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Eugenio Cinquanta, Eva Arianna Aurelia Pogna, Lorenzo Gatto, Salvatore Stagira & Caterina Vozzi

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REVIEW

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Charge carrier dynamics in 2D materials probed by ultrafast THzspectroscopy

Eugenio Cinquanta (Dª, Eva Arianna Aurelia Pogna (Dª, Lorenzo Gatto (Dª, b, Salvatore Stagira (Dª, b and Caterina Vozzi (Dª)

^alstituto di Fotonica e Nanotecnologie, Consiglio Nazionale delle Ricerche, Piazza Leonardo da Vinci, Milano, Italy; ^bDipartimento di Fisica, Politecnico di Milano, Piazza Leonardo da Vinci, Milano, Italy

ABSTRACT

In this review, we discuss the rich ultrafast response at terahertz (THz) frequencies of two-dimensional (2D) materials. Thanks to their unique optoelectronic properties and exceptional tunability, van der Waals organic and inorganic 2D materials, such as graphene, transition metal dichalcogenides (TMDs), and 2D perovskites, are emerging as promising platforms for the development of nano-electronic and nano-photonic devices in the THz range. The investigation of the ultrafast charge carriers dynamics resulting from their reduced dimensionality is crucial for guiding the engineering route towards novel nanotechnologies. Here, we first give a brief overview of the state-of-the-art experimental schemes for inspecting the ultrafast response of 2D materials in the THz range, including the generation and the detection of THz light and the prototypical optical pump THz probe setup. Then, we present and discuss the most relevant results, reviewing the THz ultrafast signatures of charge carriers and excitons dynamics in graphene, TMDs, and 2D perovskites. Finally, we provide a vision of the emerging tools for characterizing the ultrafast THz dynamics at the nanoscale.



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KEYWORDS

2D materials; ultrafast THz spectroscopy; charge-carriers dynamics; THz nonlinearities; excitons; graphene; transition metal dichalcogenides



CONTACT Eugenio Cinquanta e ugenioluigi.cinquanta@cnr.it Stituto di Fotonica e Nanotecnologie, Consiglio Nazionale delle Ricerche, Milano, Italy

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1. Introduction

Since the discovery of graphene in 2004 [1], 2D materials have become one of the most explored playgrounds for the ultra-scaling of devices for different applications [2–4]. Layered 2D materials, such as graphene, hexagonal boron nitride (hBN), transition metal dichalcogenides (TMDs), and related van der Waals (vdW) heterostructures, have gained great attention as a promising platform for the development of THz photonic and optoelectronic devices [5,6], able to address, with innovative solutions, the high demand for reliable and practical THz technologies. 2D materials present distinctive, highly-tunable optical and electronic properties, support extreme light confinement, and ultrafast charge carriers dynamics, that can be exploited for the manipulation, propagation, generation, and detection of THz waves with an unprecedented level of control [6].

The exploitation of light as a probe of new exotic phenomena in these emerging materials, gives access to delicate details of the physical processes under investigation. The comprehension of the fundamental mechanisms that govern the functionalities of 2D materials is a mandatory step toward their effective exploitation in practical devices. In this respect, ultrafast spectroscopy continuously increased its area of applications becoming one of the most explored characterization techniques for the study of extended as well as isolated systems, due to its capability to investigate physical effects with unprecedented details. In the last decade, time-resolved optical spectroscopy in the near-infrared (NIR) and visible (VIS) range, has been extensively applied to probe the transient response of 2D materials and related heterostructures, unveiling the effects of the strong enhancement of many-body Coulomb interactions that results from quantum confinement and reduced screening in 2D [7,8], testified by the large exciton binding energies (~hundreds meV) and the formation of multiparticle excitations, such as trions and biexcitons [9]. In particular, time-resolved spectroscopy has allowed to observe large renormalization of the optical transition energies upon photoexcitation [10,11], to track the dynamical exciton response [12–14], to probe the strong electron–phonon interaction [15,16], and many other non-linear processes (e.g. harmonics generation) in atomically-thin materials. In this framework, ultrafast spectroscopy in the THz range (1 THz = 300 μ m = 4 meV) represents a powerful tool among the timeresolved techniques, as many fundamental excitations, in strongly-correlated electronic systems and nanoscale materials, occur at such low energies. Free carriers and electron-phonon related phenomena as well as low-energy collective oscillations of conduction charges, show their fingerprint in the THz spectral range. Moreover, by lying close to the Fermi level, the charge carriers that are photoexcited by THz waves are closely connected to DC transport. By detecting in the time domain, the amplitude and phase of a THz pulse that has interacted with the 2D material, its complex dielectric function, or complex conductivity, can be directly obtained, without the need for Kramers-Kronig relations. The access to the optical conductivity, combined with the time resolution that results from the short temporal duration of the THz time-domain sources, makes ultrafast THz spectroscopy the key spectroscopy technique for the study of fundamental physical phenomena occurring in 2D materials, where the dynamical response of free charges and bound quasiparticles, together with their interplay with the lattice, are strongly influenced by the dielectric landscape and quantum confinement.

In this review, we focus on the out-of-equilibrium THz response, investigated by optical pump – THz probe (OPTP) spectroscopy, of graphene, atomically thin TMDs, and 2D perovskites, which represent three prototypical classes of 2D materials. In section 2, we discuss different schemes for the generation and detection of THz pulses, the experimental configurations for measuring the charge-carriers dynamics, and the procedures routinely exploited to retrieve the pump-induced optical conductivity in the frequency domain from the THz fields acquired in the time-domain. In section 3, we discuss the most significant results concerning the charge carriers in graphene (section 3.1), and exciton dynamics in TMDs (section 3.2) and 2D perovskites (section 3.3). In the *Perspective* we discuss ultrafast THz nanoscopy as the emerging tool to enable the extension of ultrafast THz spectroscopy to the nanoscale, explore regimes of strong-light matter interaction and resolve spatial inhomogeneities in the THz response of 2D materials.

2. Ultrafast THz spectroscopy

2.1. Generation of THz radiation

THz radiation can be generated as continuous waveforms (CW) or broadband pulses. CW THz generation can be obtained by exploiting electronic sources, photomixing, and quantum cascade lasers (QCL) [17]. In this review, we focus on the generation of broadband THz pulses that, in contrast with CW THz generation, requires the use of femtosecond laser sources and enable the investigation of ultrafast dynamics by time-resolved THz spectroscopy (TRTS).

The underlying idea is that the large bandwidth and the ultrashort duration of the driving pulse, provided by femtosecond lasers, can be leveraged to obtain transient electromagnetic fields with frequencies in the THz range. This is usually obtained following two different main strategies. The first one is inducing a transient current density of free charges, while the second one is inducing a transient polarization field in a crystal lattice [18].

The first strategy can be accomplished by photoconductive (PC) antennas [19] and this solution is typically employed when the laser system is an oscillator without any additional amplification stage. The reason is that PC antennas do not require high intensities to be operated. In fact, high pulse energies can easily damage them, unless some care is dedicated to choosing a PC antenna with a large enough aperture (and a correspondingly high bias voltage) or to adjusting the pulse energy of the generation beam below the damage threshold [20,21]. PC antennas represent also an attractive solution for the development of on-chip integrated THz systems [22,23].

A growing interest has been recently dedicated to an emerging class of THz sources called spintronic emitters [24]. In these devices, based on layered heterostructures, the photoinduced spin-current in a ferromagnetic material is converted into a transient charge-current in a non-magnetic material. This conversion relies on the inverse spin Hall effect (ISHE).

Concerning the second strategy, based on polarization effects, the generation of THz radiation is typically obtained by exploiting non-linear processes in solids. In this case, high laser intensities are in general beneficial and, the handling conditions of solid-state emitters are usually less demanding, compared to those of PC antennas. For this reason, crystals are the most frequent choice to obtain THz pulses from amplified laser systems.

The working principle of solid-state emitters usually relies on optical rectification (OR). The THz bandwidth obtained by OR is mainly limited by phase-matching conditions and by the absorption in the crystal itself, due to the interaction with phonons in the relevant spectral range. A careful choice of the crystal is therefore critical to extend the bandwidth of the emitted THz pulse up to 10 THz and beyond.

THz generation by laser-induced plasma in air is currently a fairly common technique when a bandwidth of tens of THz needs to be explored. This technique was originally interpreted as four-wave mixing (FWM) rectification in laser-ionized air, due to the coupling of ω and 2ω components by the third-order susceptibility $\chi^{(3)}$ [25,26]. A different semi-classical model based on a transient current density of the ionized electrons was then demonstrated to better explain the process [27,28]. A further quantum mechanical model can be found in reference [29].

2.1.1. Photoconductive antennas

A typical PC antenna [30,31], or PC switch, comprises an undoped semiconductive substrate and two metallic electrodes, lithographically fabricated on top of its surface. An ultrashort optical pulse photoexcites electrons to the conduction band of the substrate. In this way the PC switch is turned on and a DC bias voltage applied to the electrodes can drive a current density.

The temporal evolution of this current density is then determined by the recombination dynamics of the photo-excited free charge carriers. The acceleration of the charge carriers is responsible for the emission of radiation, consequently, the switch-on and recovery times determine the frequency range of the emitted pulse. A phenomenological model, clarifying the main physical aspects of THz generation in PC antennas can be found in reference [32]. Monte Carlo simulations were also carried out in references [33,34].

The ultrafast dynamics of the transient current density is consequently responsible for the emission of a THz pulse and, to increase its bandwidth, the optical pulse duration and the recombination time must be as short as possible. For this reason, the most successful substrate operating at an excitation wavelength of 800 nm is made of low-temperature grown GaAs (LT-GaAs). The first PC antennas employed semi-insulating GaAs (SI-GaAs) or radiation-damaged silicon-on-sapphire (SOS). Low-bandgap materials like InGaAs and InGaAs/InAlAs heterostructures are currently attracting more interest, thanks to their higher absorption at the telecom wavelength 1.55 μ m.

The emitted THz pulse propagates in the substrate, it is then collected by a high-resistivity float-zone (HRFZ) silicon lens attached to the back of the antenna and eventually transmitted to free-space. The combination of the HRFZ silicon lens with an off-axis parabolic (OAP) mirror allows for obtaining a collimated THz beam.

2.1.2. Optical rectification

Optical rectification refers to the generation of a polarization with nonvanishing time-average inside a crystal interacting with intense laser beams [35]. The non-vanishing time-average is due to the zero-frequency, or "DCcomponent", of the non-linear polarization. Thanks to this phenomenon, the transient electrical dipole induced by an ultrashort pulse inside a crystal can radiate a THz pulse. This can also be seen as a difference frequency mixing of the components within the spectrum of the ultrashort pulse.

Second-order optical rectification is usually the dominant contribution to the emission of THz radiation [36], therefore it is convenient to use noncentrosymmetric crystals, where the second-order non-linear susceptibility is non-vanishing. The highest frequency of the emitted THz pulse is virtually limited by the bandwidth of the driving optical pulse, but in practice, the emitted bandwidth is cut off by the phase-matching conditions and by the absorption due to the phonon modes in the crystal itself.

Thanks to the good phase matching at 800 nm and the high second-order non-linearity, the most common crystal used for THz generation with Ti:Sa

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based laser systems is ZnTe. This crystal offers a bandwidth up to 3 THz, limited by the absorption by IR-active phonons in the crystal. The most straightforward way to extend the bandwidth is to use crystals with optical phonons lying at higher frequencies. A popular choice is GaP, which gives access to a larger bandwidth up to 7 THz [37].

Another possibility is represented by thin GaSe crystals. If care is dedicated to meeting the correct phase-matching conditions, the generated radiation can be tuned in a frequency range approaching 100 THz [38–40]. Efficient THz generation and large bandwidths can be also obtained by organic crystals like DAST, DSTMS and OH1 [41]. These crystals often present a better phase-matching when pumped in the 1200 – 1700 nm range [21].

The generation of high-power THz pulses by optical rectification was also studied in organic crystals [42] and in LiNbO₃ [43]. A pulse front tilting scheme is required in LiNbO₃ to overcome the high-velocity mismatch between the 800 nm and the generated THz radiation [44].

2.1.3. Laser-induced air plasma

This method relies on the ionization of molecules in the gas phase by a two-color laser pulse and allows to obtain a bandwidth of tens of THz. The asymmetric oscillation of the two-color electric field can induce a drift current of the ionized electrons and this is the main contribution to the emission of a THz pulse [27,45]. This is usually obtained by focusing the 800 nm beam of a Ti:Sa laser through a thin BBO crystal (typically type I, 100 μ m-thick) for second harmonic generation (SHG). The generated 400 nm pulse can then interact with the residual 800 nm pulse at the beam waist, forming an intense two-color electric field.

The second harmonic and the fundamental pulse must be synchronized at the beam waist and it is important to maximize the polarization component of the 400 nm parallel to the polarization of the 800 nm [26]. The relative orientation of the polarization can be optimized either by rotating the BBO around the normal direction to find a compromise with SHG efficiency [28], or by using a dual-wavelength wave plate (DWP) [46,47]. Fine-tuning of the relative phase between the fundamental and the second harmonic is also necessary to maximize the generated THz field [27]. Different gases and pressure values were also considered, demonstrating a further possibility to optimize the efficiency of the process [28].

The generation of intense THz pulses by relativistic laser-induced plasma is also attracting increasing attention in recent years [48]. This can provide unprecedented THz field strengths (>GV/m) and pulse energies ($\geq mJ$).

2.2. Detection of THz radiation

Time-resolved THz spectroscopy takes advantage of coherent detection techniques, or in other words, of the possibility of accessing both amplitude and phase information from the detected THz pulse. This enables the reconstruction of the complex response of the material under study, without the use of Kramers-Kronig relations. Some incoherent detection methods like Golay cells, pyroelectric sensors, and bolometers exist as well. These detectors can be also coupled to a Michelson interferometer to characterize the power spectrum of the THz pulse, similarly to what is done in Fouriertransform infrared spectroscopy (FTIR). Here, we are going to focus on coherent methods based on PC antennas, electro-optic crystals, and gas photonics.

PC antennas identical to the ones used for THz generation can be also used for THz detection. In this case, the electrodes are not biased by a constant voltage, because the THz electric field itself is responsible for the generation of a measurable photocurrent. This can flow only when free charge carriers are generated by an ultrashort gate pulse in the semiconductive substrate between the electrodes. The measured current is then proportional to the convolution of the THz electric field and the substrate transient photoconductivity [38]. Consequently, it is possible to sample in the timedomain the waveform of the THz electric field, scanning the delay between the THz and gate pulses. The gate pulse and the lifetime of the photogenerated carriers determine the spectral sensitivity of the PC antenna.

The most common detection technique in time-resolved THz spectroscopy is electro-optic sampling (EOS). The electric field of the THz pulse can modify the polarization of an ultrashort optical pulse when these are overlapped in a crystal with a large second-order non-linear response. This can be related to the Pockels electro-optic effect and a complete frequencydomain description of its application for THz detection can be found in reference [49].

The polarization-change is proportional to the THz field and can be measured by a lock-in amplifier, using an ellipsometry setup as explained in reference [49]. The reconstruction in the time-domain of the full THz waveform is then obtained by scanning the delay between the THz and the gate pulse. The same crystals used for THz generation by OR can be used for THz detection with EOS. The absorption inside the crystal, the phase-matching conditions, and the walk-off limit the measurable bandwidth. Thinner crystals can be used to improve the detected bandwidth, at the cost of a smaller electro-optic signal, as a consequence of the shorter interaction length [50,51].

Coherent detection techniques based on gas photonics were also developed to obtain efficient broadband detection. The most common approach 8 😸 E. CINQUANTA ET AL.

is called air-biased coherent detection (ABCD) [52,53]. The detection is based on the third-order coupling of the THz electric field with an ultrashort 800 nm gate pulse and an alternating external bias, applied between two electrodes at the focal point. This interaction generates a second harmonic (400 nm) signal, which is proportional to the THz field and can be measured by a photomultiplier tube (PMT).

2.3. Experimental configurations

Figure 1 shows a typical configuration for performing an OPTP spectroscopy experiment. An ultraviolet-visible (UV-VIS) or NIR light pulse ($\hbar\omega \sim eV$) is used as a *pump* to photoexcite the sample generating a highenergy charge carriers distribution. The temporal evolution of the photoexcited charge carriers distribution, is then monitored by measuring the absorption of a second pulse, the *probe*, which is in the THz range ($\hbar\omega \sim$ 1 - 10 meV). Probe detection can be performed in transmission or reflection geometry, depending if the substrate that supports the sample is semi-transparent or reflects the THz light. Common substrates for static transmission measurements on 2D materials include quartz, HR Si, SiC, sapphire, and many polymeric materials (such as polyethylene terephthalate, PET) [54]. To perform OPTP experiments, the choice of the most



Figure 1. Prototypical setup for OPTP spectroscopy experiments. An amplified laser system delivers *fs* pulses to the THz generation and detection arms and to the wavelength conversion setup exploited for exciting the samples. The two choppers (C1 and C2) modulate the THz generation and the pump pulses at different frequencies to obtain $E_{off}(t)$ and $\Delta E(t, \tau)$, respectively. Off-axis parabolic mirrors are exploited to collimate and focus THz radiation. Balanced photodiodes are used to acquire the signals that are demodulated at the relevant frequencies by a lock-in amplifier. The setup is enclosed in a purged N₂/ dry air or kept in vacuum to remove THz absorption by water vapor. D1 and D2 are the delay stages used to control the pump-probe and EOS delay respectively.

suitable

substrate is limited to those materials, which do not present a pump-probe signal.

The relative time delay between pump and probe pulses τ is usually controlled by changing the path length of one of the two pulses, and the temporal resolution is determined by the time duration of the pump and probe pulses. In TMDs, the use of a circularly polarized pump allows for valley-selective excitation or detection. The spot size of the THz probe pulse in OPTP measurements is typically ~ 1 mm imposing a limitation on the sample size and, accordingly, on the type of 2D material that can be investigated.

The pump beam is usually modulated by an optical chopper, so that a lock-in amplifier can be used to measure the pump-induced change of the THz field $\Delta E = E_{on} - E_{off}$, where E_{on} and E_{off} represent the detected THz field with the pump beam on and off, respectively. The relative change of the probe can be evaluated by combining the differential signal ΔE with an independent measurement of E_{off} , obtained by modulating the THz generation beam with a different chopper. The pump and generation beams can be also simultaneously modulated at different frequencies to obtain both the ΔE and E_{off} signals with a single measurement, as explained in reference [55]. In the cited article, the double modulation technique is proposed using a single chopper wheel, with two different sets of slots. The rotation speed of the chopper is such that the outer set modulates the pump beam at 500 Hz, while the inner one modulates the generation beam at 333 Hz.

The relations used in [55] to compute the waveforms are here derived in a simple way, with the assumption of an instantaneous response of the photodetectors. In realistic modeling of the process, one has to take into account the temporal shape of the detected electric signal, which depends on the acquisition system, as well as on the actual modulation functions of the choppers at the two frequencies. Since all these signals are periodic in the temporal domain, they can be decomposed in the Fourier series with infinite contributions. However, the main mechanism behind the technique can be captured just by stopping these Fourier series to the first two terms, i.e. the CW and the fundamental frequency components. In this case, the current difference provided by the balanced photodetectors and sent to the lock-in amplifier at fixed pump-probe and EOS delays, can be represented by sinusoidal functions. In the absence of the pump beam the signal is written as a cosine function oscillating at 333 Hz = $\omega_g/(2\pi)$:

$$I_{off}(t) = \frac{I_{off}}{2} [\cos(\omega_g t) + 1]$$
(1)

When the pump beam, modulated at frequency $\omega_p = 3\omega_g/2 = 2\pi \cdot 500$ Hz, is photoexciting the sample, the THz probe pulses are modulated by the

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transient response of the material. The modulation function can be written in this framework as:

$$T(t) = \frac{1}{2} \left[(1 - \frac{I_{on}}{I_{off}}) cos(\omega_p t) + (1 + \frac{I_{on}}{I_{off}}) \right]$$
(2)

where the frequency of the modulation is 500 Hz = $\omega_p/2\pi$. Therefore, the signal sent to the lock-in amplifier can be modeled as $I(t) = T(t)I_{off}(t)$:

$$I(t) = \frac{I_{off} - I_{on}}{8} cos[(\omega_g + \omega_p)t] + \frac{I_{off} - I_{on}}{8} cos[(\omega_g - \omega_p)t] +$$
(3)

$$+\frac{I_{off}+I_{on}}{4}\cos(\omega_{g}t)+\frac{I_{off}-I_{on}}{4}\cos(\omega_{p}t)+\frac{I_{off}+I_{on}}{4}$$

When the pump beam is blocked, we can take $I_{on} = I_{off}$, therefore T = 1 and the lock-in demodulates at frequency ω_g the amplitude $I_{off}/2$ of the oscillation in Equation (1). When the pump beam is unblocked, we can demodulate two signals at different frequencies: I_1 at frequency ω_g and I_2 at frequency ω_p . Looking at Equation (3), we can conclude that the output given by the lock-in demodulators will be:

$$I_1 = \frac{I_{off} + I_{on}}{4} \tag{4}$$

$$I_2 = \frac{I_{off} - I_{on}}{4} \tag{5}$$

following Iwaszczuk et al., the effect of the non-instantaneous response of the photodetectors can be modeled by dividing Equation (5) by a calibration constant A, which can be experimentally determined. Since the output of the lock-in with the pump beam blocked is interpreted as the reference THz field, we can also take $E_{ref} = I_{off}/2$, $E_{pump} = I_{on}/2$, $E_1 = I_1$ and $E_2 = I_2$ to be consistent with the reference [55]. Therefore, with these definitions, we find:

$$E_{ref} = E_{off} = E_1 + AE_2 \tag{6}$$

$$E_{pump} = E_{on} = E_1 - AE_2 \tag{7}$$

which are the relations used by Iwaszczuk et al. The differential signal is consequently given by $\Delta E = E_{pump} - E_{ref} = -2AE_2$.

Two different acquisition modes can be used. First of all, it is possible to keep a constant EOS delay and scan the pump-probe delay, obtaining the pump-probe dynamics $\Delta E(t_0, \tau)$. This acquisition mode gives frequency-averaged information about the THz probe pulse. The second possibility is to acquire a complete 2D map $\Delta E(t, \tau)$, scanning both the EOS and the pump-probe delays.

Let us now introduce how the data are processed in a typical OPTP experiment in transmission geometry. The differential $\Delta E(t, \tau)$ and pumpoff $E_{off}(t)$ signals can be Fourier-transformed and used in this case to compute the transient transmission function of the sample $T^*(\omega, \tau) = \frac{\Delta \tilde{E}(\omega, \tau)}{\tilde{E}_{off}(\omega)} + 1 = \frac{\tilde{E}_{on}(\omega, \tau)}{\tilde{E}_{off}(\omega)}$. The transmission function can be theoretically modelled and compared with the experimental data. The transient response function of the material can be then retrieved by different numerical approaches [56–58]. When the photo-excited material can be considered a thin-film, as it is usually the case for samples of 2D materials, the photoconductivity is commonly directly related to the transient transmission function by the following relation [59,60]:

$$\Delta\sigma(\omega,\tau) = \frac{n_1 + n_2}{Z_0 d^*} \left(\frac{1}{T^*(\omega,\tau)} - 1\right) \tag{8}$$

where n_1 and n_2 are the real refractive indices of the media surrounding the thin-film, d^* is the thickness of the photo-excited material, and $Z_0 = 376.7$ Ω is the impedance of free-space. When the photoinduced change is small, so that $\left|\frac{\Delta \tilde{E}(\omega,\tau)}{\tilde{E}_{off}(\omega)}\right| \ll 1$, Equation (8) is often replaced by the further approximated relation [60,61]:

$$\Delta\sigma(\omega,\tau) = -\frac{n_1 + n_2}{Z_0 d^*} \frac{\Delta \tilde{E}(\omega,\tau)}{\tilde{E}_{off}(\omega)}$$
(9)

The photoconductivity is related to the photoinduced change of the dielectric function by $\Delta \sigma = -i\omega\epsilon_0\Delta\epsilon$. Therefore, if the THz response of bound charges is changing because of the pump, this will also appear in the photoconductivity spectra [58].

We now consider only the pump-induced change due to the free charge carriers. In the simplest case, the THz conductivity of the sample before and after the photo-excitation can be described by the Drude model with $\omega \tau \ll 1$. With this assumption, the response of the free carriers is approximately the real frequency-independent conductivity $\sigma = e\mu n$ and Equation (9) relates the optical pump-THz probe dynamics to the photoconductivity. We can also write an approximated relation for the photoconductivity, at the first order in the variations Δn and $\Delta \mu$:

$$\Delta\sigma(\tau) = e\mu\Delta n(\tau) + en\Delta\mu(\tau) \tag{10}$$

where μ and *n* are respectively the mobility and the carrier density when the pump beam is off. If *n* is negligible with respect to the photogenerated carrier density Δn , as it typically happens in intrinsic semiconductors, Equation (10) becomes $\Delta \sigma(\tau) = e\mu\Delta n(\tau)$.

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The applicability of these hypotheses must be always carefully evaluated because the role of bound charges and collective excitations can significantly complicate the physical interpretation of OPTP dynamics [60,62]. The most complete set of information can be retrieved by the frequency-resolved photoconductivity spectra.

3. Ultrafast THz response of 2D materials

In this section, we present a compendium of the most recent studies of charge carrier dynamics in 2D materials, including graphene and graphene-based heterostructures, TMDs, and 2D perovskites, as investigated by ultrafast THz spectroscopy. For graphene, the response to THz driving fields is dominated by the interaction with mass-less Dirac electrons, giving rise to strong non-linear response, mostly facilitated by intraband conductivity [63]. In the following, we present the key experiments on graphene charge carrier dynamics, showing dependence on intrinsic (e.g. doping) and extrinsic parameters (e.g. defect, substrate), with a focus on the several results of OPTP that have significantly contributed to the understanding of graphene THz nonlinearity. For TMDs, OPTP spectroscopy revealed the free charges, excitons, and trions dynamics. Moreover, it allowed understanding of the impact of the growth conditions, e.g. the role of defects, substrate, and doping. For the 2D perovskites case, OPTP allowed observing the presence of excitons at room temperature and their interplay with the polaronic unbound carriers, confirming the excitonpolaron picture.

3.1. Graphene

Graphene is an elemental semimetal endowed with large electron mobility (>100000 cm²V⁻¹s⁻¹ [64]), broadband optical absorption [65], easily-tunable Fermi energy E_F (up to ~ 1 eV) by electrostatic gating, ensued by the linear dispersion of its conduction (CB) and valence (VB) bands, crossing at the Dirac points, at which there is a vanishing electron density of states [66], see Figure 2(a).

Steady-state transport properties of graphene are controlled by charge carriers residing near the Fermi level, while, the transport in high-speed devices depends on higher-energy charge carriers, whose temperature is increased by the presence of dynamic external fields.

First studies of the relaxation dynamics of the charge carriers in singlelayer graphene (SLG) were mainly driven by the scientific interest in the effect of reduced dimensionality on the electron–electron and electron– phonon interactions [67–71]. More recently, the interest has multiplied by the emergence of highly promising optoelectronic applications exploiting



Figure 2. (a)Electronic band structure of graphene. Conduction and valence bands cross at six K points, at which they have linear dispersion relation forming Dirac cones, corresponding to the energy spectrum of massless Dirac fermions [83]. (b) Calculated optical absorption of graphene as a function of the photon energy at different Fermi level/ sheet resistance R [84]. (c) Broadband THz frequency-dependent conductivity of graphene measured with commercial and air plasma TDS systems and fit with Drude model of Equation 12 (dashed black line), from Ref. [54].

the ultrafast dynamics of graphene carriers, such as ultrafast THz photodetectors [72–75], high-speed large-bandwidth optical modulators [76,77], ultrafast THz saturable absorbers [78,79], efficient terahertz frequency multiplier [80] and phase-shifters [81,82].

Light absorption in SLG is defined by the interplay of interband [85,86] and intraband [87–90] electronic transitions, see Figure 2(b). The first dominates in the VIS, NIR and mid-infrared ranges and can be described by the interband optical conductivity: σ_{inter} [91]:

$$\sigma_{inter} = \frac{e^2}{8\hbar} \left[\tanh(\frac{\hbar\omega + 2\mu}{4k_B T_e}) + \tanh(\frac{\hbar\omega - 2\mu}{4k_B T_e}) \right]$$
(11)

where *e* is the electron charge, k_B is the Boltzmann constant, μ is the chemical potential and T_e is the charge-carriers temperature describing the Fermi-Dirac (FD) distribution of charge-carriers in SLG. For large photon energies ($\hbar\omega > 2\mu$, $\hbar\omega > 4k_BT_e$) σ_{inter} reduces to $\frac{e^2}{4\hbar}$ giving a frequency independent absorbance A = $\pi\alpha \sim 2.3\%$ where α is the fine structure constant [65].

At THz frequencies, interband transitions are usually inhibited by Pauli blocking since a SLG is always endowed with finite doping (>30 meV), i.e. $\hbar\omega < 2 \mu$ and intraband carrier transitions dominate [92]. The intraband

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absorption σ_{intra} is well-described by the Drude model of a 2D electron gas [93]:

$$\sigma_{intra} = \frac{\sigma_{DC}}{(1 - i\omega\tau)} \tag{12}$$

where σ_{DC} is the DC conductivity at $\omega = 0$ and τ is the scattering time. The Drude weight D of graphene differs from that of conventional metals and semiconductors, being $D = v_F e^2/\hbar \sqrt{\pi n}$ with v_F the Fermi velocity and *n* the charge carrier density [90,94,95] and *h* the Planck constant. A microscopic description of intraband optical conductivity can be found in Ref. [96], where long-range Coulomb scattering with impurities is included in the calculation of the energy-momentum dependent scattering rate $\Gamma = 1/\tau$. Semiclassical Boltzmann transport theory [91] allows to extract *n* and the carrier mobility μ_e from the σ_{DC} and τ as follows [54]:

$$n = \frac{\pi h^2}{e^4 v_F^2} \left(\frac{\sigma_{DC}}{\tau}\right)^2 \tag{13}$$

$$\mu_e = \frac{e^3 v_F^2}{\pi h^2} \frac{\tau^2}{\sigma_{DC}} \tag{14}$$

Accordingly, THz spectroscopy measurements, in transmission and reflection mode [54,97], enables for a non-destructive characterization of the electrical properties of SLG, see Figure 2(c).

The charge carrier dynamics in SLG has been extensively investigated using pump-probe techniques in which the charge carriers are brought outof-equilibrium by impulsive excitation with an electrical or optical pulse, and, the equilibrium recovery is studied in the time-domain.

Impulsive excitation *via* interband absorption of femtosecond VIS or NIR pulses can be exploited to induce non-thermal distributions of charge carriers [68,69], peaked at half of the pump photon energy from the Dirac point, in the CB for electrons and the CV for holes. This first stage of the relaxation dynamics, needed by the carriers to assume FD distributions with defined temperature T_e and a larger momentum spreading, is referred in the literature as *thermalization*. The subsequent decay through quasi-equilibrium states described in terms of a hot T_e is called *cooling*.

The thermalization occurs on sub-100 fs timescale, and it has been rationalized in terms of electron-electron Coulomb scattering processes [68–70,98], such as interband and intraband carrier-carrier scattering and Auger processes. The latter includes impact ionization or carrier multiplication, and the Auger recombination [69,70,98,99]. This ultrafast carrier heating was first observed by time-resolved ARPES by Gierz *et al.* [70] in 2013, as a smearing of the electron FD distribution in the energy-

momentum space. Interestingly, the thermalization is accompanied by hot carrier photoluminescence (HotPL) [100–102], first discovered by Lui *et al.* [103], due to the radiative recombination of the excited charge carriers emitting a broad spectrum extended from the VIS [104] to the THz [105] with a frequency-dependence resembling the emission of a gray body at temperature T_e [102,103], see Figure 3.

The cooling of the hot thermalized charge carriers has been studied by OPTP spectroscopy [71,96,106–112] which allows following the timeevolution of T_e as an ultrafast electron thermometer, monitoring the induced modifications to the THz absorption. In intrinsic graphene ($E_{\rm F} < 0.1$ eV), the ultrafast photoexcitation produces an increase of the THz conductivity, as expected for a semiconductor after interband photoexcitation. In doped graphene ($E_{\rm F} > 0.1$ eV), instead, a reduction in the THz conductivity was observed by several groups [71,96,106,110,111,113], detected in transmission geometry as a transmission increase $\Delta T > 0$, that scales roughly linearly (or weakly sub-linearly) with the change ΔT_e in the temperature of the charge-carrier distribution. The negative THz photoconductivity, which indicates that the ultrafast interband excitation



Figure 3. (a-c) Schematic illustration of the HotPL emission mechanism in n-doped SLG from Ref. [102]. Dirac cones populated by e (red)-hole (h,blue) with thermal distribution at T₀ at the equilibrium (a); out-of-equilibrium charge distributions formed within 20 fs after ultrafast laser excitation (b), thermalized e-h distributions at $T_e>T_0$ with two different chemical potentials formed on a time scale comparable with the pump pulse duration leading to radiative electronhole recombination, i.e. HotPL, restoring the initial equilibrium h density (c). (d) HotPL spectrum at different T_e from Ref. [[103]].(e-f) Photoinduced change in the THz transmission through graphene (e) and inferred change in the graphene conductivity (f) from Ref.[106] [].(g) Temporal evolution of the change in the maxima of the transmitted THz waveforms normalized by the THz probe for different pump fluences [106]. The inset shows the decay times extracted by fitting the dynamics with single exponential function. Panels e-g are reprinted with permission from Ref.[106]. Copyright {2022} American Chemical Society.

makes doped graphene more resistive to THz fields, was explained in terms of *hot-carrier multiplication* [71,96,114], i.e. the scattering of photo-excited electrons with the background electrons in the Fermi sea which increases their energy. This electron-electron scattering has a rate that can largely exceed that of electron-phonon scattering and is related to the ultrafast thermalization processes producing the increase of T_e by supporting fast energy and momentum exchange within the entire electron population.

The ultrafast electron transport in graphene, on timescales longer than that of the thermalization, is described considering time-dependent quasiequilibrium within the electron population, using a thermodynamic model of the conductivity [96,115,116]. Charge carrier heating can be induced also by intraband excitation with strong THz fields acting on the free carriers in the vicinity of the Fermi level [115,117]. Since the thermalization is much faster than the duration of the THz field cycle, the THz energy is quasiinstantaneously converted into the thermal energy of the electron population, which is simply described by its instantaneous temperature T_e , which accumulates during the THz field oscillation as shown in Figure 4(a).

The carrier heating must obey the conservation of carrier density, which implies that the chemical potential μ decreases with increasing T_e while the FD distribution broadens, as depicted in Figure 4(b). The downshift of the chemical potential, in turn, causes an increase in the number of optically active interband transitions at energies 2 μ and explains the reduction of intraband conductivity due to conservation of the total spectral weight, as sketched in Figure 4(c). The heating efficiency per photoexcitation energy E_{phot} has been evaluated in Ref. [108] as $\rho = \Delta Q/E_{phot}$ with ΔQ corresponding to the heat absorbed by the electron bath, and varies with the excitation



Figure 4. (a) The calculated time-dependent T_e in graphene during the interaction with the THz fields in the range 2.3–120 kVcm⁻¹ compared to the absolute value of the experimentally measured ultrafast electric field transient as a gray area, from Ref. [115]. (b) Sketch of the evolution of the FD distribution (f_D) and charge carrier population N_e with T_{er} , T* T₀, from Ref. [116], in case of adherence (dashed red solid line) or violation (solid red line) of particle number conservation, implying constant or down-shifted chemical potential, respectively. (c) Change in the complex-valued conductivity spectrum of doped graphene with T_e in case of spectral weight conservation, from Ref. [116].

fluences, from 1, standing for total energy conversion, to an asymptote of ≈ 0.5 approached at high fluences due to the competition with energy dissipation channels such as the optical phonon emission [108].

The combination of the reduced THz absorption due to the highly efficient THz-induced carrier heating, and the ultrafast heating/cooling dynamics of hot electrons, results in an intense nonlinear response of graphene at THz energy which allowed observing THz harmonics with multicycle THz pulse [118].

Hot charge carriers cooling proceeds via energy transfer to the lattice through the emission of strongly coupled optical phonons (SCOP) till a common temperature is reached. Then, the thermalized electron-optical phonon bath relaxes via anharmonic coupling to the acoustic phonons.

Optical phonons emission requires minimum excess energy, corresponding to G Raman mode (wavevector q = 0, $\omega_G \sim 1560 \text{ cm}^{-1}$, equivalently $E_G \sim 190 \text{ meV}$) [15] of the lowest-energy optical phonon branch, and it can involve either intraband transitions causing the electrons to cascade down to Dirac point, or electron-hole recombination or intervalley scattering by zone boundary phonons. Despite this energy constraint, this cooling channel is rather efficient thanks to the continuous re-thermalization of the electronic system through carrier-carrier scattering, which guarantees the presence of carriers with enough energy to emit optical phonons [112,119], see Figure 5(a).

The direct coupling of the hot electronic system to acoustic phonons is in principle an inefficient cooling channel, which would lead to hundreds of picoseconds recovery time, due to the moment conservation constraints limiting the interaction to low-energy acoustic phonons with momentum $q < 2k_F$, with the Fermi momentum $k_F = \frac{E_F}{h_{VF}}$. However, defects or disorder can mediate



Figure 5. (a) Schematic overview of the hot-carrier relaxation channels in graphene including the *thermalization* to hot FD distributions, hotPL, *cooling* via emission of optical phonons anharmonically coupled with acoustic phonons, near-field coupling to the substrate phonons, from Ref. [120]. b)Schematic illustration of electron relaxation via *supercollision cooling* with acoustic phonons without (left) and with (right) disorder, where the presence of disorder supports the emission of finite momentum q_d acoustic phonons, from Ref. [121]. Panel a is reproduced from Ref. [[120]] with permission from the Royal Society of Chemistry.

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the scattering with high-energy acoustic phonons providing the required extra momentum. The disorder-assisted scattering is referred to as *supercollision cooling* which is expected to be especially relevant for carriers near the Dirac point and in high-disorder low-electrical mobility graphene [121,122], see Figure 5(b).

Following Ref. [121], which is based on the theory developed in Ref. [123], the supercollision cooling time can be estimated as $\tau_{cool}^{SC} = (3\frac{A}{\alpha} \cdot T_l)^{-1}$, where T_l is the lattice temperature, and $\frac{A}{\alpha} = \frac{2\lambda k_B}{3k_F \ell \hbar}$. Here the mean free path is $\ell = v_F \tau_{ms}$, with the momentum scattering time given by $\tau_{ms} = \frac{\mu E_F}{ev_F^2}$, and $\lambda = \frac{D_e^2 2E_F}{\rho v_s^2 \pi (\hbar v_F)^2}$, where ρ is the mass density, v_s is the sound velocity and D_e is the electron-phonon deformation potential which can be determined by transport measurements assuming phonon-limited momentum scattering [124]. Accordingly, supercollision theory predicts a decrease in the cooling time for increasing defects density, as it has been observed by NIR pump-probe spectroscopy on SiO₂-supported graphene [125]. Currently, there are contrasting views in the literature regarding supercollision theory with some studies, such as Ref. [126] questioning its role.

3.1.1. Graphene-based heterostructures

Graphene deposited on standard SiO₂ presents a high degree of crystal disorder. Charge carriers scattering with impurities, defects and SiO₂-optical phonon reduces the carrier mobility. Moreover, graphene tends to conform to the silicon oxide substrate forming strong spatial perturbations which have been observed by scanning tunneling microscopy [127]. Charged impurities trapped in the substrate or at the graphene/substrate interface result in an effective p- or n-doping. OPTP measurements revealed that in high-quality ($\mu_e > 10\ 000\ \text{cm}^2\text{V}^{-1}\text{s}^{-1}$) suspended graphene [112] the charge-carrier relaxation is dominated by the cooling through optical phonons, which is more efficient than disorder-assisted supercollision cooling for which a much slower cooling time (>10\ ps) is predicted.

Encapsulation with different dielectric materials has been proposed to preserve the superior properties (i.e. high-quality graphene with mobility as large as $\mu_e > 30000 \text{ cm}^2 \text{V}^{-1} \text{s}^{-1}$ and low intrinsic doping) of suspended graphene without losing the advantages of having a substrate. Among the different encapsulants, hexagonal boron nitride (hBN) has been identified as a promising candidates being an isomorph of graphite with a small lattice mismatch ($\delta = a_{hBN}/a_{graphene} = 1.8\%$ [127]), transparent to VIS and NIR thanks to the large electronic gap ($E_g \sim 6 \text{ eV}$). Furthermore, due to the strong ionic nature of the in-plane bonding, it is relatively inert and free of dangling bonds and surface charge traps. Its surface is planar on the atomic scale and suppresses graphene rippling reducing the charge carrier scattering induced by surface roughness.

The hot electrons cooling in hBN-encapsulated graphene has been investigated by several techniques, including transport [128], time-resolved photocurrent, NIR-transient absorption and OPTP [129]. These studies demonstrate the appearance of an additional relaxation channel: the near-field coupling of the graphene charge carriers with the hyperbolic phonons (HP) of hBN whose theoretical definition is provided in Ref. [130]. HP in hBN have mid-infrared energies ($E_{HP} \sim 110$ and 140 meV) and are associated with large phonon densities boosting the hot electrons cooling dynamics by supporting a very efficient Super-Planckian out-of-plane heat transfer [129,130].

To suppress the hyperbolic cooling and obtain longer-living charge carriers excitation, the encapsulation with thin films of TMDs, which have reduced hyperbolicity [112] compared to hBN, has been proposed. High-quality devices with mobility up to 100 000 cm² V⁻¹ s⁻¹ [131] have been obtained using WSe₂. However, the cooling dynamics of high-quality graphene is intrinsically limited by the relaxation *via* emission of SCOPs, which dominates the cooling when competitive channels are suppressed, imposing cooling time of few picoseconds. Similar dynamics have been observed by OPTP measurements in suspended high-quality graphene [112] where interaction with the substrate is inherently excluded, see Figure 6(c).



Figure 6. (a) hBN encapsulated graphene showing near-field energy transfer from hot electrons to the encapsulant explaining the observed picosecond OPTP dynamics, from Ref. [129].(b) Calculated dielectric constants of hBN (blue lines) and WSe2 (red lines) along the in-plane directions (xx and yy, solid lines) and along the c-axis (zz, dashed lines) showing reduced hyperbolicity, from Ref. [112]. (c) OPTP dynamics of suspended graphene from Ref. [112]. Further permission related to the material excerpted to produce panels b-c, should be directed to the ACS.

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In high-quality graphene, diffusive cooling becomes competitive with optical phonon emission as an additional relaxation channel [132] that however is observed only with μ m size pump and probe spots.

3.1.2. Gate-tunability

By tuning the chemical potential *via* electrostatic gating, below or above the half frequency of impinging radiation, large tunability of the static and transient [133] interband absorption of graphene has been demonstrated in the VIS and NIR.

At THz frequencies, modulation of the charge carrier density using either electrostatic gating or optical excitation allows for achieving large tunability of the intraband absorption (Figure 2(b)).

OPTP measurements as a function of E_F , revealed that the THz photoconductivity of graphene can be continuously tuned from semiconductorlike positive values, displayed at near-zero carrier density reached at gate voltage $V_g = V_{CN}$, to metal-like negative values at high carrier densities [110,111,134], see Figure 7.

The multiple sign change in the THz photoconductivity (see Figure 7(b)), has been related [110] to the temperature-dependence of the Drude weight and of the scattering rate Γ , see Figure 7(c,d). It was shown [96] that the ultrafast electronic thermalization in photoexcited undoped graphene ($E_F <$ 0.1 eV), can also lead to free carrier multiplication via interaction with the VB states, in addition to the hot carrier multiplication, which instead preserves the total number of free carriers. Carrier multiplication was first predicted by Winzer et al. [135], observed by Plötzing et al. [136] and rationalized in terms of electron kinetics by Tomadin et al. [96].



Figure 7. (a) Sketch of the OPTP experiment on gated SLG, where the differential transmission of a THz probe pulse is investigated as a function of the Fermi level controlled by field-effect with a gate voltage V_g [110]. (b) Temporal evolution of the change in the THz field transmission $\Delta E/E$, proportional to SLG differential conductivity σ , as a function of the Fermi energy E_F . (c) Instantaneous $\Delta E/E$ signal at zero pump-probe delay showing the change of sign as increasing the doping from the charge neutrality point (E_F =0). The corresponding change in THz conductivity $\Delta \sigma$ is shown at the right axis. Panel a is adapted from Ref. [110]. Panels b-c are adapted with permission from Ref. [111] Copyright {2022} American Chemical Society.

The doping allows also to tune the fluence dependence of the signal, exploiting the variation of the *Te*-dependence of the electronic heat capacity, which is quadratic at the charge-neutrality point CNP and becomes linear at high doping [111].

The conversion efficiency of THz third-harmonic generation [137] can be controlled by electrostatic gating exploiting the dependence of electronic heat capacity on doping. The THz nonlinearity of graphene has been indeed attributed to the collective thermodynamic response of free-carriers to the THz field. Increasing the free carriers, i.e. E_F , enhances the power absorption leading to an increasing nonlinearity, scaling approximately with the square of E_F . However, the THz-induced electronic temperature rise, decreases with increasing doping because the absorbed THz energy is shared among a very large number of carriers, diminishing the material's thermodynamic nonlinearity. Accordingly, there is an optimal doping to enhance graphene's THz nonlinearity [137].

3.2. Transition metal dichalcogenides

Atomically thin TMDs are inorganic 2D crystals emerging as novel quantum materials, that, differently from graphene, exploit a direct band gap to conduct electricity whenever the electrons absorb enough energy through heat, light, and other means, therefore promising groundbreaking impacts in nano-electronics and nano-photonics. In analogy with graphite, TMDs can be easily thinned down to one atomic layer using mechanical as well as chemical exfoliation techniques. Moreover, they can be synthesized as single and multilayers with bottom-up approaches like Chemical Vapour Deposition (CVD) and Metal-Organic Chemical Vapour Deposition (MOCVD) techniques. At the single-layer limit, the physical properties of TMDs are dominated by the Coulomb interaction between the carriers resulting from the reduced dielectric screening and strong quantum confinement. Therefore, in contrast with other inorganic semiconductors, a single layer of TMDs shows a rich and robust exciton population at room temperature that promotes them as one of the most explored classes of materials for beyond state-of-the-art (opto)electronics. Both geological and synthesized TMDs are naturally doped, with the consequent electron/ hole excess that has a sizeable impact on their electronic and optical properties.

Ultrafast THz spectroscopy emerged as a powerful technique to study the carrier dynamics in monolayer and few-layers TMDs, thanks to its peculiarities. The THz probe strongly couples to charged particles like free carriers and intra-exciton transitions as far as the probe photon energy matches some allowed 1*s* bright to *np* dark states transition [138,139]. When these transitions are not accessible by the probe bandwidth or cannot be resolved at room temperatures, it is still possible to obtain an indirect signature of the presence of the excitons in the THz transient photoconductivity. Indeed, bound e-h pairs induce a negative Im ($\Delta\sigma$) due to the exciton dielectric polarization and the related capacitive response [140,141].

As mentioned in Sec. 1, THz spectroscopy can access the complex photoconductivity spectrum by acquiring the entire waveform of the THz electric field at different pump-probe delays. Alternatively, it is possible to obtain the frequency-averaged real and imaginary photoconductivity by measuring the change of two special points of the THz waveform, as a function of the pump-probe delay. One point is the peak of the electric field, which is sensitive to amplitude variations and is slightly affected by small phase changes. A pure amplitude change is given by the low-frequency response of a photoexcited free-carrier gas. Therefore, in the absence of a large temporal shift induced by the pump, the dynamic of the peak reflects the conductive response (current in phase with electric field) of the sample and it is routinely described as the frequency-averaged real part of the photoconductivity. The other point is the zero-crossing, which is sensitive to phase shifts but not to amplitude variations. A phase shift can be induced by the polarizability of an exciton population [141], which is related to a positive change of the real dielectric function and a correspondingly negative change of the imaginary photoconductivity. For this reason, the dynamics of the zero-crossing point reflects the capacitive response (polarization in phase with the electric field) and it is routinely described as the frequency-averaged imaginary part of the photoconductivity [62].

Molybdenum Disulphide (MoS₂) has been the most studied TMD, since the observation of the 10^4 enhancement of the PL emission and the crossover from indirect to a direct band gap in the limit of the single monolayer [142]. Concerning the carrier and exciton dynamics, a debate started after the observation of trion, a three-bodies quasiparticle formed by two electrons (holes) and one hole (electron), in the optical response of monolayer MoS₂ [143].

C. H. Lui and coworkers in their seminal work exploited OPTP spectroscopy to investigate the THz dynamic response of CVD grown monolayer MoS_2 [[144]]. The top panels of Figure 8(a) show the THz probe electric field in the time domain together with a sketch describing the pump-probe experiment. The bottom panels report an unexpected positive pumpinduced THz response that would mean a persistent decrease of the conductivity of the material after the photo-injection of additional charge. In details, the bottom left panel shows a 150 ps long positive $\Delta T/T$ ($\frac{T_{on}-T_{off}}{T_{off}}$) composed by a short (~1 ps) and long (~42 ps) component, while the transient conductivity spectra reported in the bottom right panel shows negative real (Re($\Delta \sigma$)) and imaginary (Im($\Delta \sigma$)) parts. This counter-intuitive response observed regardless of the temperature (4 – 350 K) and the fluence



Figure 8. (a) The THz electric fields exploited as probe and (b) a sketch of the OPTP experiment; (c) and (b) the frequency-averaged and frequency-resolved pump induced THz response of monolayer MoS₂, respectively (adapted from Ref. [144]).

 $(0.4 - 170 \ \mu \ J \ / \ cm^2)$, was explained by considering the presence of a doping-induced extra charge in the pristine monolayer that couples with the photo-generated el-h pairs forming negative trion (two electrons and one hole). The effective mass m^* of this quasiparticle is higher as compared to the one of the unexcited material with the consequent decrease of the carrier mobility $\mu = \frac{er}{m^*}$. From Equation 10, it follows that the transient response $\Delta\sigma$ becomes negative in presence of heavier carriers in the photo-excited material.

The attribution of the negative conductivity to trion has been confirmed by exploiting different pump photon energies in OPTP experiments. When the pump photon energy falls below the *A* excitonic resonance, neither free carriers nor excitons can be generated in the material, and the THz response is totally quenched. As the pump photon energy approaches the *A* exciton, then the formation of exciton occurs and THz response sharply increases, in the absence of photo-generated free carriers. As the THz probe cannot couple with charge-neutral species and intra-exciton transitions lie at much higher energy with respect to the \sim 4 meV probe used for the experiments, the resulting negative THz dynamics can be solely ascribed to the presence of a charged exciton, i.e. the trion. In [[144]] the authors further confirmed this claim by tuning the doping of the pristine monolayer. Remarkably, the effect of the *p* doping, which reduces the initial free charge, is the suppression of the trion formation, while *n* doping partially restores the pristine monolayer PL and THz dynamic response. These results support that trions form from the coupling of the doping-induced charge with the photo-generated e-h pairs population.

Although strong evidence supports the picture developed in [[144]], C. J. Docherty et al. observed a different ultrafast THz response for a CVD grown monolayer MoS₂. In particular, in [145] a positive transient THz conductivity ($\Delta T/T < 0$, see Equation10) has been reported, with a signal that recovers within hundreds of fs with a simple exponential decay, regardless of resonant or non-resonant photo-excitation, suggesting that monomolecular processes are at play. Doping and fluence-dependent measurements allowed to assign of the fast recombination to trap-assisted processes at the surface due to the presence of a trap density higher than 12×10^{13} cm⁻². The frequency-resolved transient conductivity spectra reported in Figure 9(a) confirm the differences observed in the frequencyaveraged THz response. In [145] indeed, the authors observed a positive Re $(\Delta \sigma)$ accompanied by a negative Im $(\Delta \sigma)$. This behavior typically occurs when a restoring force acts on the photo-generated free carriers, or when the probe photon energy matches a resonance, such as phonons, plasmons, or excitons. In this respect, trions in monolayer MoS₂ have a binding energy of



Figure 9. (a) Top and bottom panels reports the photoconductivity of monolayer and trilayer MoS₂, respectively (adapted with permission from Ref. [145]). (b) Frequency-averaged (top-right and bottom-left panel) and frequency-resolved (bottom-right panel) photoconductivity of monolayer MoS₂ adapted with permission from [147].

~18 meV (4.4 THz) that falls in the 0 – 5 THz (0 – 20 meV) spectral range of the probe used in [145]. Therefore, J. C. Docherty *et al.* fitted the $\Delta\sigma(\tau, \omega)$ with a Drude-Lorentz model, where an oscillator at ~3.6 THz (15 meV) is needed to account for the trion resonance and the Drude term accounts for a small fraction of free charges. For the trilayer MoS₂, J.C. Docherty *et al* observed a similar response but with the resonance located at much higher energy outside the THz probe region. As the trion is expected to occur for the monolayer only, this resonance can be related to excitons only. For the trilayer, the exciton binding energy is ~100 meV (15 THz) and matches the dominant 1*s*-2*p* intra-excitonic transition [146], thus inducing the Lorentz lineshape for the $\Delta\sigma(\tau, \omega)$.

In 2016, P. D. Cunningham *et al.* investigated CVD grown monolayer MoS_2 with OPTP in the 0.5 - 6 THz spectral region [147]. Here the authors observed an increased photo-conductivity with an hundreds of fs dynamics similar to the one reported in [145] (see the top-right panel of Figure 9(b)), but sizeable differences in the conductivity spectra, i.e. a positive Re $(\Delta\sigma)$ with an almost negligible Im $(\Delta\sigma)$ (see bottom-right panel of Figure 8(b)). This outcome points to a free-carrier dominated conductivity, and the deviation from a pure Drude dispersion has been ascribed to carrier back-scattering at defects or grain boundaries where random disorder causes destructive interference among carrier scattering events. This leads to a non-Drude conductivity that can instead be described by the phenomenological Drude – Smith model [148]. The authors suggested that the absence of the signature for the trions, i.e. the negative photo-conductivity and the resonance at ~15 meV, can be explained by considering the oxygen neutralization of the n – type doping at the grain boundaries.

The striking differences observed in the dynamical THz responses suggest that the parameters adopted for the CVD growth of monolayer MoS_2 play a crucial role in determining the morphology and the doping level of the material. These in turn influence the formation of trions. Another detrimental aspect is the role of the substrate and the capping layer if present. Indeed, they determine the dielectric landscape for the monolayer and consequently the exciton and trion binding energies.

For the multi-layer case, the ultrafast THz dynamics of a chemically exfoliated free-standing MoS_2 laminate consisting of 1-6 layers were studied by S. Kar *et al.* In [149] the authors performed a fluence-dependent study and analyzed the frequency-averaged THz response of the sample. The authors interpreted the results by developing a system of rate equations incorporating defect-assisted Auger scattering of photo-excited electrons, holes, and excitons. The dynamics of the laminate resulted to be composed of a fast (<3 ps) and a slow (hundreds of ps) relaxation channel. It emerged that defect states are used on a sharing basis by the free carriers as well as the excitons in relaxation kinetics. The rate coefficients of

the free carrier and exciton capture obtained by the fit of the dynamics with the solutions of the rate equations were almost comparable and excitons bound to defects emerged to have a slower relaxation with respect to the free charge.

Concomitantly with the work of C. H. Lui *et al.* on the monolayer, J. H. Strait *et al.* presented their ultrafast THz investigation of bulk MoS_2 [150]. In this work, the authors performed OPTP spectroscopy to study the carrier mobility and the dynamics associated with carrier relaxation. Here, by varying the temperature in the 30 - 300 K range, a free-carrier like THz conductivity was observed. From the fit with the Drude model, the authors extracted carrier mobility of $4200 \text{ cm}^2 \text{V}^{-1} \text{s}^{-1}$ at 30 K. From the temperature dependence of the scattering time, they found how the mobility is intrinsically limited by acoustic phonon scattering below 200 K, while at the higher temperature the contribution of optical phonon becomes evident. Concerning the carrier recombination, by varying the fluence and fitting the dynamics with the solutions of a system of rate equations, the authors obtained that the carrier relaxation lasting tens of nanoseconds mainly attributed to Auger scattering into defects.

In the family of TMDs, the carriers and excitons ultrafast dynamics of CVD grown 3 - 4 layers WS₂ laminate ha been investigated using OPTP spectroscopy. In [151], the authors reported the methodology adopted to disentangle the free carriers and exciton contributions. As summarized in the introduction of this section, by following the peak and the zero-crossing points it is possible to grab the bound and unbound carrier contributions to the frequency-averaged complex photoconductivity (see Figure 10). In this case, a restoring force provided by the excitons was also proposed as the origin of the localization of the unbound carriers. Figure 10(a) shows the frequencyaveraged $\operatorname{Re}(\Delta\sigma)$ and $\operatorname{Im}(\Delta\sigma)$, both characterized by a 4 ps formation time that is, the concomitant ultrafast formation of free carriers and excitons, respectively. Remarkably, the relaxation dynamics of the $Im(\Delta\sigma)$ emerged to be much faster than the Re($\Delta \sigma$), suggesting that excitons recombine faster than free carriers. As shown in Figure 10(c), Re($\Delta\sigma$) was fitted with a triexponential curve which describes the contribution of el-ph scattering $(\tau_1 \sim 1 - 2 \text{ ps})$, many-body effects $(\tau_2 \sim 16 \text{ ps})$, unbound el-h recombination $\tau_3 \sim 110$ ps. An offset has been also included in the model to consider mid-gap trap-assisted processes (1 ns). The relaxation dynamics of $Im(\Delta\sigma)$ were conversely fitted with a bi-exponential curve that returns two fluenceindependent decay times of $\tau_1 \sim 1 - 2$ ps and $\tau_1 \sim 16$ ps assigned to excitonphonon coupling and exciton lifetime, respectively (see Figure 10(d)).

The frequency-resolved transient THz conductivity of WS_2 shown in Figure 11(a) returns a Drude-Lorentz response, in analogy with the MoS_2 laminate case, where the authors added an oscillator to the fit function to



Figure 10. (a) Reference (black curve) and the pump modulated (red curve) electric fields in the time-domain. The vertical dashed lines lie at those points in the time domain used to acquire the frequency-averaged real (peak of the electric field) and imaginary (zero-crossing point) photoconductivity. (b) Pump-probe scans were performed at the peak and at the zero-crossing points. (c) and (d) Fluence-dependent scans of the real and imaginary photoconductivity of WS₂, respectively [151].



Figure 11. (a) Photoconductivity of multilayer WS₂ obtained with (left panel) different fluences and at (right panel) different pump-probe delays (adapted with permission from [151]). (b) Photoconductivity of monolayer WS₂ acquired at different pump-probe delays in the 1 - 8 THz spectral range (adapted with permission from [152]).

follow the negative $Im(\Delta \sigma)$. While for MoS₂ the Lorentz line-shapes were directly attributed either to trion or exciton resonances, for WS₂ X. Xing *et al.* invoked a restoring force acting on the free carriers induced by the exciton polarization field as the origin of the charge localization.

For the monolayer WS_2 , the ultrafast THz response of CVD-grown samples has been investigated and the results have been published in 2019 by S. Xu *et al.* and J. K. Gustafson *et al.* [152,153].

In Ref. [153] the first measurement of the THz photo-conductivity of monolayer WS₂ is reported. Here the authors, from the fit with a biexponential curve of the 15 ps long dynamics, obtained fast (hundreds of fs) and slow (few ps) components assigned to charge carrier loss induced by the exciton formation and Auger recombination plus surface trapping, respectively. The THz conductivity spectrum reported by S. Xu *et al.* shows a positive Re($\Delta \sigma$) accompanied by a small but negative Im($\Delta \sigma$), which points to a free carriers conductivity. The authors fitted this spectrum at different delays with the Drude – Smith model and obtained a localization parameter of c = -0.6, which indicates a relevant localization of the free carriers at the grain boundaries of the CVD-grown materials. The author also argued that the influence of excitons through their polarization field cannot be ruled out; however, if this were the case, the Drude – Smith model should have returned a *c* parameter around – 1, which would have led to a shape identical to the Lorentz oscillator.

In Ref. [152] the authors performed ultrafast THz spectroscopy of monolayer WS₂ at 20 K with a probe pulse covering the 0 - 8 THz spectral region and with pump photons resonant and non-resonant with the A exciton at \sim 2 eV. J. K. Gustafson *et al.* directly excited the A resonance and observed an increase of the conductivity that recovers within tens of ps. The authors argued that no free carriers can be generated using a sub-bandgap excitation and that the 1s - 2p intra-exciton transition falls outside the probe region. Therefore, the observed positive photo-conductivity can be solely ascribed to trions. This charged quasiparticle has an overall positive charge and forms through the coupling of excitons with a free holes population. This excess of positive charge in the valence band is photo-excited into trap states by the sub-band gap pump photon energy, which, as a result, creates free holes in the valence band [152]. By increasing the fluence up to 1.3×10^{14} photons/cm², the frequency-averaged THz photo-conductivity saturates, which suggests that defect-mediated processes are at play. The recovery of the frequency-averaged dynamics has been fitted with a bi-exponential curve that returns fast (hundreds of fs) and slow (few ps) components tentatively addressed to trion-phonon scattering and trion dissociation, respectively. Further evidence for the trion origin of the positive transient conductivity can be obtained by looking at the photo-conductivity spectra reported in Figure 11(b). Here the complex $\Delta\sigma$ presents a sharp and broad resonance at ~8 THz and ~4 THz, respectively, plus a non-vanishing trend when $\omega \rightarrow 0$. Therefore, the authors adopted a Drude-Lorenz model consisting of i) a Drude-term, describing the motion of the charged trion as heavy *free* – *carriers* under the influence of the THz field; ii) a Lorentz oscillator at ~4 THz accounting for the accumulation of an extra-charge (trions) at the grain boundaries of the material that gives rise to a restoring force acting on the other trions; iii) a Lorentz oscillator at ~7.4 THz (30 meV) that matches the trion binding energy. J. K. Gustafson *et al.* ruled out the picture proposed by X. Xing *et al.* about the polarization field sustained by excitons as the origin of the negative Im($\Delta\sigma$) and non-Drude conductivity as they demonstrated that charge-neutral exciton did not form in their experiment.

The results obtained with OPTP on multilayer and monolayer WS₂ can be rationalized by considering the different analyzed samples. X. Xing et al. and S. Xu *et al.* exploited the same THz probe (0.2 - 3 THz, 1 - 12 meV)but studied two different systems, the multilayer, and the monolayer WS_{2} , respectively. It is known that for TMDs the exciton binding energy largely blueshifts from the bulk to the monolayer case. Therefore, the indirect presence of excitons observed in [151] for the multilayer is compatible with the reduced exciton binding energy. In [153], the authors studied the ultrafast THz response of the monolayer WS₂ and it is reasonable that the presence of excitons and trions could not be detectable, not even indirectly. J. K. Gustafson et al., on the other hand, studied the monolayer in different experimental conditions as compared with S. Xu et al. Indeed, in [152], the authors used pump photons resonant with the excitons (no free carriers generation) and exploited a broader THz probe spectrum that covers the trions binding energy. Therefore, they were able to directly observe the spectral fingerprint of trions at \sim 7 THz and confirm the presence at room temperature of bound carriers.

In the family of Tungsten-based TMDs, the ultrafast THz response of bulk WSe₂ has been investigated by C. He *et al.* [154]. On one hand the authors reported frequency-averaged dynamics for both Re($\Delta\sigma$) and Im($\Delta\sigma$) in analogy with the WS₂ laminate [151]. On the other hand, they found different formations times for the two dynamics. As previously discussed, while the former is related to the contribution of free carriers, the latter reflects the presence of excitons. Free carriers are photo-generated via phonon-assisted processes and then undergo rapid cooling. After the hot carrier thermalization, excitons can be formed thanks to the binding energy which is greater than thermal energy at room temperature and then can dissociate after 700 fs. A system of rate equations has been developed considering the free carriers, defect-assisted, and phonon-assisted recombination pathways plus a term for free carriers generation via exciton dissociation. Concerning exciton dynamics, beyond the dissociation term, the rate equation includes the exciton recombination via phonon-assisted processes. The photo-conductivity spectrum of WSe_2 reflects the concomitant presence of free carriers and excitons. A Drude-Smith term considers the effect of back-scattering on mobile charge, while the Lorentz term describes the presence of excitons due to the effect of the polarization field on the localization of the free carriers.

3.2.1. TMDs heterostructures

VdW heterostructures (HS) are made up of vertically stacked TMDs monolayers. These systems, thanks to the weak VdW force between the layers, preserve the electronic band structure of the single constituents but acquire additional features. TMDs HS typically show a type-II band alignment with the valence band maximum and the conduction band minimum belonging to different layers. For TMDs HS after the photoexcitation, a rapid intralayer exciton dissociation occurs with the consequent charge separation and interlayer exciton formation. These quasiparticles are bound el-h pairs where the pristine carriers belong to different constituents and have lifetime orders of magnitude longer than intralayer ones making them promising candidates for those applications where an efficient charge extraction is needed (e.g. optoelectronic and light-harvesting).

S. Kumar *et al.* investigated the ultrafast THz response of a VdW HS composed of two semiconductive monolayers, namely CVD-grown MoSe₂ /MoS₂ HS. The authors observed an efficient charge transfer (\sim 90%) from the MoSe₂ to the MoS₂ layer [155] with the consequent formation of interlayer excitons. Frequency-averaged dynamics returned a manifold enhancement of the THz conductivity change and similar relaxation time for the photo-injected carriers in the single-layer and the HS (see Figure 12(a)) [155].

K. Lee *et al.* studied the THz response of a semiconductor/semimetal HS composed by CVD grown MoTe₂ and WTe₂ laminates in their 2 H and T_d phase, respectively [156]. By exploiting sub- and above-bandgap excitation and varying the fluence, the authors reported significantly shorter decay times for Re($\Delta\sigma$) in MoTe₂/WTe₂ as compared to the decays observed for the two distinct components. Remarkably, ground-state charge transfer (CT) and band gap renormalization were excluded as a possible origin for the faster recombination because i) the static conductivity (σ_{dc}) of the HS returned to be the summation of the two individual contributions; ii) the presence of a Schottky Barrier higher than thermal energy which hampers thermionic emission without photoexcitation. By following the dynamics of Im($\Delta\sigma$), the authors also invoked the presence of the exciton contribution in the recombination processes of the HS. From the analysis of Im($\Delta\sigma$) for the individual constituents and the HS as reported in Figure 12(b), it emerged a weak exciton contribution to the overall carrier relaxation for the



Figure 12. (a) Top panels: Differential THz reflectivity of monolayer $MoSe_2$ and of the heterostructure obtained with 1.55 eV pump photon energy; bottom panels: decay times and related amplitudes obtained from the fit with a bi-exponential curve of the ΔR (adapted with permission from ref [155]. © The Optical Society); (b) top-left panel static (red curve) and pumpinduced (blue curve) THz waveforms exploited for the experiments and top-right panel differential transmission through the heterostructure; bottom panels transient THz transmission signals of (bottom-left) monolayer WTe₂ and (bottom-right) monolayer MoTe₂ (adapted with permission from [156]).

semimetallic WTe₂, a strong intralayer exciton contribution for semiconductive MoTe₂, and a strong exciton contribution in MoTe₂/WTe₂, confirming that the charge species population was dominated by excitons. The overall sub-*ps* dynamics observed for the HS has been assigned to the formation of interlayer exciton via CT across the interface and their consequent dissociation via interfacial effects with a weak phonon contribution.

3.3. 2D perovskites

Metal halide perovskites (MHPs) are 'soft – lattice' ionic 3D semiconductors with a direct band gap with very interesting physical properties that promote them as a promising candidate for photovoltaics and light-emitting applications. The fascinating physical properties of MHPs are mainly due to the coupling of the anharmonic modes of the soft lattice with the photo-injected charge resulting in the formation of large polarons [157–160]. 2D MHPs consist of layers of inorganic Pb–I octahedra alternated with layers of organic cations. Thanks to the quantum confinement and the reduced Coulomb screening, 2D MHPs show a robust exciton population at room temperature with binding energy in the order of 200 - 300 meV [161]. Remarkably, these bound pairs are made of the photogenerated polaronic carriers, that are free carriers that bear the coupling with the lattice [162,163].

A first study on the carriers dynamics on 2D perovskites revealed the effect of excitons on the THz conductivity [164]. In details, R. L. Milot *et al.*



Figure 13. (a) and (b) Cartoon and frequency-averaged THz response of the bulk (left), intermediate (center) and 2D perovskite structure (adapted with permission from Ref. [164]; (c) photo-conductivity of 2D perovskites at different pump-probe delays together with the Drude-Lorentz fit (adapted with permission from Ref. [166]).

exploited OPTP spectroscopy to study the role of the reduced screening in determining the charge carriers recombination dynamics of 2D perovskites by varying the chemical composition of the system. Figure 13(a) shows the evolution from the bulk to 2D perovskite passing from an intermediate phase as a function of the phenethylammonium (PEA) content. The authors performed a fluence-dependent study of the frequency-averaged THz response of 3D, intermediate, and 2D samples, as shown in Figure 13(b). Here the author fit the $\Delta T/T$ dynamics with the following rate equation:

$$\frac{dn}{dt} = -k_1 n - k_2 n^2 - k_3 n^3 \tag{15}$$

where k_1 , k_2 and k_3 are the monomolecular, bimolecular and many-body decay rates, respectively. The global fit with Equation 15 of the data acquired at different fluences showed that a lower density of traps and a larger population of excitons are obtained, moving from the bulk to the pure 2D systems. Indeed, while for bulk-like and intermediate systems the monomolecular recombination rate constant k_1 ($\propto n$, where n is the carrier density) decreases indicating a lower trap density, for confined systems it starts to increase under the influence of the emerging exciton population. Moreover, bimolecular ($\propto n^2$) processes monotonically increase and become dominant up to the intermediate system due to the increase in the electronic density of states near the band edge, and then drop for the purely 2D MHP as the photon-to-charge branching ratio ϕ counteracts the rise in k_2 in the measured quantity ϕk_2 because of the exciton formation. A similar trend has been observed for the Auger ($\propto n^3$) recombination rate ϕk_3 , whose increase has been attributed to stronger Coulomb interaction. This trend is inverted for the purely confined system because of the decrease of ϕ .

The presence of excitons in 2D MHPs has been then demonstrated by exploiting broadband THz generation with a THz pulse covering the 3 - 16THz (15 - 65 meV) spectral range [165]. The sample analyzed in this work is PEA₂PbI₄ which shows an excitonic peak at \sim 510 nm and the band-edge onset at ~450 nm. The authors observed different dynamics by directly pumping the excitonic resonance or pumping at the band edge. Indeed, if large excess energy is transferred to the carriers, then they need to thermalize and condense to reach equilibrium. After this rapid cooling and the formation of excitons, the frequency-averaged dynamics follows the same recombination dynamics and recovers within 100 ps. Concerning the conductivity spectra, A. Burgos-Caminal et al. observed a non-Drude response composed by a positive and increasing $\text{Re}(\Delta \sigma)$ accompanied by negative Im $(\Delta \sigma)$. Invoking multiple intra-excitonic transitions, the observed transient conductivity has been modeled by a Drude-Smith term that describes the motion of free carriers under back-scattering events plus a Lorentz oscillator that accounts for intra-excitonic transitions. Remarkably, the global fit analysis returned that the density of free carriers has an instantaneous formation time peaked \sim 500 fs after the photo-excitation, and then rapidly decreases to a stable value of 50% of its maximum value. At the same time, the excitonic resonance takes \sim 500 fs to stabilize to its maximum value, confirming the dynamics extracted via the analysis of the frequencyaveraged response.

Together with the work of A. Burgos-Caminal et al., G. Folpini et al. obtained an indirect confirmation of the concomitant presence of free carriers and exciton in 2D perovskites. In [166], the ultrafast THz response of NBT₂PbI₄ was investigated in the 0 - 3 THz spectral region. The ultrafast response of the sample showed a buildup of the signal within the instrument response function followed by a fluence independent recovery within the first ~ 10 ps. Here the data were fitted with a rate equation that consider bimolecular processes only, as linear and cubic terms in the carrier density (trap-assisted and Auger processes, respectively) resulted to be negligible, suggesting that the signal arises from the minority free carriers that do not contribute to the sub-ps formation of excitons. Differently from the data reported in [165], in [166] the narrower THz probe is not directly sensitive to the exciton resonance, hence returning the contribution of intraband absorption only. This picture was supported by analyzing the spectral THz response of the sample, reported in Figure 13c. Firstly, the pump-induced conductivity confirmed the polaronic nature of the free carriers due to the presence of phonon signatures, in analogy with the 3D bulk counterpart [160]. This confirms the exciton-polarons pictures proposed by S. Neutzner et al by exploiting linear and non-linear optical spectroscopy [162]. Moreover, $\operatorname{Re}(\Delta\sigma)$ vanishes for $\omega \longrightarrow 0$ and monotonically grows with the increase in frequency, while $Im(\Delta \sigma)$ is negative at all frequency. This points to the presence of a restoring force acting on the polaronic carriers, as for the TMDs case discussed in Section 3.2. The conductivity spectra were thus fitted with a combination of a Drude term plus a Lorentz oscillator to describe the motion of the free carriers under the influence of the restoring force originated by the polarization field sustained by excitons playing the role of an ensemble of interacting dipoles.

4. Perspectives

4.1 THz near-field imaging

A major challenge to the investigation of the THz response of van der Waals 2D materials is the long free-space wavelength of THz radiation that, due to diffraction, severely limits the achievable spatial resolution (0.3 mm at 1 THz, according to the Rayleigh criterion), and imposes constraints to the sample size, hindering the investigation with far-field optics of few-microns samples, like those prepared by exfoliation, or prevents the study of una-voidable spatial inhomogeneities at the nanoscale.

Near-field microscopy [167–174] represents a powerful tool for circumventing diffraction limits and investigate the infrared light–matter interaction in the far infrared, in graphene [175–177] and other 2D materials [178– 181]. Among the different near-field probing schemes developed to date to probe the THz optical response, apertureless, scattering-type scanning nearfield optical microscopy (s-SNOM) achieves the highest spatial resolution, which is wavelength-independent and reaches the tens of nanometers range [182–184], corresponding to field confinements as large as $\lambda/4600$ [185].

Near-field approaches have allowed mapping charge carrier density distribution with unprecedented spatial resolution [186-189], exploiting the inherent sensitivity of THz fields to free-carriers. Broadband THz sources, such as TDS (covering the range 0.4-1.8 THz) [184,190], free-electron lasers [191], multimode THz frequency combs [192], and synchrotron radiation [193,194], allow the characterization with spatial and spectral control, while the 3D confinement of THz fields provides the tomographic sensitivity for retrieving depth-dependent dielectric properties, as recently done with thin films of topological insulators, in the mid-infrared [195,196] and THz ranges [197,198]. THz near-field experiments applied to investigate lowdimensional materials includes also THz control of photoemission [199] and THz photocurrent nanoscopy. The latter directly gives access to the transport properties of the material and is based on the detection of the currents arising from local near-field photoexcitation. It was used to reveal acoustic plasmons in graphene [177] and to identify the photodetection mechanisms in semiconductor nanowires [200], with straightforward implications for the development of efficient room-temperature THz detectors.

Moreover, THz s-SNOM enables to probe collective excitations in 2D materials [201–204] exploiting the strong light confinement achieved with light compression into sub-diffractional volumes, to access regimes of strong light-matter interaction in which THz photons can couple with material excitations to form polaritons, which are quasiparticles composed of a THz-photon and an oscillating charge [201] that can be either free charge carriers within metals and doped-semiconductors (plasmon polaritons) or ionic charges on a polar crystal lattice (phonon polaritons, PPhs) [197,198]. Most importantly, s-SNOM allows to map in the real space the propagation of THz polaritons giving access to their energy dispersion relation and damping channels [191,205], as recently done in graphene [177,206] and topological insulators [197,198]. The THz polaritons supported by 2D materials, offer unique opportunities for extreme THz light confinement at the atomic limit with groundbreaking perspectives for the development of ultra-sensitive plasmonic devices [207] and tailored nanophotonics [208].

Ultrafast studies exploiting the near-field approach to study photoinduced dynamics in 2D materials with nanoscale spatial resolution can play a pivotal role in guiding the development of nanotechnologies in the THz range. Pump-probe scheme with NIR pump and mid-IR probe pulses coupled to a s-SNOM has allowed investigation of the ultrafast local photoconductivity of graphene [209], insulator-metal phase-change materials [210], transient surface plasmons in graphene [211] and interface polaritons in black phosphorus on SiO₂ [178].

At THz frequencies, sub-picosecond charging dynamics of single nanoparticles was resolved by coupling a few THz field to a scanning tunneling microscope for electrical detection of local currents [212]. However, the first demonstration of an all-optical time-resolved near-field experiment in the THz range was presented in 2014, using a near-field pulse as pump and a multi-THz pulse as probe, detected by EOS, to monitor the transient dielectric properties of semiconducting InAs nanowires [213] with sub-cycle temporal resolution and nanometer tomographic sensitivity, see Figure 14.

More recently, ultrafast THz nanoscopy has been applied to investigate 2D materials [179,214] using time-resolved THz polarization nanoscopy [179,214] and laser terahertz emission nanoscopy [214], see Figure 14(b,c).

The former is based on time-resolved detection of the pump-induced local THz emission, which is outcoupled by the s-SNOM tip and electrooptically sampled in the far-field. Applied to WSe_2/WS_2 heterostructure, it has allowed to detecting the 1 ps THz emission generated by the out-ofplane current associated with interlayer exciton transfer. The latter is based on the measurement of amplitude and phase change of a tip-scattered THz pulse and it has been applied to interrogate excitons of TMDs. The THz field concentrated beneath the tip extends out-of-plane and is able to displace



Figure 14. (a) Schematics of the experimental set-up for ultrafast THz s-SNOM from Ref. [213] used to measure THz nano-movie frames of nanowire photoexcitation (maps at the bottom), using NIR (1.56 μ m) pump pulses and multi-THz probe pulses, focused onto the atomic force microscope (AFM) tip of a s-SNOM, detecting the scattered electric near field by EOS or with a MCT photodiode to perform FTIR spectroscopy; (b) Schematic of OPTP near-field experiments on a WS₂/WSe₂ vdW heterostructure interrogating photo-generated e–h pairs (blue and red spheres) using a THz probe pulse, detected as a function of the pump–probe delay t_p, see Ref. [214], together with a schematic of the type-II band alignment of the heterostructure and the tunneling of charge carriers through the vdW gap. The map shows the magnitude of the pump-induced changes in characteristic lifetime τ . (c) Ultrafast THz polarization nanoscopy of monolayer and bilayer to study exciton Mott transition from Ref. [179], together with the carrier density dynamics in bilayer WSe₂ for increasing pump fluence, extracted from the peak of the pump-induced change to the THz scattered field.

photoexcited excitons [214], depending on their polarizability. Interestingly, THz field is sensitive to the entire exciton population, differently from interband techniques only probing the excitons that obey specific selection rules, such that this technique has allowed to track the Mott transition of TMDs excitons into electron-hole plasma in monolayer and stacked twisted bilayer WSe₂ [179].

These pioneering near-field studies highlighted the crucial role of nanoscale inhomogeneities in defining the ultrafast THz response of 2D materials, *e.g.* the spatial variation of Mott transition threshold leads to the inhomogeneous broadening observed by spatially averaged techniques [179]. Many promising research opportunities still have to be explored, including the study of the optical control of THz polaritons and electronic waves with important perspectives for ultrafast plasma electronics, as recently demonstrated with black-phosphorus-based heterostructure, where mid-IR plasmon waves have been activated with ultrafast NIR light with femtosecond switching times [178]. Therefore, ultrafast THz nanoscopy offers great opportunities to expand our understanding of the optoelectronic properties of 2D materials in a path towards the design of efficient nanotechnologies for the detection, modulation and emission of THz radiation, especially of ultrafast components like ultrafast switch, saturable absorbers and amplitude and phase modulators. On the other hand, it appears the ideal tool for probing local quantum phenomena in low-dimensional materials.

In conclusion, in this review we reported the most recent results obtained on 2D materials by exploiting ultrafast OPTP spectroscopy. We highlighted the effectiveness of THz-based investigation techniques in addressing ultrafast processes in 2D materials, which are at the base of their optical response over a broad spectral range.

OPTP measurements of graphene have revealed the efficiency of the collective response of Dirac electrons in converting THz fields into electronic heat, giving rise to broadband strong THz nonlinearities, to be exploited for nonlinear wave conversion and efficient THz high harmonics generation [137]. Moreover, OPTP spectroscopy has allowed to follow the picoseconds cooling dynamics of the hot charge carriers which enables a number of applications such as ultrafast broadband detection of THz light by photo-thermoelectric and bolometric effects [75], ultrafast THz modulation [76,77] and saturable absorption [78] for THz pulses generation. The investigations of the cooling dynamics of graphene hot carriers as a function of different control knobs, such as defects density, doping and dielectric environment, has unveiled a large tunability ultimately constrained by the strong coupling of the hot electrons to graphene optical phonons.

In TMDs, studies by OPTP has targeted the dynamics of both charge carriers and excitons, the latter being associated to strong absorption resonances in the VIS. For TMDs, the interplay between free carrier and excitons emerged to be not only layer-dependent, but also strongly influenced by the growth protocol and doping.

The technological relevance of the observed ultrafast processes is enhanced by the compatibility of 2D materials with existing electronic technologies, like CMOS-based electronics. Accordingly, a large number of applications to generate, detect and manipulate THz waves has been envisioned ranging from telecommunications, based on the ability of THz frequency carriers to enable higher data-rate communication for next-generation technologies, to gas and bio-sensing, based on both the extreme THz confinement enabled by waveguided modes, plasma waves and polaritons, and on the exploitation of the THz molecular fingerprints.

Disclosure statement

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ORCID

Eugenio Cinquanta http://orcid.org/0000-0002-4721-5215 Eva Arianna Aurelia Pogna http://orcid.org/0000-0003-4779-3549 Lorenzo Gatto http://orcid.org/0000-0002-7195-4317 Salvatore Stagira http://orcid.org/0000-0002-8457-3185 Caterina Vozzi http://orcid.org/0000-0002-0212-0191

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