersion of multipole plasmons with particle size. Curves: Mie theory. (d) Plasmon width, obtained from Mie theory.

FIG. 29 (Color online) Comparison of EELS and CL intensities in a gold nanodisk. Two different electron beam trajectories are considered (see inset) separated 10 nm from the disk in both cases. The disk edges have been rounded with a curvature radius of 3 nm. These results have been obtained using the retarded BEM with the dielectric function of gold taken from Johnson and Christy (1972).

which the electron source consists of a photocathode being excited by external light pulses, so that the time-resolved CL emission can be assigned an excitation time origin determined for each of the pulses, taking into account the time of flight delay of the electron in its way to the target. This technique has been recently utilized to resolve charge-carrier dynamics in quantum dots and quantum wells with 50 nm spatial resolution and 10 ps time resolution (Merano et al., 2005; Sonderegger et al., 2006). Similarly, picosecond electron pulses have been used to develop ultrafast electron microscopy (Lobastov et al., 2005), so far applied just for simultaneous dynamical and structural characterization (Barwick et al., 2008; Grinolds et al., 2006). These studies open an avenue towards a whole range of interesting phenomena in the domain of the ultrafast nanoworld.

V. RELATED PHENOMENA AND SUGGESTED EXPERIMENTS

Electron microscopy is a rather versatile discipline that finds in the nanoworld its natural domain. We can certainly think of many new phenomena to be explored with focused electron beams, particularly when low-energy, valence excitations are involved. In this section, we briefly go through some of them.
A. Mechanical momentum transfer

A fast electron beam has the potential to act as a nanotool that manipulates molecules and nanoparticles by transferring linear and angular momentum. This should become an excellent complement to optical tweezers, currently employed to trap small particles ranging in size from nanometers to several microns (Meiners and Quake, 2000), and to study their mechanical behavior by exerting electromagnetic forces (Grier, 2003; Korda et al., 2002). Studies based on optical tweezers range from elastic properties of deformable living cells (Guck et al., 2000) to fluctuation forces in biomolecules (Meiners and Quake, 2000). Likewise, an electron microscope can modify and simultaneously image small structures with far greater resolution than what is achieved by optical imaging. A more sophisticated setup would involve in-vacuo optical tweezers integrated in an electron microscope for observation of trapped particles. This system is experimentally challenging because it requires to trap particles in vacuum, for which pioneering results on optical levitation were reported three decades ago by Ashkin and Dziedzic (1976). A solution that is perhaps more compatible with trapping in vacuum consists in relying on evanescent fields (e.g., localized SPs), as proposed by Novotny et al. (1997) and recently demonstrated by Righini et al. (2008).

Here, we estimate the magnitude of the momentum transfer imparted on small objects using the methods elaborated in previous sections (García de Abajo, 2004). The electromagnetic force is obtained by integrating Maxwell’s stress tensor over a surface surrounding the object (Jackson, 1999). Performing the time-integral of this force, we find

\[
\text{Momentum transfer} = \int F(t) \, dt = \int_0^\infty F(\omega) \, d\omega, \tag{60}
\]

where the spectral decomposition in the last identity is similar to the one used for the EELS probability in Sec. II.B.

For simplicity, we first limit our discussion to a small isotropic particle described by its electric polarizability \( \alpha(\omega) \). The force exerted by an external field \( \mathbf{E}(\mathbf{r}, \omega) \) acting on such particle is given by the expression (Gordon and Ashkin, 1980)

\[
\mathbf{F}(\omega) = \text{Re} \left\{ \alpha \sum_j E_j(\mathbf{r}, \omega) \nabla [E_j(\mathbf{r}, \omega)]^* \right\}, \tag{61}
\]

where the sum runs over Cartesian components and the external field supplied by the electron is given by Eq. (4). Using this in Eq. (61), one finds

\[
\mathbf{F}(\omega) = \frac{2\varepsilon_0 \omega^3}{v^4 \gamma^4} \left[ - \text{Re}\{\alpha\} f' \left( \frac{\omega b}{v \gamma} \right) \hat{x} \right. \tag{62}
\]

\[
\left. + 2 \gamma \text{Im}\{\alpha\} f \left( \frac{\omega b}{v \gamma} \right) \hat{z} \right],
\]

where the function \( f \) is the same as in Eq. (35) and the particle is located at a distance \( b \) from the electron beam.

Symmetry considerations lead to the conclusion that Rayleigh scattering by the particle produces a radiation pattern independent of the direction of \( \mathbf{v} \). This means that the overall transfer of momentum in the form of induced radiation is zero in the small-particle limit, so that Eq. (60) accounts for all momentum transfer emanating from the moving electron along the \( z \) axis, parallel to the trajectory. Then, the contribution of each \( \omega \) component to the electron energy-loss probability is \( v F_z(\omega) \). In fact, we find from Eqs. (62) and (34) the identity \( v F_z(\omega) = \hbar \omega \Gamma_{\text{EELS, dip}}(\omega) \): it is reassuring that the mechanical energy exchanged through momentum transfer equals the loss probability multiplied by the transferred energy for small targets.

The mechanical effect can be significant in nanoparticles, as shown in Fig. 30(b), which compares Eq. (62) (dashed curves) with the results obtained when all multipoles are included (solid curves). For instance, a typical STEM beam with a current of 0.1 nA passing 10 nm from the surface of a 40 nm alumina particle exerts a macroscopic attractive acceleration equal to 0.13 times the gravity. The transfer of angular momentum \( \Delta \mathcal{L} \) can be written

\[
\Delta \mathcal{L} = \int_0^\infty \tau(\omega) \, d\omega,
\]

where the torque \( \tau \) is also given by the integral of the Maxwell stress tensor (Jackson, 1999) over a surface surrounding the target. For the small particle considered

\[
\tau = \frac{\omega b}{v \gamma} \mathbf{E}(\mathbf{r}, \omega) \times \mathbf{E}(\mathbf{r}, \omega) - \frac{\omega b}{v \gamma} \mathbf{E}_{\text{scat}}(\mathbf{r}, \omega) \times \mathbf{E}_{\text{scat}}(\mathbf{r}, \omega),
\]

where \( \mathbf{E}_{\text{scat}}(\mathbf{r}, \omega) \) is the scattered field produced by the particle. However, for a small particle, the scattered field is negligible compared to the incident field, so that \( \tau \) reduces to

\[
\tau = \frac{\omega b}{v \gamma} \mathbf{E}(\mathbf{r}, \omega) \times \mathbf{E}(\mathbf{r}, \omega).
\]
above, one obtains
\[ \tau(\omega) = \frac{8\sqrt{2}\omega^2}{v^4\gamma^2} K_0 \left( \frac{w_b}{v\gamma} \right) K_1 \left( \frac{w_b}{v\gamma} \right) \left[ \text{Im}\{\alpha\} - \frac{2k^3}{3}|\alpha|^2 \right]. \]

Interestingly, the expression inside the square brackets of this equation vanishes for a non-dissipative particle.\(^{20}\) Equation (63) shows that the torque produced on a small particle drops exponentially with impact parameter [see Fig. 30(c)], but it is still sufficient to cause an astonishing 3.9 MHz/s angular acceleration when a 0.1 nA electron beam passes 20 nm away from a 40 nm alumina particle. The type of experiment needed to demonstrate this prediction should be attainable using the combination of optical tweezers and TEM suggested above. Perhaps the reader shares the curiosity of the author to see what happens to the particle as it spins increasingly faster until centrifugal forces compete with atomic binding.

B. Vicinage effects

At half the speed of light, electrons in a typical 100 keV, 0.1 nA STEM beam are separated by an average distance of 26 cm along the propagation direction and they impact the specimen with a time delay of 1.6 ns. The interaction among them is therefore negligible in vacuum, but a leading electron produces excitations in the sample that a second trailing projectile can pick up, whenever they last more than the delay time. This situation is discussed in Sec. V.B.1.

Another type of vicinage effect involves the electron self-interaction, taking place when its wave function extends over different regions of the sample. Current technology makes the observation of this effect feasible, and it should find direct application to measure nonlocal screened interactions, as we show in Sec. V.B.2.

1. Interaction between two electrons

This effect bears close relation to the amply discussed topic of swift charged-molecule dissociation after traversing thin foils, in which the interaction among the resulting ionic fragments affects their stopping (Gemmell et al., 1975) and the SE yield (Tomita et al., 2006). Likewise, mutual interaction between bunched electrons is at the heart of super-radiant Smith-Purcell emission (Li et al., 2006; Urata et al., 1998), as discussed in Sec. IV.E.1.

We can easily find from Eq. (8) the overall loss probability for two electrons moving a distance \(a\) apart from each other along the same straight-line trajectory. The ratio of this probability to that of a single electron is exactly given by
\[ 4\cos(\omega\Delta R/2v) \]
with independence of the nature of the sampled excitations. When the electrons move in close proximity as compared to \(v/\omega\), a two-fold increase in the combined loss probability is produced. An interesting situation occurs under the condition \(a = \pi v/\omega\): the energy released from the leading probe is picked up by the trailing one and the overall probability vanishes. This may find application to shadow intense energy-loss or light-emission features in order to make weaker spectral details more visible, in the spirit of the masking interferometry techniques utilized in astronomy (Ireland et al., 2008).

2. Electron self-interaction

The quantum nature of the electron probe can be exploited to retrieve nonlocal information on the dielectric response if we separate the incident beam into two or more coherent spots at the position of the sample.\(^{21}\) This is routinely done in TEM holography by inserting a prism in the electron optics, basically exploiting partial coherence of the energy-loss signal (Herring, 2008; Schattschneider and Werner, 2005; Verbeeck et al., 2008).

Inelastic holography has been performed both in the real image plane (Lichte and Freitag, 2000) and in the diffraction plane (Herring, 2005), although the current theoretical understanding of the obtained results suggests that further development of the technique is required in order to extract new information on the coherence of inelastic excitations (Verbeeck, 2006). This technique permits addressing profound questions on quantum mechanics, such as the Feynman thought experiment (Potapov et al., 2007).

Here, we discuss a similar scenario, consisting of an electron beam split into two coherent beams that are focused on identical neighboring spots. This situation is well described by Eq. (17) if one assumes that the lateral wave function of the incident electron has the form
\[ \psi_{i\perp}(R) = \frac{1}{\sqrt{2}} \left[ F(R - R_1) + F(R - R_2) e^{i\delta} \right], \]
where \( F \) gives the spot profile (e.g., a Gaussian of width \( \Delta R \)), \( R_j \) are the centers of the two separate spots (\( |R_1 - R_2| \gg \Delta R \)), and the phase \( \delta \approx m_\text{e}va/h \) accounts for the difference in path length between the two beams \( a \), which can be controlled through the electron optics of the microscope.

We now make two assumptions regarding the spot size: (1) the induced part of the screened interaction \( W^{\text{ind}} \) is considered to be a smooth function that does not vary significantly over the extension \( \Delta R \); and (2) the lateral

\(^{20}\) This is a consequence of the optical theorem for small particles, stating that \( \text{Im}\{-1/\alpha\} \geq 2k^3/3 \), with the equal sign only occurring for non-absorbing particles (van de Hulst, 1981).

\(^{21}\) Nonlocality in this paragraph refers to \( W(r, r', \omega) \) for \( r \neq r' \), and not to spatial dispersion in \( \epsilon \).
variation of the detected plane wave components across a distance $\Delta R$ is negligible, that is, $p_{f\perp}\Delta R \ll 1$ in Eq. (17). The first assumption limits our analysis to excitations extended over large regions as compared to the size of each beam spot. This is quite reasonable for a realistic value of $\Delta R \approx 2\text{Å}$, since most valence excitations including plasmons are extended over much larger distances (see, for example, Fig. 16). The second assumption defines the angular region of transmitted electrons in which we are interested (for instance, $p_{f\perp} \ll \Delta R \approx 0.5\text{Å}^{-1}$, or equivalently, collection angles $\ll 15\text{ mrad}$ for 100 keV electrons).

Inserting Eq. (64) into Eq. (17), one finds

$$
\frac{d\Gamma_{\text{NR}}^\text{EELS}(\omega)}{dp_{f\perp}} \propto \Gamma_{\text{NR}}^\text{EELS}(\mathbf{R}_1, \omega) + \Gamma_{\text{NR}}^\text{EELS}(\mathbf{R}_2, \omega) + \Gamma_{\text{INR}}(\mathbf{R}_1, \mathbf{R}_2, \omega),
$$

where

$$
\Gamma_{\text{INR}}(\mathbf{R}_1, \mathbf{R}_2, \omega) = \frac{2\nu^2}{\pi \hbar v^2} \int dz_1 dz_2 \text{Im}\{-W(\mathbf{r}_1, \mathbf{r}_2, \omega)\}
$$

$$
\times \cos \left[ \delta + p_{f\perp} \cdot (\mathbf{R}_1 - \mathbf{R}_2) - \frac{\omega}{v} (z_1 - z_2) \right]
$$

contains the interference between the two spots.

At variance with the EELS probability $\Gamma_{\text{NR}}^\text{EELS}(\mathbf{R}, \omega)$ of Eq. (10), obtained when the entire inelastic signal is collected, the quantity $d\Gamma_{\text{NR}}^\text{EELS}(\omega)/dp_{f\perp}$ offers direct access to the nonlocal dependence of the screened interaction, $\text{Im}\{W(\mathbf{r}_1, \mathbf{r}_2, \omega)\}$. This requires to have exquisite wavevector resolution, $\Delta p_{f\perp} \ll 1/|\mathbf{R}_1 - \mathbf{R}_2|$, in order to prevent cancelations of the interference signal produced by the oscillations of the cosine function in $\Gamma_{\text{INR}}$. Using 100 keV electron with an achievable angular resolution of $\varphi_{\text{out}} \approx 1\text{ mrad}$, one should be able to retrieve information for spots separated by a distance of $\approx 50 \text{ nm}$, which is sufficient to study excitations of interest in nanoparticles and nanostructured materials.

We illustrate this formalism by considering two coherent beams of impact parameters $\mathbf{R}_1 = b\mathbf{x}$ and $\mathbf{R}_2 = -b\mathbf{x}$ relative to the center of a sphere of radius $a$, described by Eq. (B6). We obtain

$$
\frac{d\Gamma_{\text{NR}}^\text{sph}(\omega)}{dp_{f\perp}} \propto \sum_{l=-1}^{1} \sum_{m=-l}^{l} (\omega/v)^{2l} \frac{1}{(l+m)!(l-m)!}
$$

$$
\times K_m^2 \left( \frac{\omega b}{v} \right) \text{Im} \left\{ \alpha_l^{\text{NR}} \right\} [1 + (-1)^m \cos(\delta + 2p_{f\perp}b)],
$$

where $\alpha_l^{\text{NR}}$ is defined in Eq. (32). Figure 31 represents the results obtained from Eq. (65) for a 200 keV electron split into two beams that are passing at a distance of 2 nm from a 20 nm gold particle, as shown in the inset. Interestingly, $d\Gamma_{\text{NR}}^\text{sph}/dp_{f\perp}$ is periodic in $p_{f\perp}$ with period $\pi/b$ and is dominated by a dipolar feature. Furthermore, the interference disappears under the condition $\delta + 2p_{f\perp}b = (2n + 1)\pi/2$ for integers $n$, leading to a loss probability proportional to Eq. (31).

![FIG. 31](image-url) (Color online) Loss probability according to Eq. (65) for two coherent beams passing close to opposite sides of a 20 nm gold sphere (see inset). The gold is described with the dielectric function of Johnson and Christy (1972). The electron energy is 200 keV.

C. Electron energy-gain spectroscopy

The width of the ZLP limits the resolution of EELS and masks low-energy features sitting in the elastic peak tail. These problems are partially solved as we show below if one is capable of reversing the process, so that energy gain rather loss is recorded. Electron acceleration events were actually reported over 40 years ago when Boersch et al. (1966b) observed absorption of thermal phonons by electrons transmitted through thin LiF films, as illustrated in Fig. 32. The gain signal was later shown to depended on temperature as

$$
\Gamma_{\text{EEGS}}(\omega) = n(\omega) \Gamma_{\text{EELS}}(\omega),
$$

where $n(\omega)$ is the EELS probability discussed in Sec. II and $n(\omega) = 1/\left[\exp(h\omega/k_B T) - 1\right]$ is the Bose-Einstein distribution function at temperature $T$ (García de Abajo and Kociak, 2008a). Actually, Boersch et al. (1966b) managed to suppress the measured $\Gamma_{\text{EEGS}}$ by cooling their samples. Unfortunately, thermal transfers are limited to modes of energies $\approx k_B T \approx 26 \text{ meV}$ at room temperature.

A more flexible strategy consists in exciting through external illumination only those modes that we want to probe, as already suggested by Howie (1999). The proposed technique is illustrated in Fig. 33 and can be termed electron energy-gain spectroscopy (EEGS) (García de Abajo and Kociak, 2008a), in which the electrons absorb photons from an external light source. However, the energy-momentum mismatch between freely propagating charges and photons prevents the linear coupling of fast electrons and light plane waves in vacuum (for example, an electron moving in vacuum cannot emit
In-vacuum optical acceleration of an electron moving along the main (Rapoport and Khattak, 1988). This is a 100-fold the external illumination, typically in the sub-meV domain. The energy gain process can occur near the Cherenkov cone, as proposed by More (1966) and observed fifteen years later by Edighoffer et al. (1981). The inverse Cherenkov effect can find application to accelerate charged particles (Kimura et al., 1995). In a related context, the observation of the inverse Smith-Purcell effect (Mizuno et al., 1975, 1987) demonstrated the coupling of free electrons to evanescent components of momentum produced by interaction of external light with a grating.

Here, we consider a more suitable scenario for spectroscopy. Light scattered from a nanostructure, and in particular evanescent components in its vicinity, possesses the necessary momentum to couple to the electron. Consequently, the energy-gain process can occur near the specimen and should yield information on its spectral response, directly mapping the evanescent light strength. In such setup, the full width of the gain peak must correspond to electrons that have absorbed one photon out of the external light, and therefore the energy resolution of this technique is only limited by the width of the external illumination, typically in the sub-meV domain (Rapoport and Khattak, 1988). This is a 100-fold increase in spectral resolution with respect to currently available EELS technology. [Notice that low-energy excitations (e.g., sub-meV modes) are difficult to resolve due to the ZLP background. Here we refer instead to the ability to resolve higher-energy excitation features (e.g., 1 eV) separated by minute energy differences (e.g., 1 eV versus 1.001 eV).]

In order to evaluate EEGS probabilities, we consider an energetic electron focused near the sample and describe it through the wave function of Eq. (16) (Sec. II.C.2). We adopt the non-recoil approximation and assume that the normalized transversal wave function $\psi_{\perp}(\mathbf{R})$ is focused within a narrow spot centered around $\mathbf{R} = \mathbf{R}_0$, so that only the motion along the beam direction $z$ needs to be considered. The electron wave function is perturbed by interaction with the external light field and picks up components of higher energy to become

$$
\psi_{\parallel}(z) = \frac{1}{L^{1/2}} e^{i p_i z} e^{-i \varepsilon_i t} + \delta \psi_{\parallel}(z) e^{-i \varepsilon_f t},
$$

where we now have $\varepsilon_f = \varepsilon_i + \omega$ and $\omega$ is the photon frequency. Using first-order perturbation theory, we find

$$
\delta \psi_{\parallel}(z) = \int dz' G_f(z - z') H_{\omega}(z') \psi_{\parallel}(z'),
$$

where $H_\omega(z) = (-e\hbar/m_\omega) E_z(\mathbf{R}_0, z) \partial_z$ is the energy-gain part of the electron-photon coupling Hamiltonian, $E_z$ is the $z$ component of the light electric field (parallel to the trajectory), $\partial_z$ denotes differentiation with respect

---

22 In-vacuum optical acceleration of an electron moving along the axis of a Bessel beam has been recently proposed (Zhao and Lü, 2008).
Direct application of Eq. (67) yields

\[ G_f(z - z') = \int \frac{dp}{2\pi} \frac{e^{ip(z - z')}}{(m_e \varepsilon_f / \hbar - p^2/2 + i0^+)} \]

\[ = -i\hbar^2 \frac{e^{ip(z - z')}}{m_e p_f} \]

is the non-relativistic Green function of the free electron propagating along the \( z \) direction with final energy \( \hbar \varepsilon_f \). The integral in Eq. (66) must be performed in the \( z \rightarrow \infty \) limit in order to ensure that the electron has already abandoned the region of interaction with the nanostructure. Then, \( \delta \psi_{||} \) can be expressed as a coefficient multiplying a plane wave function of energy \( \hbar \varepsilon_f \). Finally, the gain probability is given by the square of that coefficient as the dimensionless quantity

\[ P_{\text{EEGS}}(\omega) = \left( \frac{e}{\hbar \omega} \right)^2 \left| \int dz \, e^{-i\omega z/v} \frac{E_z(R_0, z)}{E_{\text{ext}}(\omega)} \right|^2, \quad (67) \]

where we have used the non-reciprocal approximation \( p_f - p_i \approx \omega/v \ll p_i \) for the electron moving with high velocity \( v = \hbar p_i / m_e \). If the incident field is a plane wave and the electron moves in vacuum, only scattered components contribute to the EEGS probability, because the momentum gained by the electron (\( \hbar \omega / v \)) is always larger than the free-space momentum of light. Therefore, we can substitute the induced field for the total field in Eq. (67).

Useful analytical formulas can be derived for small particles, in which the dipolar response is dominant, via their electric polarizability \( \alpha \). The EEGS probability is given by integration of the dipole scattered field,

\[ E_{\text{z}}^\text{ind} = \alpha \left( \omega^2 / c^2 \right) E_{\text{ext}} + (E_{\text{ext}} \cdot \nabla) \frac{\partial E_{\text{z}}}{\partial t} \exp(i\omega r/c / r). \]

Direct application of Eq. (67) yields

\[ P_{\text{EEGS, dip}}(\omega) = \left( \frac{2e \omega \alpha}{\hbar \omega^2 \gamma} \right)^2 \left( \frac{\omega b}{\gamma \gamma_e} \right) + \left( \frac{|E_{\text{ext}}|^2 \gamma^2 / 2 - \hbar^2 \omega / \gamma^2 \gamma_e^2} \right), \]

where \( \mathbf{b} \) is the impact parameter of the electron relative to the particle \( (\mathbf{b} \perp \hat{z}) \) and \( \mathbf{b} = \mathbf{b} / b \). Interestingly, for an external field that is perpendicularly oriented with respect to the electron trajectory, the scattered field picks up \( z \) components that couple to the electron, giving rise to the \( K^2 \) contribution in Eq. (68). This expression needs to be compared with the EELS probability for a small particle \( \Gamma_{\text{EELS, dip}} \) (Eq. (34)). For a lossless particle, the optical theorem \( \text{Im} \{\alpha\} = (2\omega^3/3c^3)|\alpha|^2 \) allows us to write \( \Gamma_{\text{EELS, dip}} = (P_{\text{EEGS, dip}}) \rho^2 f \), where \( \rho^2 = \omega^2 / 2c^3 \) is the free-space local density of photonic states, the factor \( f = 2\pi \hbar \omega \) corrects for the number of photons contained in a normalized external field \( E_{\text{ext}} \) (i.e., \( |E_{\text{ext}}| = 1 \)), and the average over orientations of this field has been performed. In other words, the EELS probability, or equivalently the CL probability \( (\Gamma_{\text{CL}}) \) in our lossless particle, coincides with the EEGS probability for an incoherent source containing one photon per photon state. For a dissipative particle, we have \( \Gamma_{\text{EELS}} > \Gamma_{\text{CL}} \), but the relation \( \Gamma_{\text{CL}} = (P_{\text{EEGS, dip}}) \rho^2 f \) still holds.

We are now ready to make a quantitative comparison of EEGS and EELS. Incidentally, \( \Gamma_{\text{EELS}} \) has units of inverse energy, whereas \( P_{\text{EEGS}} \) is dimensionless, so that the comparison requires integrating \( \Gamma_{\text{EELS}} \) over a finite energy-loss range \( \hbar \Delta \omega \). We have \( \Gamma_{\text{EEGS}}/ \Gamma_{\text{EELS}} \sim \pi |E_{\text{ext}}|^2 / (\hbar \Delta \omega \text{Im}(-1 / \alpha)) \). For instance, for a 100 nm gold nanoparticle illuminated with a laser of moderate intensity \( \sim 10^6 \text{W/cm}^2 \), tuned to the particle plasmon energy \( \hbar \omega = 2.5 \text{ eV} \), and taking \( \hbar \Delta \omega = 0.1 \text{ eV} \) (the plasmon width), we find \( P_{\text{EEGS}}/ \Gamma_{\text{EELS}} \approx 40 \). This must be compared with \( \Gamma_{\text{CL}}/ \Gamma_{\text{EELS}} < 1 \).

We conclude that EEGS should provide an excellent probe of local photonic properties, combining the spatial resolution of TEMs and the energy resolution of optical sources. The EEGS signal is free from bulk features that dominate the loss spectrum as soon as the electron trajectory intersects the sample. Control over the intensity of the pumped light adds an extra handle to facilitate the detection of the gain signal, in contrast to CL. Finally, EEGS is sensitive to light polarization and it can be applied to investigate nonlinear optical response with nanometer resolution.

### D. Surface plasmon launching

As microchip features continue to shrink, lithographically-patterned metal structures are becoming natural candidates to replace electronic microcircuits. The new devices are expected to operate at frequencies above the THz, rather than GHz, and to encode information in plasmon signals rather than electric currents. The field of plasmonics is actually experiencing a tremendous expansion in preparation for this emerging technology (Barnes et al., 2003) but is still facing the lack of suitable plasmon sources that provide the versatility demanded by highly integrated plasmonic circuits, which conventional techniques like frustrated total internal reflection and NSOM cannot satisfy.

Fast electrons are a promising alternative for generating plasmons with exquisite spatial control and relatively high efficiency. Plasmon launching by swift electrons was already demonstrated several decades ago by Teng and Stern (1967) and Heitmann (1977) in metallic gratings. These authors utilized the periodic corrugation to couple plasmons to propagating light, which was analyzed in energy and angle of emission.

These pioneering studies have been recently supplemented by monitoring SPP generation in metallic planar surfaces. The SPPs were eventually decoupled by means of a grating placed at some distance from the position in which they were generated (Bashevoy et al., 2006; van Wijngaarden et al., 2006), thus yielding information on...
SPP propagation lengths through the dependence of the resulting emission intensity on the distance between the beam and the grating, and conclusively demonstrating the excitation of SPPs by fast electrons. Subsequent studies have added more insight into the spectrally- and spatially-resolved propagation of SPPs (Bashevoy et al., 2007; Kuttge et al., 2009; Yamamoto and Suzuki, 2008) and the increase in plasmon losses produced by grain boundaries in polycrystalline gold compared to single-crystal gold (Kuttge et al., 2008). The excitation near the plasmon threshold $\omega_s$ has been also addressed (Peale et al., 2008).

Plasmons can be identified in Eq. (51) with the pole in the denominator of $D$ [Eq. (54)], which yields the dispersion relation given by Eq. (27). The $q_{\parallel}$ integral of Eq. (55) can be then performed in the $R \gg \lambda$ limit by only retaining the contribution of the $q_{\parallel} = q_{\parallel}^{\text{SP}}$ pole. This is the so-called plasmon-pole approximation (Ford and Weber, 1984), which enables us to approximate

$$D \approx \frac{C}{q_{\parallel} - q_{\parallel}^{\text{SP}}} \quad (69)$$

with

$$C = \frac{2ieq_{z_1}}{c} \frac{\epsilon_2}{\epsilon_1^2 - \epsilon_2^2}. \quad [\text{This comes from the Taylor expansion of } \epsilon_2 q_{z_1} + \epsilon_1 q_{z_2} \text{ around the } q_{\parallel} = q_{\parallel}^{\text{SP}} \text{ pole.}]$$

Now, we proceed by inserting the plasmon-pole expression back into the reflected components of the fields [Eq. (55)] and by approximating the integral over $q_{\parallel}$ using the $q_{\parallel} = q_{\parallel}^{\text{SP}}$ pole of Eq. (69). Finally, Eq. (55) reduces to

$$\mathbf{H}^{\text{ref}}(\mathbf{r}, \omega) \approx A_j e^{iq_{z_2}x} H_1^{(1)}(q_{\parallel}^{\text{SP}} R) \hat{\varphi}, \quad (70)$$

where

$$A_j = \pi q_{\parallel}^{\text{SP}} \text{sign}(z) C \mu_j \left( q_{\parallel}^{\text{SP}} \right),$$

and $\mu_j$ is defined in Eqs. (52) and (53). Here, $H_1^{(1)}$ is a Hankel function, dying off as $\approx \left( 2/\pi q_{\parallel}^{\text{SP}} R \right)^{1/2} \exp(iq_{\parallel}^{\text{SP}} R)$ at large distance $R$ from the electron trajectory (Abramowitz and Stegun, 1972). Equation (70) indicates that the induced plasmon field is a cylindrical wave of $m = 0$ symmetry and parallel wavevector $q_{\parallel}^{\text{SP}}$.

Figure 34(a) shows the real part of the reflected magnetic induction ($\mathbf{B}^{\text{ref}}(\mathbf{r}, \omega)$), calculated from Eqs. (54) and (55) as a function of $R$ and $\omega$ for $\mathbf{r} = (R, 0, 10\text{nm})$ along a silver surface bombarded by a 50 keV electron. Surface plasmons produce strong modulations of the field at energies below Ritchie’s non-retarded limit ($h\omega_s \approx 3.7$ eV for Ag). The period of these oscillations increases with decreasing $\omega$ in accordance with the $H_1^{(1)}$ behavior noted above. The surface field exhibits faster oscillations for $\omega > \omega_s$, which rapidly decay away from the trajectory.

The full spatial dependence of $\mathbf{B}^{\text{ref}}(\mathbf{r}, \omega)$ is represented in Fig. 34(b) for $h\omega = 1.5$ eV, clearly showing the excitation of a cylindrical SPP wave (see intensity maxima near the surface) and TR (oscillatory wave components evolving away from the surface). The TR region is dominated by oscillations of short wavelength ($2\pi/k$) compared to the SPP wavelength ($2\pi/Re\{q_{\parallel}^{\text{SP}}\}$), in consonance with the larger momentum of SPPs ($Re\{q_{\parallel}^{\text{SP}}\} > k$).

We are ultimately interested in determining how many plasmons are launched per incident electron, which can be calculated from the integral of the Poynting vector derived from Eq. (70) over a cylindrical surface centered around the electron trajectory. This leads to

$$\Gamma_{\text{SP}}(\omega) = \frac{1}{2\hbar k^2} \sum_{j=1,2} \frac{|A_j|^2}{\text{Im}\{q_{z_2}\}} \text{Re}\left\{ \frac{q_{\parallel}^{\text{SP}}}{\epsilon_j} \right\} \quad (71)$$

for the probability normalized per incident electron and per unit of plasmon frequency range.24

The prediction of Eq. (71) is represented in Fig. 23(a) (solid curves) for Au and Ag surfaces bombarded by 200 keV electrons, and normalized per unit of emission wavelength $\lambda$ [i.e., the plotted quantity is $(\omega/\lambda)\Gamma_{\text{SP}}(\omega)$]. Plasmon production is boosted near $\omega_s$ in both cases, and it actually surpasses the production of TR in that region. Roughly speaking, $\sim 10^{-3}$ gold plasmons are produced for each normally-incident electron within the 600-1200 nm spectral range, a figure that reduces to $\sim 1.7 \times 10^{-4}$ for 20 keV incident energy and monotonically grows with electron velocity until it levels up at a value of $\sim 2 \times 10^{-3}$ around 2 MeV [Fig. 23(b)]. This behavior is qualitatively consistent with the spatial extension of the external electron field $\sim \nu/\omega$ [see Eq. (5)], and one should expect maximum coupling to the plasmon field when that extension is of the order of the SPP wavelength. The dependence of the emission on primary electron-energy, plotted in Fig. 23(b), confirms this explanation.

Good agreement between the above theory and experiment has been recently obtained by Kuttge et al. (2009), who inferred the absolute SPP yield by first normalizing the measured CL signal to the calculated TR in a flat surface, and then analyzing the interference between SPPs decoupled by a grating and TR. The results are shown in Fig. 23(c).

23 The $1/\sqrt{R}$ dependence of the field is the signature of a surface mode, since the resulting Poynting vector scales as $1/R$, so that its integral over a circle of radius $R$ centered at the electron source stays independent of $R$, except for the exponential attenuation due to non-vanishing SPP absorption, $\exp(-2\text{Im}\{q_{\parallel}^{\text{SP}}\}/R)$.

24 It should be noted that the approximation of Eq. (69) is fully justified for $R \gg \lambda$, that is, in the limit that we have examined to derive the plasmon energy flux far away from the electron trajectory (García de Abajo, 2007).
Propagating plasmons can be generally excited with electron beams in any plasmon-supporting system, including metal films, nanowires, and particle arrays (Cai et al., 2009). An example of plasmon excitation in a linear particle array is discussed in Fig. 17, whereas experimental observation of electron coupling to plasmons in nanowires was obtained by Yamamoto et al. (2006b) and Vesseur et al. (2007).

The emission mechanism so far considered is coherent in the sense explained in Sec. IV.A.1. However, there are at least two other incoherent mechanisms of emission involving plasmons: (1) plasmon-assisted decay of localized excitations produced by the impinging electron close to its trajectory and (2) SEs, which can in turn couple to the plasmon, acting as primary electrons. These contributions are only effective within a distance of the order of the skin depth inside the metal. We can invoke Bethe’s theory of charge stopping (Bethe, 1930) to conclude that the energy deposited within that region scales as $\sim 1/E$ with the primary electron energy $E$. Consequently, one should observe relatively more incoherent emission at low electron energies and an increase in the degree of coherence with faster electrons.

E. Nonlocal effects in nanostructured metals

The interaction between closely spaced particles has recently attracted much interest because they produce large electric field enhancement in the gap region, which is beneficial for molecular sensing applications (Xu et al., 1999). Gap distances in the range of one to a few nanometers have actually been used to measure surface-enhanced Raman scattering (SERS) signals, but nonlocal effects have been largely neglected in this context until very recently (García de Abajo, 2008; Zuloaga et al., 2009). Spatially-resolved EELS is well suited to experimentally study these unexplored nonlocal effects in dimers, and therefore we discuss this problem in this section.

Spatial dispersion in the material response generally produces blue shifts and decreasing intensity of plasmon features. Since the early work of Fujimoto and Komaki (1968) on spheres, nonlocal effects in EELS have been theoretically addressed for several geometries in the non-retarded limit, and they have been predicted to show up for electron impact parameters in the nanometer range (Aizpurua and Rivacoba, 2008; Zabala and Echenique, 1990).

Nonlocal effects in the optical response of small metal spheres ($\lesssim 10$ nm) have been known for a long time [see, for example, Ouyang et al. (1992) for an EELS experimental study of small silver particles], and they are well described by a phenomenological model incorporating conduction-electron scattering at the particle boundary as a source of plasmon decoherence (Kreibig and Vollmer, 1995). This model consists in supplementing the damping rate $\eta$ in Eq. (A1) with a size-dependent term, $v_F/a$. However, the $v_F/a$ quantum-confinement model is inconsistent with the fact that an electron state trapped inside a finite cavity (mimicking a conduction electron in a nanoparticle) has infinite lifetime. Therefore, many-body effects rather than reflection at the boundaries seem to be the source of dephasing in small particles, facilitated by spatial dispersion, which is responsible for the breakdown of the local response approximation at distances.
below a few nanometers (García de Abajo, 2008).

The hydrodynamic model offers an alternative way of accounting for nonlocal effects, as discussed in Appendix A.2 for a general case and in Appendix C for spheres. The results of this model for single aluminum spheres are shown in Fig. 14(a) (dashed curves), compared with a local description (solid curves). As expected, nonlocal calculations are blue shifted with respect to the local theory.

For small gold dimers at optical frequencies, it is reasonable to describe the particles using Eq. (37), and to plug the resulting multipolar polarizability into the multiple-scattering formalism of Sec. III.E.4. We have used Eq. (21) for \( \epsilon(q, \omega) \). The calculated loss probability is represented in Fig. 35 for 20 nm particles (see García de Abajo, 2008, for more details on the model). The loss spectra show a dominant feature that corresponds to the lowest-order longitudinal dipole mode of the structure (i.e., this mode corresponds to the co-axial dipole-dipole mode characterized by a charge pattern 
\[ (+) \cdot (--) \] at large separations). Significant enhancement of the electric field is produced near the gap region at the plasmon mode frequency (Fig. 35, lower-left inset), so that the details of the dielectric response at small distances become critical. The local description of the sphere (dashed curves) is clearly insufficient at distances below 2 nm, since they produce redshift and increase of intensity as compared to the nonlocal results (solid curves). The response of the dimer is extremely sensitive to the gap distance, but electron microscopes are ideally suited to simultaneously determine this parameter and the plasmon energy.

VI. PROSPECTS FOR PLASMONICS

Browsing the figures of this review, we observe that the historical development of plasmons has been intimately correlated with advances in electron microscopy and spectroscopy. Since the early days of SPs, which Ritchie (1957) predicted to show up as characteristic energy losses in electrons traversing metallic thin foils (see Fig. 10), up until now that particle plasmons can be mapped with nanometer precision (see Fig. 16), these collective excitations have captured the imagination of electron microscopists and material scientists alike.

Plasmons have now taken off as excellent tools in ultrafast technology for processing information encoded as plasmonic signals (Barnes et al., 2003; Ozbay, 2006; Zia et al., 2006) and for applications in biology (Schuster et al., 1993) and biomedicine (Burda et al., 2005). In particular, localized plasmon resonances (LPRs) play a relevant role because they are typically accompanied by large enhancement of the electromagnetic field that is useful in molecular sensing (for example, using SERS (Xu et al., 1999)). It is of course important to have experien-

25 Localized plasmons can be confined down to a few nanometers in small particles, for which retardation effects are negligible, so that LPRs are safely described by Poisson’s equation. Actually, the minimum size of well-defined LPRs is only limited by broadening due to nonlocal effects (see Sec. V.E). Now, the question arises, are metals (in the optical sense) necessary to produce localized electromagnetic resonances? In other words, do we need to have negative \( \epsilon \) materials? A positive answer to this question comes from examining the eigenvalues \( A = 2\pi(\epsilon_2 + \epsilon_1)/(\epsilon_2 - \epsilon_1) \) that appear in the solution of Poisson’s equation, recast into Eq. (38) (i.e., the solution corresponding to vanishing external field): one finds that the eigenvalues are real and they must satisfy \( |A| < 2\pi \), which is only possible if \( \epsilon_1 \) and \( \epsilon_2 \) have opposite sign. An alternative explanation based upon physical intuition can be found in the formal equivalence between Poisson’s equation and the heat transport equation under stationary conditions, with \( \epsilon \) and \( \phi \) playing the role of the thermal conductivity and the temperature, respectively: real materials have positive thermal conductivity, which guarantees that the second law of thermodynamics if fulfilled (heat does flow towards regions of lower temperature); but we do not know that heat cannot be spatially confined...
tal tools that permit us to visualize LPRs in patterned metallic surfaces and nanoparticles. One of these tools is NSOM (see Kim and Song, 2007, and references therein), which has been successfully applied to map plasmons (Bozhevolnyi et al., 2006; Drezet et al., 2007), although it is limited to around several tens of nanometers in spatial resolution. In this context, absorption and scattering spectroscopy has been extensively used to characterize the optical properties of metallic nanoparticles in colloids (Burda et al., 2005), but it requires narrow distributions of particle size and morphology, and it can only access the average properties of particle assemblies. Similarly, dark-field microscopy (DFM) is becoming a popular technique in the colloidal chemistry community, allowing localized plasmons to be studied in individual nanoparticles (Mock et al., 2002). However, unlike EELS, the DFM technique cannot provide details on the local distribution of these modes.

Spatially-resolved EELS and CL offer alternative methods to characterize plasmons with unmatched resolution. Actually, no other available technique has achieved the degree of detail obtained by the plasmon studies that we review in Figs. 16, 28, and 36. Furthermore, the theoretical tools that are necessary for interpreting EELS and CL spectral images have already reached a mature status that allows addressing arbitrarily complex geometries (see Sec. III.E).

The plasmon energies and widths obtained from EELS and optical extinction are in excellent agreement [Fig. 36]. In particular, the spectra of dumbbells that we show in Fig. 36(c) provide similar information in both techniques. However, EELS adds the spatial resolution that allows mapping plasmons in real space [see Figs. 16 and 36(d)]. The resulting EELS images compare reasonably well with the near-field enhancement that is expected upon conventional optical illumination [Fig. 36(e)]. Moreover, EELS can access plasmon modes that are not resolvable with light (Chu et al., 2009). For example, multipoles of order \( l > 1 \) in small metal spheres [Fig. 36(f)].

Like far-field optical spectroscopy, the EELS and CL techniques benefit from the use of relatively non-invasive probes (the electrons) that produce negligible changes in the optical properties of what is being observed. In contrast, the NSOM tip exerts a strong perturbation on the sample. The spatial resolution of EELS and CL is now pushed down to the sub-Å level, but even a modest SEM setup equipped with CL can focus an electron beam down to \(< 2 \) nm spots. The photon-energy resolution of CL is similar to that of other optical techniques, although the signal-to-noise ratio becomes annoying, particularly when the electron does not penetrate the specimen but simply interacts through electromagnetic fields at a short distance from it (aloof geometry). Regarding EELS, the 0.1 eV resolution achieved by forefront commercial STEMs is already sufficient for most plasmonic structures, and this situation should improve in coming years with further advances in electron monochromators and spectrometers.

Finally, it would be unfair to conclude this section without a summary of the impressive amount of information that has been gathered on both localized and extended plasmons using electron microscopes. We cannot review the vast literature here, but we present a succinct overview with some relevant references in Table I.

VII. CONCLUSION

Overall, valence excitations, probed by electron microscopes, have been a continuous source of information on electronic structure and optical response of materials on the nanometer scale over the last six decades. The energetic electrons employed in these microscopes permit obtaining structural and spectroscopic details of the investigated samples with sub-nanometer resolution, and they can be regarded as relatively contactless probes compared to other techniques such as STM of NSOM, particularly when the aloof configuration is employed, in which the electron beam passes outside the targeted materials and interacts with them via electromagnetic coupling. Optical excitations are the most intense in the energy-loss spectra, so that they can be collected using low beam-currents, and therefore, sample damage is reduced to small levels, even for penetrating trajectories. However, electron microscopes are versatile instruments in which the beam current can reach doses that cause substantial changes in the specimen, including reshaping of the morphology of metallic nanoparticles (Batson, 2008).

In the context of electron microscopy, CL constitutes an inexpensive addition to electron microscopes that can be very valuable when combined with appropriate theoretical tools, the versatility of which we have tried to illustrate in this review. Electron microscopes in scanning mode of operation, combined with either EELS or CL analysis, are suitable tools for studying the optical response of nanostructures, and no other existing technique can match them in combined space-energy resolution. Furthermore, these instruments should facilitate the eventual application of swift electrons to light and plasmon sources for nanoscale devices. Finally, there is a huge background of both experimental and theoretical expertise that constitutes a valuable guide for future research making use of electron microscopes in nanophotonics studies.
FIG. 36 (Color online) Comparison between EELS, CL, and optical extinction spectroscopy. (a) Electron energy-loss spectrum (EELS) of the dumbbell gold nanoparticle shown in the dark-field TEM-image inset. (b) Plasmon maps obtained from the intensity of electrons that have lost an energy corresponding to the dipolar longitudinal mode (1.7 eV) and the dipolar transversal mode (2.4 eV). (c) Calculated EELS probability (left scale) and extinction cross section (right scale) of a nanoparticle mimicking the dumbbell of (a). The cross section is obtained for external electric field orientations both parallel and perpendicular to the particle axis, in order to excite longitudinal (l) and transversal (t) plasmons, respectively. The EELS spectra correspond to positions of the electron beam as shown in (d) by white circles. The longitudinal mode is red-shifted in the experiment, as compared to the calculation, presumably due to the effect of a mica substrate, which is not accounted for here. (d) Calculated EELS plasmon maps, with the particle contour shown as a shaded area. Notice that the experimental images of (b) exhibit depletion of electron intensity for trajectories passing through the particle, due to elastic scattering of the electrons with gold atoms. (e) Near-field plots for longitudinal and transversal polarization of the incident electric field under external optical illumination. (f) Comparison of EELS and CL probabilities (solid and dashed curves, respectively) with the optical extinction cross section $\sigma_{ext}$ (dash-dotted curve) for a metal sphere described by the dielectric function of Eq. (A1). The damping rate is $\eta = 0.04 \omega_p$ (similar to Al), the sphere radius is a tenth of the free-space wavelength at frequency $\omega_p$, and the impact parameter is $b = 1.1 a$. The experimental data of (a) and (b) are taken from Bosman et al. (2007). The calculations of (c)-(e) are adapted from Myroshnychenko et al. (2008b).

Acknowledgments

It is a pleasure to express my gratitude to Archie Howie for countless suggestions, and to Benito Hombra-dos Madrid and Carmen Martín Nieto for their generous technical assistance. I would also like to acknowledge many enjoyable and stimulating discussions with Harry A. Atwater, Cheng-Hsuan Chen, Ming-Wen Chu, Christian Collieux, Pedro M. Echenique, Archie Howie, Mathieu Kociak, Martin Kuttge, Viktor Myroshnychenko, Albert Polman, Alberto Rivacoba, Peter Schattschneider, Naoki Yamamoto, Nerea Zabala, and Nikolay Zheludev. This work was supported in part by the Spanish MICyT (MAT2007-66050 and Consolider NanoLight.es) and by the EU (NMP4-2006-016881 "SPANS" and NMP4-SL-2008-213609 "ENSEMBLE").

APPENDIX A: Dielectric response of homogeneous media

The local response approximation yields excellent results when comparing theory and experiment both in the context of nanophotonics and in numerous EELS and CL studies that focus on the role of sample geometry. The different materials forming a structure are then described by frequency-dependent dielectric functions, usually taken from tabulated optical data (see, for example, Palik, 1985).

Several metals, including some alkali elements and aluminum, are remarkably well described by a free-electron gas model, in which the local dielectric function takes the
The dependence of $\omega_p$ on $n_0$ has been experimentally monitored by Imbusch and Niedrig (1970) using EELS and changing $n_0$ through thermal expansion in the 3 – 295 K range.

Logical relaxation rate that accounts for electromagnetic damping in the material (e.g., $\hbar \eta = 0.6 \text{ eV in Al}$). This model gives $\hbar \omega_p = 15.8 \text{ eV}$ for Al, in good agreement with the value of 15 eV obtained from EELS measurements (see Fig. 8 and Table II).

Some ionic materials, and alkali-halide crystals in particular, are better described by assuming localized ionic oscillation modes, which result in the dielectric function (Ashcroft and Mermin, 1976)

$$\epsilon(\omega) = \epsilon_{\infty} \left[ 1 - \frac{\omega_{\text{tr}}^2}{\omega(\omega + i\eta) - \omega_{\text{lon}}^2} \right], \quad (A3)$$

where $\omega_{\text{tr}}$ is the frequency of transverse vibrations and $\omega_{\text{lon}}$ signals a plasmon-like longitudinal mode characterized by $\epsilon \approx 0$. Retardation produces dispersion of these modes for momentum transfers comparable to $\hbar k$, as shown in Fig. 5.

Neglecting the influence of the atomic lattice, the ratio

 表 I Historical overview of experimental surface plasmon studies making use of electron beams.

| Structure                        | Technique | $\hbar \omega$ (eV) | Reference                          | Comments |
|---------------------------------|-----------|----------------------|-----------------------------------|----------|
| SP Al planar surfaces           | REELS     | 5 – 100              | Powell and Swan (1959)             | first observation of planar SPs |
| SPP Al grating                  | CL        | 2.3                  | Teng and Stern (1967)             | SPPs signature in CL |
| LPR Al nanoparticles            | CL        | 5 – 100              | Fujimoto et al. (1967)            | Mie modes |
| SPP Al films                    | EELS      | 3 – 10               | Pettit et al. (1975)              | SPPs dispersion relation |
| SP Al and In surfaces           | EELS      | 10 – 12              | Krane and Raether (1976)          | large-momentum dispersion |
| SPP Ag shallow grating          | CL        | 2 – 3                | Heitmann (1977)                   | SPP dispersion relation |
| LPR Al nanospheres              | EELS      | 1 – 20               | Batson (1980)                     |          |
| LPR Al nanospheres on AlO$_3$   | EELS      | 1 – 40               | Wang and Cowley (1987a)           | plasmon maps |
| LPR Al nanospheres on Al        | EELS      | 1 – 40               | Wang and Cowley (1987b)           | plasmon maps |
| LPR Ag nanospheres              | EELS      | 2 – 35               | Ouyang et al. (1992)              | nonlocal effects |
| LPR Si nanospheres              | EELS      | 2 – 35               | Ugarit et al. (1992)              | plasmon maps |
| LPR Al nanospheres              | EELS      | 2 – 40               | Stöckli (1997)                    |          |
| SP Al$_2$O$_3$ nanospheres      | EELS      | 2 – 30               | Abe et al. (2000)                 | diffraction radiation losses |
| SPP Ag gratings                 | CL        | 1 – 4                | Yamamoto et al. (2001b)           | plasmon standing waves |
| LPR Ag nanospheres              | CL        | 2 – 4                | Yamamoto et al. (2001a)           | plasmon maps of Mie modes |
| LPR WS$_3$ nanotubes            | EELS      | 3 – 30               | Kociak et al. (2001)              |          |
| LPR C nanotubes                 | EELS      | 3 – 25               | Stéphan et al. (2002)             | $\pi$ and $\sigma$ plasmons |
| LPR Ag rough surfaces           | PEEM      | 3                    | Kubo et al. (2005)                | femtosecond dynamics |
| SPP Au planar surfaces          | CL        | 1 – 3                | Baschevoy et al. (2006)           | SPP propagation |
| LPR Ag sphere dimer             | CL        | 2 – 4                | Yamamoto et al. (2006b)           | plasmon maps |
| LPR Ag nanowires                | CL        | 1 – 4                | Yamamoto et al. (2006b)           | plasmon maps |
| SPP Au atom wire array          | LEEM      | 1 – 3                | Nagao et al. (2006)               | 1D plasmon dispersion |
| LPR Ag triangular nanoprisms    | EELS      | 1 – 4                | Nelayah et al. (2007a)            | plasmon maps |
| LPR Ag nanowires                | CL        | 1 – 4                | Vesseur et al. (2007)             | plasmon maps |
| SPP Au planar surfaces          | PEEM      | 3                    | Baschevoy et al. (2007)           | hyper-spectral imaging |
| LPR Au annular grating          | CL        | 1 – 3                | Kubo et al. (2007)                | femtosecond dynamics |
| SP HfO$_2$ in stacks            | EELS      | 2 – 30               | Hofmann et al. (2007)             | plasmon maps |
| SP Si/SiO$_2$/Si stacks         | EELS      | 2 – 30               | Couillard et al. (2007)           |          |
| SPP Au planar surfaces          | CL        | 1 – 3                | Couillard et al. (2008)           |          |
| LPR Au ridge                    | CL        | 1 – 3                | Kuttge et al. (2008)              | SPP propagation |
| LPR Au nanorods, nanospheres    | EELS      | 1 – 4                | Peale et al. (2008)               |          |
| ellipsoids, dimers, and         |           |                      | Yamamoto and Suzuki (2008)        | plasmon maps |
| touching dimers                 |           |                      | Vesseur et al. (2008)             |          |
|                                 |           |                      | Bosman et al. (2007)              |          |
|                                 |           |                      | N’Gom et al. (2008)               |          |
|                                 |           |                      | Chu et al. (2009)                 |          |
|                                 |           |                      | Schaffer et al. (2009)            |          |

Equation (A1) describes the correct way of incorporating damping, via $\omega(\omega + i\eta)$ rather than $(\omega + i\eta)^2$. The latter is inconsistent with conservation of the number of electrons in the metal (Mermin, 1970).

The dependence of $\omega_p$ on $n_0$ has been experimentally monitored by Imbusch and Niedrig (1970) using EELS and changing $n_0$ through thermal expansion in the 3 – 295 K range.
between components of the external potential and the total potential (resolved in momentum and frequency) defines in general a dispersive dielectric function $\epsilon(q, \omega)$. Spatial dispersion becomes important for wavevector transfers $q$ of the order of $\omega/v_F$, where the Fermi velocity $v_F$ is typically two or three orders of magnitude smaller than the speed of light $c$. Consequently, the dependence on $q$ is typically important for $q \gg k_c$, away from the region showing retardation [see Fig. 8(b)].

1. Lindhard and Mermin dielectric functions

Spatial dispersion effects in good metals such as aluminum and silver can be accounted for by assimilating their conduction band to a gas of non-interacting free electrons (Ashcroft and Mermin, 1976). An analytical formula for $\epsilon(q, \omega)$ in the free-electron gas was derived by Lindhard (1954). The Lindhard formula, which predicts a continuum of $e$-$h$ pair excitations and collective plasmon modes as illustrated in Fig. 8(a), is obtained by perturbing the plane-wave representing each electron within the Fermi sphere by means of an external potential rather than the external potential in what has then assumed to be perturbed by the self-consistent total potential. Consequently, the dielectric function $\epsilon(q, \omega)$ is obtained by Lindhard (1954). The Lindhard formula, which predicts a continuum of $e$-$h$ pair excitations and collective plasmon modes as illustrated in Fig. 8(a), is obtained by perturbing the plane-wave representing each electron within the Fermi sphere by means of an external potential rather than the external potential in what has then assumed to be perturbed by the self-consistent total potential.

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The system is then assumed to be perturbed by the self-consistent total potential rather than the external potential in what has been termed the random-phase approximation (RPA). This leads to (Ashcroft and Mermin, 1976)

$$\epsilon(q, \omega) = \left[1 + \frac{4\pi}{q^2} \phi_{\text{ext}}\right]^{-1} = 1 + \frac{\epsilon^2}{\pi^2 \hbar q^2} \int q^2 k / h k \cdot q / m_e + \omega,$$

where the integral extends over 3D electron wavevectors $k, f_k$ is the Fermi distribution (approximately 1 for $k \lesssim k_F$ and 0 otherwise at room temperature), and $k_F = m_e v_F / \hbar$ is the Fermi wavevector. This integral admits the closed-form expression (Lindhard, 1954)

$$\epsilon^L(q, \omega) = 1 + \frac{2m_e e^2 k_F}{\pi \hbar^2 q^2} [1 + R(x, y) + R(x, -y)],$$

where $x = q/k_F$, $y = \omega/\epsilon_F$, $\hbar \epsilon_F = \hbar^2 k_F^2 / 2m_e$ is the Fermi energy, and

$$R(x, y) = \frac{1}{2x} \left[1 - \left(\frac{x^2 + y}{2x}\right)^2\right] \ln \left(\frac{x^2 + 2x + y}{x^2 - 2x + y}\right).$$

Here, $\omega$ is assumed to have a positive imaginary part. If Im{\omega} is infinitesimally small then Im{\epsilon^L(q, \omega)} takes nonzero values only for $x(x - 2) \leq y \leq x(x + 2)$ (i.e., in the so-called $e$-$h$ region), which corresponds to the shaded area of Fig. 8(a).

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A naive way of introducing a damping rate $\eta$ in the electron-gas motion consists in adding a finite imaginary part to $\omega$, but this results in a net loss of the number of electrons with time. The Lindhard formula was extended by Mermin (1970) to include finite damping while maintaining the number of electrons constant. He found the expression

$$\epsilon^M(q, \omega) = \frac{1 + (\omega + i\eta) \left[\epsilon^L(q, \omega + i\eta) - 1\right]}{\omega + i\eta \left[\epsilon^L(q, \omega + i\eta) - 1\right] / \epsilon^L(q, 0) - 1!},$$

where

$$\epsilon^L(q, 0) = 1 + \frac{2me^2 k_F}{\pi \hbar^2 q^2} \left[1 + \frac{4 - x^2}{4x} \ln \left|\frac{x + 2}{x - 2}\right|\right]$$

is the static limit of $\epsilon^L$. It is typically important for $q_c$, away from the region showing retardation [see Fig. 8(b)].

2. Hydrodynamic model

A simpler approach, which can be extended to nonlocal structured metals, consists in describing the conduction band as a classical electron plasma (Bloch, 1933). We generalize this approach here to incorporate retardation effects. The force equation for the hydrodynamic flow takes the form

$$\frac{\partial}{\partial t} \rho + \nabla \cdot \vec{j}_{\text{ind}} = \frac{(e^2/m_e) n_0 \vec{E}}{\hbar} + \frac{(\hbar/m_e)^2 [\beta^2 - (1/4) \nabla^2] \nabla \rho_{\text{ind}}}{\hbar},$$

where $\rho_{\text{ind}}$ and $\vec{j}_{\text{ind}}$ are the charge density and current density perturbations induced in the electron gas, $\vec{E}$ is the self-consistent electric field, $n_0$ is the unperturbed electron density, and the hydrodynamic pressure is related to the Fermi wavevector $k_F$ through the coefficient $\beta = (3/5)^{1/2} k_F$ (Ritchie, 1957). We have supplemented this equation with the cutoff term $-(\hbar/m_e)^2 (1/4) \nabla^2 \nabla \rho_{\text{ind}}$, which accounts for $e$-$h$ pair excitations in an effective way. The response of the system is then obtained by combining Eq. (A5) with Maxwell’s equations for the total charge and current ($\rho_{\text{ext}} + \rho_{\text{ind}}$ and $\vec{j}_{\text{ext}} + \vec{j}_{\text{ind}}$, respectively) in the presence of the ionic background, the response of which is accounted for through a permittivity $\epsilon_0$. This formalism is a direct generalization of the common non-retarded approach that is based upon Maxwell’s equations.

28 The linearized Bloch hydrodynamic equations constitute a suitable approximation when $\omega$ is much smaller than $\omega_0$. The mean free path of $e$-$h$ pairs is much smaller than the mean free path of $e$-$e$ pairs.

29 Interband transitions in aluminum do not contribute significantly to $\rho_{\text{ind}}$, but $\epsilon_0 = 1$ works well for that material, indicating that they are not strongly polarized near the bulk plasmon frequency (Raether, 1980). In contrast, interband transitions play a dominant role in noble metals such as Ag, Au, and Cu (Johnson and Christy, 1972), where $\epsilon_0$ differs considerably from 1 (see Table II).

30 Notice that the continuity equation $\nabla \cdot \vec{j} = i\omega \rho$ is implied by Maxwell’s equations.
upon Poisson’s equation and expresses the current density through a velocity potential $\xi$ as $j^{\text{ind}} = \eta_0 \nabla \xi$ (see, for instance, Ritchie, 1957, for application to planar surfaces, Fujimoto and Komaki, 1968, Crowell and Ritchie, 1968, and Barberán and Bausells, 1985, for spheres, and Wang and Cowley, 1987c, for supported particles). The customary boundary conditions of the electromagnetic field at the metal surface must be now supplemented by the condition that the normal component of $j^{\text{ind}}$ is zero. The results of this formalism for planar surfaces and spheres are presented in Appendices B.1 and C, respectively.

In homogeneous media, it is convenient to project the hydrodynamic and Maxwell equations in the space of wavevector $q$ and frequency $\omega$, from which we can work out the longitudinal and transversal dielectric functions $\epsilon_{\text{lon}}$ and $\epsilon_{\text{tr}}$ using the definition of electric displacement

$$D = \epsilon_{\text{lon}}E_{\text{lon}} + \epsilon_{\text{tr}}E_{\text{tr}}$$

in terms of electric field components parallel and perpendicular to $q$, respectively. We obtain

$$\epsilon_{\text{lon}}(q, \omega) = \epsilon_b + \frac{\omega_p^2}{(\hbar/m_e)^2 [\beta^2 q^2 + q^3/4]} - \omega(\omega + i\eta)$$

and

$$\epsilon_{\text{tr}}(q, \omega) = \epsilon_b - \frac{\omega_p^2}{\omega(\omega + i\eta)}. \quad (A6)$$

Eq. (A6) was derived by Ritchie (1957) from an expansion of Lindhard’s function for large $\omega$. The dispersion relation of longitudinal and transversal bulk plasmons is represented in Fig. 8(b) for aluminum.

**APPENDIX B: Non-retarded screened interaction in simple geometries**

The electrostatic screened interaction $W(r, r', \omega)$ is defined as the amplitude of the potential created at $r$ by a charge of magnitude $\exp(-i\omega t)$ placed at $r'$. For materials described by a local, frequency-dependent dielectric function $\epsilon(r, \omega)$, Poisson’s equation leads to

$$\nabla \cdot \epsilon(r, \omega) \nabla W(r, r', \omega) = -4\pi \delta(r - r'). \quad (B1)$$

In free space, one has $W^0(r - r') = |r - r'|^{-1}$. While Eq. (B1) is valid for arbitrary, smooth spatial dependence of the permittivity, most situations encountered in practice are reliably modeled by sharp boundaries. We can then obtain $W$ from the Coulomb interaction in each homogeneous region after imposing the continuity of both the potential and the normal displacement at the interfaces. Next, we offer analytical expressions for $W$ in the simple cases of planes, cylinders, and spheres.

1. **Planar surface**

A planar interface at $z = 0$ is best described by expressing $W$ in the space of parallel wavevector $q_{||}$. In particular, the bare Coulomb interaction can be written

$$\frac{1}{|r - r'|} = \int \frac{d^2q_{||}}{(2\pi)^2} e^{iq_{||}(R - R')} W^0(q_{||}, z, z'), \quad (B2)$$

where $r = (R, z)$ and

$$W^0(q_{||}, z, z') = \frac{2\pi}{q_{||}} e^{-q_{||}|z - z'|}.$$ 

The full screened interaction $W$ admits an expansion similar to Eq. (B2), in which $W^0(q_{||}, z, z')$ must be replaced by

| Material | Equation | $\hbar\omega_0$ or $\hbar\omega_{\text{ion}}$ (eV) | $\hbar\eta$ (eV) | $\epsilon_b$ or $\epsilon_{\infty}$ | $\hbar\omega_{\text{tr}}$ (eV) | Energy range (eV) |
|----------|----------|--------------------------------|---------------|-----------------|-----------------|-----------------|
| Ag (a)   | (A7)     | 9.17±0.19                      | 0.025±0.01    | ≈4              | –               | < 1.8           |
| Al (b)   | (A1)     | ≈15                            | ≈0.6          | –               | –               | 8 – 25          |
| Au (a)   | (A7)     | 9.06±0.18                      | 0.071±0.007   | ≈9.5            | –               | < 1.8           |
| Cu (a)   | (A7)     | 8.88±0.18                      | 0.096±0.01    | ≈8              | –               | < 1.8           |
| KCl (c)  | (A3)     | ≈0.0274                        | ≈0.0014       | ≈2.19           | ≈0.0185         | 0.01 – 0.025    |
| SiC (b)  | (A3)     | ≈0.075                         | ≈0.012        | ≈6.15           | ≈0.0999         | 0.01 – 0.2      |

**TABLE II** Dielectric parameters of selected materials, described by Eqs. (A1), (A3), and (A7), and chosen to fit the measured dielectric functions reported in the following references within the specified light-energy range: (a) Johnson and Christy (1972); (b) Palik (1985); (c) Ashcroft and Mermin (1976). The reader is referred to the book of Raether (1980) for comprehensive tables of bulk and surface plasmon energies and widths.
\[ W(q_{∥}, z, z', ω) = \frac{2\pi}{q_{∥}} \times \begin{cases} \frac{1}{\epsilon_1} \left( e^{-q_{∥}|z-z'|} + \frac{\epsilon_1(0)}{\epsilon_1(0)+1} e^{-q_{∥}(z+z')} \right), & z, z' > 0, \\ \frac{1}{\epsilon_2} \left( e^{-q_{∥}|z-z'|} - \frac{\epsilon_1(0)}{\epsilon_1(0)+1} e^{-q_{∥}(z+z')} \right), & z, z' < 0, \\ \frac{2}{\epsilon_1+\epsilon_2} e^{-q_{∥}|z|+|z'|}, & \text{otherwise}, \end{cases} \]

using the notation of Fig. 9.

Nonlocal effects can be incorporated through the hydrodynamic model outlined in Appendix A.2. The result in the non-retarded approximation reduces to

\[ W(q_{∥}, z, z', ω) = \frac{2\pi}{q_{∥}} \times \begin{cases} e^{-q_{∥}|z-z'|} + \frac{\epsilon_{s}(0)-1}{\epsilon_{s}(0)+1} e^{-q_{∥}(z+z')}, & z, z' ≥ 0, \\ \epsilon_{s}(z-z') + \epsilon_{s}(z+z') - \frac{2\epsilon_{s}(z)\epsilon_{s}(z')}{\epsilon_{s}(0)+1}, & z, z' < 0, \\ \frac{2\epsilon_{s}(z)}{\epsilon_{s}(0)+1} e^{-q_{∥}z}, & z ≥ 0, z' < 0, \\ \frac{2\epsilon_{s}(z)}{\epsilon_{s}(0)+1} e^{-q_{∥}z'}, & z < 0, z' ≥ 0, \end{cases} \]

where

\[ \epsilon_{s}(z) = \int dq_{z} \frac{e^{i q_{z} z}}{q^{2} \epsilon_{\text{lon}}(q, ω)}, \]

and \( q^{2} = q_{∥}^{2} + q_{z}^{2} \). An analytical expression for \( \epsilon_{s}(z) \) can be obtained using Eq. (A6) (García de Abajo and Echenique, 1992). This approach is formally equivalent to the specular reflection model (Ritchie and Marusak, 1966; Wagner, 1966), which applies to any \( \epsilon_{\text{lon}}(q, ω) \).

2. Cylinder

For axially-symmetric problems, it is convenient to write the bare Coulomb interaction as

\[ \frac{1}{|r - r'|} = \frac{1}{(2\pi)^2} \int_{-\infty}^{\infty} dq_{z} e^{i q_{z} (z-z')} \sum_{m=-\infty}^{\infty} e^{i m(ω-ω')} W_{m}^{0}(R, R', q_{z}, ω), \]

where

\[ W_{m}^{0}(R, R', q_{z}) = 4\pi I_{m}(q_{z} R_{<}) K_{m}(q_{z} R_{>}), \]

\( R_{<} = \min\{R, R'\} \), and \( R_{>} = \max\{R, R'\} \). In a homogeneous cylinder centered around the z axis (see Fig. 13), the screened interaction is found to be

\[ W_{m}(R, R', q_{z}, ω) = 4\pi \times \begin{cases} \frac{1}{\epsilon_{1}} \left[ I_{m}(q_{z} R_{<}) + \frac{1}{\Delta_{m}} (\epsilon_{1} - \epsilon_{2}) I_{m}(x) K_{m}(q_{z} R_{<}) K_{m}(q_{z} R_{>}) \right] K_{m}(q_{z} R_{>}), & \text{if } R, R' > a, \\ \frac{1}{\epsilon_{2}} \left[ K_{m}(q_{z} R_{>}) + \frac{1}{\Delta_{m}} (\epsilon_{1} - \epsilon_{2}) K_{m}(x) K'_{m}(x) K_{m}(q_{z} R_{>}) \right] I_{m}(q_{z} R_{<}), & \text{if } R, R' < a, \end{cases} \]

where \( \Delta_{m} = \epsilon_{2} I'_{m}(x) K_{m}(x) - \epsilon_{1} I_{m}(x) K'_{m}(x) \) and \( x = q_{z} a \).

3. Sphere

The bare Coulomb interaction can be expanded in spherical harmonics as

\[ \frac{1}{|r - r'|} = \sum_{lm} \frac{4\pi}{2l+1} W_{l}^0(r, r') Y_{lm}(Ω) Y_{lm}^{*}(Ω'), \]

where

\[ W_{l}^0(r, r') = \frac{r_{l}^l}{r_{>}}, \]
where \( r_+ = \min\{r, r'\} \), \( r_- = \max\{r, r'\} \), and \( \Omega \) denotes the direction of \( \mathbf{r} \). Using the notation of Fig. 13, the interaction near a homogeneous sphere is given by the above expansion with

\[
W_l(r, r', \omega) = \begin{cases} 
\frac{1}{c_1} W_l^0(r, r') + \left( \frac{2l+1}{\eta_1} - \frac{1}{c_1} \right) \frac{a^{2l+1}}{(rr')^{l+1}}, & r, r' > a, \\
\frac{1}{c_2} W_l^0(r, r') + \left( \frac{2l+1}{\eta_2} - \frac{1}{c_2} \right) \frac{a^{2l+1}}{(rr')^{l+1}}, & r, r' < a, \\
\frac{2l+1}{\eta_i} W_l^0(r, r'), & \text{otherwise,}
\end{cases}
\]

substituted for \( W_l^0(r, r') \). Here, \( \eta_i = (l+1)\epsilon_i + \iota \epsilon_2 \). It is also useful to write the \( r, r' > a \) part in terms of the multipolar polarizability

\[
\alpha_l^{NR}(\omega) = a^{2l+1} \frac{l(\epsilon_2 - \epsilon_1)}{\iota \epsilon_2 + (l+1)\epsilon_1}
\]

and

\[
W_l(r, r', \omega) = \frac{1}{\epsilon_1} \left[ W_l^0(r, r') - \alpha_l^{NR} \right].
\]

### APPENDIX C: Retarded scattering matrices and coupling coefficients in a sphere including nonlocal effects

The coupling coefficients of Eqs. (33) and (58) are dimensionless functions of \( v/c \) that admit the following closed-form expressions:

\[
C_l^{im} = \frac{1}{l(l+1)} \left| \frac{2mv}{c} N_{lm} \right|^2
\]

and

\[
C_l^{E} = \frac{1}{l(l+1)} \left| \frac{c}{v\gamma} M_{lm} \right|^2,
\]

where

\[
N_{lm} = \sqrt{\frac{2l+1}{\pi} \frac{(l-|m|)!}{(l+|m|)!} \frac{(2m-1)!}{(v\gamma/c)^{|m|}}} C_{l-|m|+1/2} \left( \frac{c}{v\gamma} \right),
\]

\[
M_{lm} = N_{lm+1} \sqrt{(l+m+1)(l-m)} + N_{lm-1} \sqrt{(l-m-1)(l+m)},
\]

and \( C_m^{(n)} \) are Gegenbauer polynomials (Abramowitz and Stegun, 1972). Similar formulas are encountered to describe the Coulomb excitation of atomic nuclei (Winther and Alder, 1979).

The scattering matrix elements of Eqs. (33) and (58) depend on the composition and structure of the spheres. For the homogeneous particle of Fig. 13, Mie theory leads to (Jackson, 1999; Mie, 1908)

\[
t_l^M = \frac{-j_l(x_1) x_2 j_l'(x_2) + x_1 j_l'(x_1) j_l(x_2)}{h_l^{(+)}(x_1) x_2 j_l'(x_2) - x_1 [h_l^{(+)}(x_1)]' j_l(x_2)}
\]

and

\[
t_l^E = \frac{-\epsilon_1 j_l(x_1) [x_2 j_l(x_2)]' + \epsilon_2 [x_1 j_l(x_1)]' j_l(x_2)}{\epsilon_1 h_l^{(+)}(x_1) [x_2 j_l(x_2)]' - \epsilon_2 [x_1 h_l^{(+)}(x_1)]' j_l(x_2)},
\]

where \( x_1 = ka\sqrt{\epsilon_1}, x_2 = ka\sqrt{\epsilon_2} \), the prime denotes differentiation with respect to the argument, and \( j_l^{(\pm)} = j_l - y_l \) are spherical Bessel and Hankel functions, respectively.

We can incorporate nonlocal effects in the sphere response using the formalism of Sec. A.2. This allows us to generalize the non-retarded study of dispersive spheres reported by Fujimoto and Komaki (1968). Considering a metallic sphere in vacuum [i.e., \( \epsilon_1 = 1 \), and with medium 2 inside the sphere described by \( \epsilon_{lon}(q, \omega) \) and \( \epsilon_{tr}(\omega) \)], a detailed analysis leads to

\[
t_l^E = \frac{-j_l(x_1) [x_2 j_l(x_2)]' + \epsilon_2 [x_1 j_l(x_1)]' j_l(x_2)}{h_l^{(+)}(x_1) [x_2 j_l(x_2)]' - \epsilon_2 [h_l^{(+)}(x_1)]' j_l(x_2)},
\]

where

\[
y_l = \frac{l(l+1)j_l(x_1)j_l(q_0a)}{q_0a j_l'(q_0a)} \left( \frac{\epsilon_{tr}}{\epsilon_b} - 1 \right),
\]

\[
x = ka, x_2 = ka\sqrt{\epsilon_2}, \text{ and } q_0 \text{ is given by the condition } \epsilon_{lon}(q_0, \omega) = 0 \text{ [see Eq. (A6)]}. \]
APPENDIX D: List of selected symbols and acronyms

A  vector potential
a  sphere or cylinder radius
B  magnetic induction
b  electron impact parameter
c  speed of light
Clek, CMlek  electron-sphere coupling coefficients
ey  minus the electron charge
E  electric field
F  electromagnetic force
f  far field amplitude
G  electromagnetic Green tensor
Gd  dipole-dipole interaction in vacuum
H  magnetic field
h  boundary current
h  Planck’s constant
h(l+)  spherical Hankel function of order l
I  modified Bessel function of order m
j  electron current density
ji  spherical Bessel function of order i
k  free-space light wavevector
kB  Boltzmann’s constant
kF  Fermi wavevector
Km  modified Bessel function of order m
L  length of the electron trajectory
l, m  orbital and azimuthal numbers
lmax  maximum orbital angular-momentum number
me  electron mass
ph, pf  initial and final electron wavevectors
q  wavevector transferred from the electron
q  wavevector parallel to a planar surface
q  wavevector in -plane or perpendicular to v
R  radial distance in cylindrical coordinates
R  in-plane or perpendicular ˆz component of r
R  cylindrical radial unit vector
r  radial distance in spherical coordinates
r = (R, z)  position coordinate vector
re(t)  electron trajectory
r0 = (R0, z0)  electron impact parameter at time t = 0
t  time
elek, ekM  sphere scattering matrices
ve  electron velocity vector
v  electron velocity
vF  Fermi velocity
W  scalar screened interaction
x, y, z  Cartesian unit vectors

α  particle polarizability
β  hydrodynamic wavevector
Γ(ω)  probability per unit frequency ω
γ  γ in vacuum (ε = 1)
γe  Lorentz’s contraction factor in a medium
ε  dielectric function
εD, εL, εM  Drude, Lindhard, and Mermin ε
εlon, εtr  longitudinal and transversal ε
ε  ħε is an electron energy
η  bulk plasmon damping rate
λ  free-space light wavelength
μj  coefficients for TR and SPP emission
ρ  electron charge density
ρn  photonic local density of states (LDOS)
σ  boundary charge
ϕ  scalar potential
ϕ  azimuthal unit vector
ϕq  azimuthal unit vector in q space, ⊥ q||, ˆz
φ  azimuthal angle
φin  electron-beam divergence half-angle
φout  electron-spectrometer aperture half-angle
χ  electric susceptibility
ψ  electron wave function
Ω  solid angle
ω  frequency
ωp  classical bulk plasma frequency
ωs  Ritchie’s surface plasma frequency
1D, 2D, 3D one-, two-, three-dimensional
BEM boundary element method
CL cathodoluminescence
CR Cherenkov radiation
DDA discrete-dipole approximation
DFM dark-field microscopy
EEGS electron energy-gain spectroscopy
EELS electron energy-loss spectroscopy
EFTEM energy-filtered TEM
e-h electron-hole
EIRE electron-induced radiation emission
FWHM full width at half maximum
HREELS high-resolution EELS
JDOS joint density of states
KK Kramers-Kronig
KKR Körringa-Kohn-Rostoker
LDOS local density of states
LEEM low-energy electron microscopy
LPR localized plasmon resonance
NIR near infrared
NSOM near-field scanning optical microscopy
PEEM photo-electron emission microscopy
REELS reflection EELS
SE secondary electron
SEE secondary electron emission
SEM scanning electron microscopy/microscope
SERS surface-enhanced Raman scattering
SP surface plasmon
SPP surface-plasmon polariton
S-P Smith-Purcell
STEM scanning transmission electron microscopy
STM scanning tunneling microscopy
STS scanning tunneling spectroscopy
TEM transmission electron microscopy/microscope
TR transition radiation
UV ultraviolet
XRM x-ray microscopy
ZLP zero-loss peak

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