

Mercury Chalcogenide Quantum Dots: Material Perspective for Device Integration

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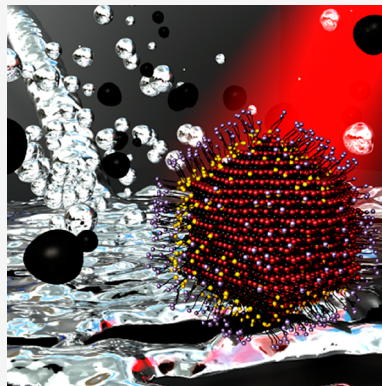
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ABSTRACT: Nanocrystals (NCs) are one of the few nanotechnologies to have attained mass market applications with their use as light sources for displays. This success relies on Cd- and In-based wide bandgap materials. NCs are likely to be employed in more applications as they provide a versatile platform for optoelectronics, specifically, infrared optoelectronics. The existing material technologies in this range of wavelengths are generally not cost-effective, which limits the spread of technologies beyond a few niche domains, such as defense and astronomy. Among the potential candidates to address the infrared window, mercury chalcogenide (HgX) NCs exhibit the highest potential in terms of performance. In this review, we discuss how material developments have facilitated device enhancements. Because these materials are mainly used for their infrared optical features, we first review the strategies for their colloidal growth and their specific electronic structure. The review is organized considering three main device-related applications: light emission, electronic transport, and infrared photodetection.



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1. INTRODUCTION

This review is related to the merging of two fields, specifically, infrared (IR) sensing, in which mercury chalcogenides have been widely applied (see Figure 1), and colloidal semiconductor nanocrystals (NCs), which are nanometer sized crystals of semiconductors. For a long period of time, these two fields evolved independently, and they started to merge only 10 years ago^{1–3} owing to the increase in the cost constraints of IR detection and the maturity of the growth and material processing techniques of NCs (toward conductive films).⁴

The infrared range can enable individuals to see in the dark in a fully passive manner (seeing without being seen) through thermal imaging. For wavelengths above 3 μm , the blackbody emission signal starts to prevail over reflection from external sources, owing to which the objects and bodies become primary light sources. The emitted radiation from these objects can be observed with a suitable (*i.e.*, wavelength matched) detector.

The first IR sensors that emerged in the 1950s were mostly based on lead chalcogenide materials, whose bulk bandgap was excessively large to realize thermal detection, as shown in Figure 1. Many candidate materials were investigated, and InSb, among the emerging III–V semiconductors, appeared to be one of the most promising material for detection in the 3–5 μm range (midwave infrared: MWIR). Nevertheless, the actual targeted

spectral range was 8–12 μm , which matched the maximum emission for a blackbody at room temperature as well as the atmospheric transparency window. This spectral range corresponds to a transition energy of 120 meV, requiring very narrow bandgap semiconductors. Moreover, a series of materials, including gray tin, HgTe, and HgSe, were investigated; however, these materials appeared to be semimetallic in their bulk form, resulting in inferior or nonexistent photoconductive properties. In 1959, HgCdTe alloys⁵ emerged as photoconductive materials with a performance comparable to that of InSb in the MWIR. In such alloys, bandgap tuning could be realized by adjusting the Cd to Hg ratio. With a 20% content of Cd, the cutoff wavelength of HgCdTe was approximately 12 μm , and the material exhibited a reasonable performance for low temperature operation. Thus, the development of HgCdTe was driven by its application (*i.e.*, passive imaging for military). At present, the motivation for this early application may appear paradoxical because HgTe/CdTe quantum well structures regained interest in academic laboratories owing to their specific electronic structure that makes such materials topological insulators. Nevertheless, this military driven application has rendered HgCdTe as one of the most investigated semiconductors, even if it notably lags Si and GaAs. To date, in the family of infrared semiconductors (InGaAs, InSb, HgCdTe) and heterostructures (QWIP and type II superlattices), HgCdTe remains the leading material for IR (3–12 μm) detection, offering a wide spectral range tunability from visible to longwave infrared, while always matching the highest performance for each spectral range.

Since 1993, monodisperse syntheses of colloidal NCs are introduced through the hot precursor injection method.⁶ This process allows the solution growth of semiconductor particles to generate quantum confined materials with an atom like spectrum. For a long time, colloidal syntheses have experienced a lack of conviction from the semiconductor community, owing to the lower control and carrier mobilities associated with these approaches than those attained using conventional methods. Nevertheless, considerable progress has been made over the last 30 years in terms of the size, shape,^{7,8} and composition control. At present, half of the periodic table can be obtained under the NC form. The key achievements include the introduction of core–shell colloidal heterostructures,⁹ as this approach can, not only achieve nanoparticles with a photoluminescence (PL) yield of nearly 100% but can also facilitate the maturity of the whole field. Owing to such developments, NCs have become one of the few representative nanotechnologies that have attained mass market status. In the last five years, NCs have been used as green and red downconverters for LCD displays,^{10,11} thereby increasing the gamut (*i.e.*, range of color) compared to that achieved using previous technologies based on lanthanide emitters. In addition to the commercial value of this market (1 billion \$ per year^{12,13}), this display application has demonstrated the viability of NCs to be included in industrial processes while enabling the realization of devices with a long lifetime (>10 y of operation). The switch from optically excited NCs to electrically pumped devices to reduce electrical consumption and enhance image contrast is expected to be the next challenge in this direction.

The introduction of mercury chalcogenide (HgX) NCs was motivated by the concept of extending the optical feature of NCs into the red and near-infrared ranges,¹⁴ as shown in Figure 1. The first report of HgX-based NCs corresponded to a CdS/HgS heterostructure.¹⁵ At this stage, type I core–shell^{9,16} colloidal heterostructures used to enhance the PL efficiency had not yet

been reported, and the method used to grow the heterostructures (*i.e.*, the cation exchange method) was not an established synthetic method^{17–19} to control the NC composition. The first targeted application of HgX NCs was their use as broadband optical amplifiers in the telecommunication range at wavelengths between 1.3 and 1.5 μm . Specifically, HgTe NCs²⁰ were first used in the telecom domain in the early 2000s. Because NCs are continuously spectrally tunable, they offer higher flexibility for the design of IR emitting devices than their III–V counterparts obtained using epitaxy, for which the substrate–lattice matching constrains the bandgap tunability. Despite the development of HgX NCs, as reviewed in the following sections, this first targeted application remains undemonstrated, although certain recent reports^{21–23} indicate that HgX NCs are promising materials to obtain infrared light sources.

Moreover, considerable progress has been made in the control of the surface chemistry of NCs^{4,24–26} to obtain large carrier mobility films ($\mu \approx 1 \text{ cm}^2 \cdot \text{V}^{-1} \cdot \text{s}^{-1}$), and the growth of HgX NCs with larger absorption wavelengths^{3,27–30} has led to the emergence of HgX NCs as a viable building block for low cost IR sensing. The basic concept of using HgTe NCs instead of HgCdTe alloy is to replace the bandgap tunability from the mixed composition to quantum confinement. For HgTe, which is a semimetal, the absorption can be tuned from 1.5 eV for the most confined form of HgTe^{31,32} to virtually zero. In practice, an absorption feature of as low as 20 meV has been reported.³⁰ In a ten-year time scale, the material has switched from seminal development with absorption and photoconduction in the short wave IR (SWIR) and MWIR^{1,3} to a versatile platform for optoelectronics. A nonexhaustive list of key developments include

- Evidence of an intraband in the steady state due to self-doping^{29,33,34}
- Large mobility films³⁵ with $\mu > 1 \text{ cm}^2 \cdot \text{V}^{-1} \text{ s}^{-1}$ ³⁶
- Large absorption^{37–40} and emission²¹ from NC films due to coupling with structures inducing a light resonance
- Multicolor detectors^{41–47}
- High detectivity devices that, for single pixels, are comparable with commercial setups⁴⁸
- Focal plane arrays in the short wave^{49,50} and midwave infrared^{51,52}

This field has likely reached a transition point from being a niche topic studied by a few groups, mostly with an NC background, to becoming a field of interest for other researchers in the photonics, μ -fabrication, and optoelectronics fields,⁵³ enabling the development of increasingly complex devices. The expansion of this field also extends to industry (SWIR Vision, IMEC, Qrv, and Emberion are working on PbS NC-based imaging setups and QDIR, New Imaging Technologies are focusing on HgTe to expand the spectral range), leading to the emergence of a viable low cost technology for IR sensing. Such an extension of this material toward a broader community compels a review aggregating the material insights and technological developments. The scope of this review is on mercury chalcogenide (HgX: HgS, HgSe, and HgTe) NCs, not on infrared NCs. In particular, the concept and performances of HgX NCs are compared with those of PbS-based NCs. However, PbS(e) is not the focus of this review. Several reviews dedicated to HgX NCs have been written. For example, Green *et al.*⁵⁴ focused on the synthetic aspect of NCs, while Rogach *et al.*⁵⁵ considered a historical perspective in their review. The

detection application is not described extensively in these documents. Several short reviews have been dedicated to HgX NCs, with a strong focus on the aspects of devices^{56,57} or the fundamental limits that can be achieved;⁵⁸ however, none of these reviews covers all the aspects from the material growth to application while discussing the electronic structure and transport properties. It is a choice of the authors not to include biological applications of the HgX NCs. Even though certain researchers have reported on the use of HgX NCs for biolabeling in the near IR,⁵⁹ the presence of Hg has been considered detrimental for biological applications, and thus, this direction has not been extensively considered.

The present review is organized into six main sections. The first section is dedicated to NC synthesis. A sub-section is dedicated to each of the chalcogenides (S, Se, and Te). As shape and dimensionality control is an emerging direction for HgX NCs, a section is dedicated to 2D nanoplatelets composed of HgX. The subsequent section discusses the challenging growth of core–shell objects based on HgX NCs. The final part of this chapter deals with the doping⁶⁰ and tunability of the carrier density. This section is of utmost importance for the discussion of optical properties (for example, the intraband absorption), transport properties (occurring at a considerably higher carrier density compared to that for other II–VI materials, to the point at which semiconductor to metal transition can occur), and device design (*i.e.*, p–n junction formation).

The second section deals with optical properties, starting with the bulk dispersion and modifications introduced by quantum confinement. Certain results of the modeling of the electronic structure are presented: in addition to the sizing curve (*i.e.*, energy *vs* size), multiband k·p model, tight binding, and DFT have facilitated the understanding of properties such as the fine structure of the conduction band,⁶¹ the origin of the stimulated emission,²² and the inverted temperature dependence of the HgTe absorption spectrum.⁶² A subsection is focused on the optical properties of the NCs, before we discuss the absolute band alignment (*vs* vacuum) of the HgX NCs. This section is critical in terms of two key aspects, specifically, the design of colloidal heterostructures and NC-based device integration. A rational design of devices requires a deep and reliable understanding of the band alignment to move beyond empirical rules. Nevertheless, this aspect is challenging because HgX NCs combine an inverted band ordering, quantum confinement, and surface chemistry dependence of their spectra, which prevents the textbook design of heterojunctions. Lastly, we dedicate a section to the carrier dynamics, wherein we discuss the specific experimental setup used to examine the relaxation of narrow bandgap semiconductors with optical features in the IR.

After defining the materials and their fundamental properties, we focus on material applications for light emission and detection. In the subsequent section, we review the light emission properties of the HgX NCs. We first discuss the interband and intraband photoluminescence and later describe recent progresses related to stimulated emission and electroluminescence.

Section 5 focuses on the transport properties of HgX NC arrays, providing a basis for the subsequent section that reports on the photoconductive properties of these materials. As high carrier densities and degenerate doping are key properties of HgX compounds, we first discuss the influence of the carrier density on transport. Moreover, a section is dedicated to the specific surface chemistries developed for HgX NCs to realize simultaneous increases in the carrier mobility and stability. The

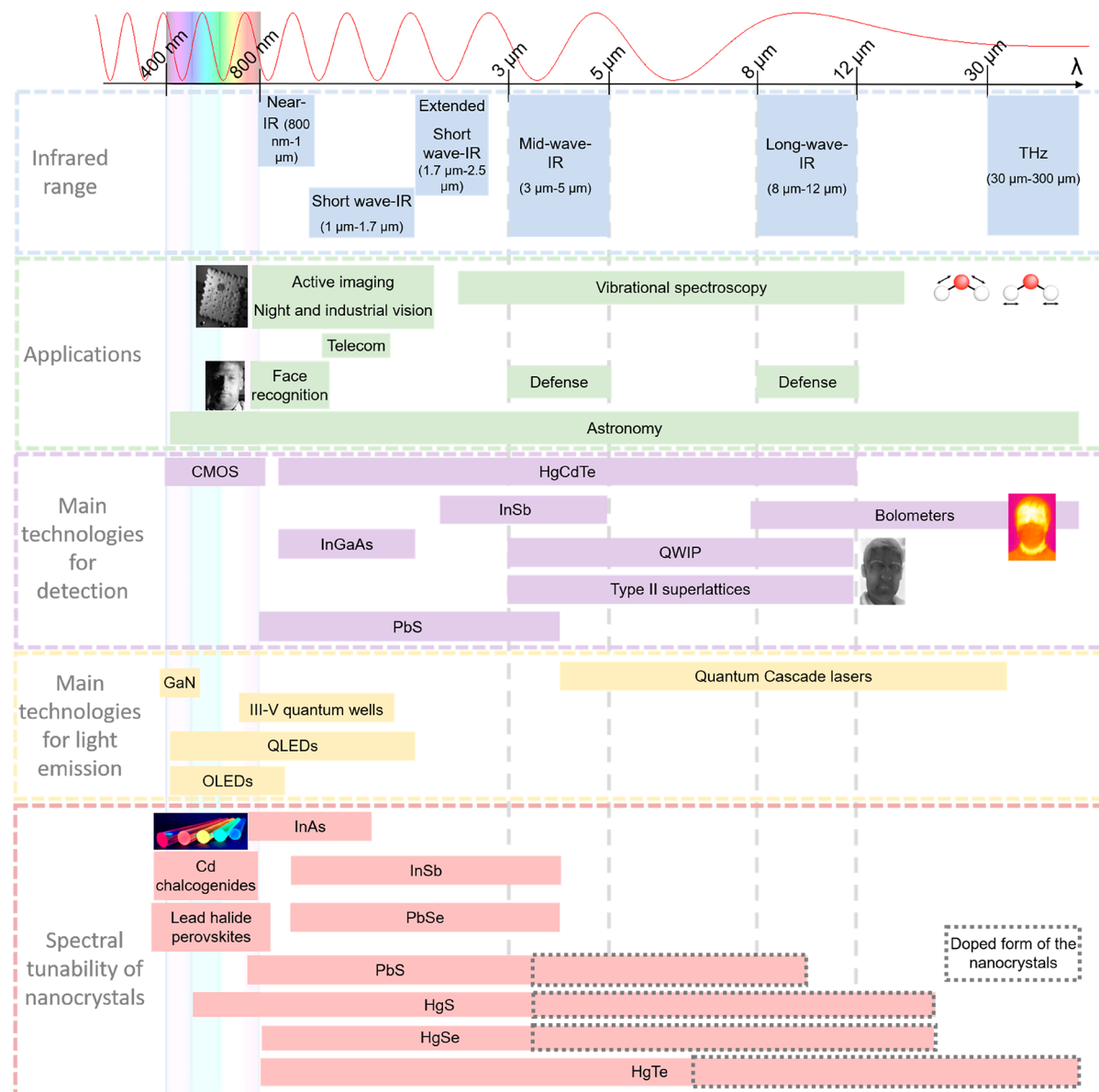


Figure 1. Electromagnetic spectrum highlighting the different parts of the infrared range. The main applications, sensor and source technologies, and tunability range for the nanocrystal optical features are also illustrated.

latter aspect is examined in depth. Finally, a section is dedicated to field effect transistors based on HgX NCs.

In the final section, we discuss the IR photoconduction and photodetection properties of HgX NC arrays. A section is dedicated to the specific figures of merit of IR detection, such as the spectral range, response time, responsivity, noise, signal-to-noise, and operating temperature. We discuss the experimental determination of these figures of merit and the specificities of IR NCs. The following section is dedicated to devices for IR detection based on HgX NCs as the active material. To avoid presenting an exhaustive list of devices, the devices are categorized based on their specific geometry. We start with photoconductive devices and later include certain refinements of this basic geometry, including the addition of gates for the design of phototransistors, the use of flexible substrates, intraband detection, and the coupling with 2D materials. Subsequently, we review photovoltaic devices, which exhibit the highest performance at present. A subsection is dedicated to the key upgrades

such as multicolor sensing, devices with enhanced light–matter coupling, and devices to realize imaging (*i.e.*, focal plane arrays).

The conclusion summarizes the main developments achieved over the past 10 years, and we highlight the main directions that remain to be addressed: material optimization, enhanced understanding of the material characteristics, and upgraded devices to facilitate the propagation of the technology in the market.

2. MATERIAL SYNTHESIS

This review first discusses the growth of NCs to be used as building block for devices. Mercury chalcogenides are II–VI semiconductors and are thus natural candidates to extend the optical features of cadmium chalcogenide nanocrystals, which are the most investigated material in the colloidal nanocrystal form. Because mercury is located below cadmium in the periodic table, the switch from Cd to Hg is associated with an increased number of electrons and a reduced bandgap. Consequently, mercury chalcogenides are a natural candidate to address the

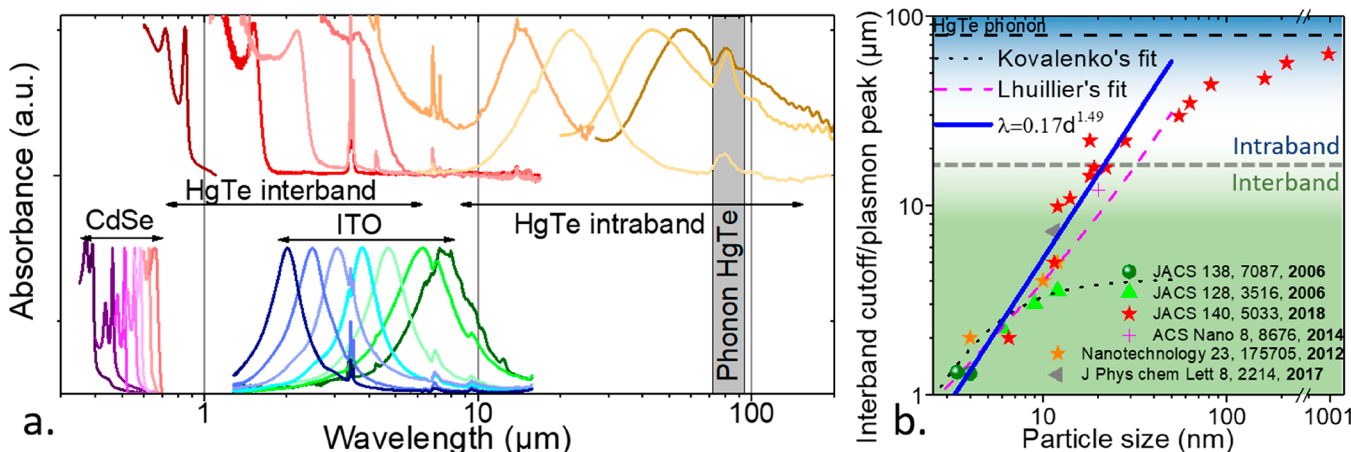


Figure 2. (a) Absorption spectra for various sizes of HgTe and CdSe and various doping values of ITO NCs. (b) Wavelength of the lowest energy absorption feature as a function of the particle size for HgTe NCs, obtained from various references.

infrared range, as discussed later. Surprisingly, the field of HgX NCs did not begin with the growth of core only nanoparticles with a spherical shape but instead through the growth of heterostructures by using a cation exchange procedure at a stage in which this procedure was not a conventionally accepted synthetic tool to grow NCs.¹⁵ This aspect led to the formation of HgS/CdS core–shell particles with absorption in the red domain. Nevertheless, it was promptly observed that core only structures offered an easier strategy to achieve such NCs.²⁰

Even though the above-mentioned materials are II–VI semiconductors, their growth differs significantly from that of Cd-based materials. In particular, colloidal synthesis of these materials is generally realized at a considerably lower temperatures (typically, 100 °C), indicating a highly different reactivity of HgX-based compounds compared to that of their Cd-based counterparts. Among mercury chalcogenides,⁵⁴ HgTe has attracted a large amount of interest, in contrast to cadmium chalcogenides, as the ease of oxidizing CdTe renders this compound more difficult to grow and study than CdSe or CdS. The vicinity of HgTe to the HgCdTe alloy, which is already used for IR detection, has facilitated the development and understanding of this compound. Colloidal growth of HgTe has attained a high level of maturity with unique size tunability (from few nanometers to micrometers),³⁰ high monodispersity, and shape control.^{31,63,64} The two other chalcogenides, HgS and HgSe, have been less investigated; however, the recent emergence of intraband transition in these materials may change this trend.^{65,66} Among the HgX compounds, HgS exhibits two specificities. First, the bulk material presents a gap, while HgSe and HgTe are semimetals in the bulk form. In addition, HgS exists in two crystalline forms: zinc blende and cinnabar. The latter form is promising to obtain chiral optical features.

This section is organized as follows: we first review the growth of core-only objects for the three chalcogenides. Then, we discuss shape control relative to the growth of HgX NCs with a focus on the growth of 2D nanoparticles (nanoplatelets)⁶⁷. A section is dedicated to HgX-based heterostructures. In addition to the traditional enhancement of the photoluminescence efficiency, the growth of the heterostructure is a key step to obtain air stable and temperature resistant NCs. Finally, we discuss the doping of HgX NCs, which is a central aspect for device integration. In this section, we review not only the

inherent doping after synthesis but also the strategies use to tune the doping magnitude through physicochemical methods.

2.1. HgTe Nanocrystal Synthesis

Among the three chalcogenides (HgS, HgSe, and HgTe), HgTe is the most studied from the synthesis and application viewpoints, which may seem paradoxical from the material growth perspective because Te-based chalcogenides are less available than their S and Se counterparts. In addition, Te-based semiconductors usually present a lower work function,⁶⁸ making them more prone to oxidation. HgTe in its bulk form has been well developed for infrared detection for over 50 years.⁶⁹ This application has encouraged the development of growth methods and has facilitated the understanding of the HgTe electronic structure.^{70–72} In particular, HgCdTe alloys exhibit broad tunability of the absorption spectrum from the near IR to longwave IR. Photocarriers exhibit a large lifetime of up to 1 μs , which is favorable for the charge collection in IR sensors, and the dark current in this material is among the lowest for IR detectors,^{73,74} enabling higher operating temperatures. Although this material is toxic, until now, the performances recorded with HgTe-based materials are higher than those of any other material. This aspect is more notable for NCs dedicated to IR. HgTe exhibits the highest spectral tunability (from below 1 to above 100 μm), as shown in Figure 2a. In comparison, quantum confinement allows the tuning of the CdSe band edge from only 400 to 700 nm, while doping in ITO (tin doped indium oxide) NCs only shifts the absorption from 2 to 9 μm .^{75,76} The tunability in both cases is considerably small (less than the 2 orders of magnitude) compared to that achieved by HgTe.

Until now, the use of less toxic compounds in IR applications has led to a dramatically reduced performance.^{76–79} As a bulk material, InSb exhibits a similar performance as that of HgCdTe in the 3–5 μm range; however, the NC counterpart is not as widely investigated as HgTe. InSb NCs are promising materials;^{80–82} however, their integration in devices remains unexplored. Under NC form, lead chalcogenides are the most notable competitors to HgTe.⁴⁹ Among these materials, PbS is the most developed, exhibiting a high synthesis maturity,^{83,84} tunable absorption in the near IR, mature transport,⁸⁵ and device integration.^{86–88} However, the material becomes challenging to manage above 2 μm ⁸⁹ and is limited by its bulk bandgap ($\approx 4 \mu\text{m}$).⁹⁰ Owing to these considerations, HgTe NCs

Table 1. Conditions for the Selection of Relevant Syntheses for HgTe NCs

Hg precursor	Te precursor	ligand	solvent/temperature	method	size (nm)	optical prop	year/ref
Hg(ClO ₄) ₂	H ₂ Te	thioglycerol	water/RT		3–6	up to 1.4 μm	1999 ²⁰
Hg(ClO ₄) ₂	H ₂ Te	thioglycerol, thioglycolic acid, L-cysteine, mercaptoethanol, and mercaptoethylamine	water/RT and then heating at 80 °C		3–12	up to 3.7 μm	2006 ²⁷
HgCl ₂	H ₂ Te	triethylamine, tripropylamine -furanmethanethiol	DMSO or DMF and diphenylether/40–80 °C		4–10	up to 4.5 μm	2020 ⁹²
HgBr ₂	TOP:Te	TOPO	TOPO/70–100 °C	hot injection	3–17	above 1.7 μm	2013 ⁹³
Hg(OAc) ₂	TOP:Te	ethanol	ethanol/−78 °C	low T injection	3–4	up to 1.6 μm	2006 ⁹⁴
Hg(OAc) ₂	Te	pyridine	butanol/0–90 °C	injection of cation on anion	5–12	up to 5 μm	2011 ³
HgCl ₂	TOP:Te	DDT	OLA/60–120 °C		5–12	up to 5 μm	2011 ⁹⁵
HgCl ₂	(TMS) ₂ Te	DDT	OLA		5–12	up to 5 μm	2017 ⁶³
HgX ₂ (X = Cl, Br, I)	TOP:Te	DDT	OLA/60–300 °C	injection of both cations and anions in a hot solvent	5–1000	2–60 μm	2018 ³⁰
liquid Hg	TOPTe	DDT	OLA/60–150 °C	<i>in situ</i> formation of Hg thiolate	3–20	1.3–15 μm	2020 ⁹⁶

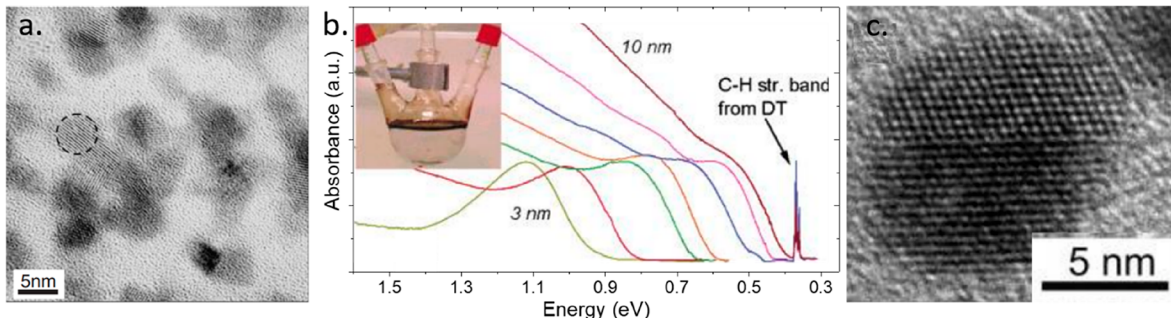


Figure 3. (a) High resolution TEM image of HgTe NCs obtained through the synthesis described in ref 20. (a) Adapted from ref 103. Copyright 2005 Japan Society of Applied Physics. (b) Representative room temperature absorption spectra of DDT-capped HgTe NCs in CCl₄. (c) TEM image of ≈9 nm DDT-capped HgTe NCs emitting at 3 μm. (b,c) Adapted with permission from ref 27. Copyright 2006 American Chemical Society.

represent a highly versatile platform for IR photonics and optoelectronics. Initially (2000–2010), the interest in HgTe NCs was not driven by IR detection as for the bulk case but rather by light amplification.⁵⁵ Only later (post 2006) did HgTe NCs emerge as a promising low cost alternative^{3,91} to bulk HgCdTe alloys.

Although HgX NCs are II–VI semiconductors, as is CdSe, their synthesis involves certain specificities. For example, the synthesis temperature is significantly reduced compared to that of CdSe NCs, as indicated in Table 1. The synthesis strategies used to obtain HgTe NCs can be categorized as the aqueous method, which has been extended to polar solvents, and the organic medium method.

2.1.1. Synthesis in a Polar Solvent. Very early reports on HgTe NCs and their alloying with HgCdTe can be found. These concepts were likely motivated by the realization of polycrystalline films by using chemical bath deposition. By mixing mercury halides with sodium telluride in methanol, large (1–100 nm) nanoparticles of HgTe and HgCdTe were obtained by Müllenborn *et al.*⁹⁷ This research was followed by a period of low interest in HgX NCs, likely corresponding to the period in which CdSe NCs became popular. Rogach *et al.*, inspired by their earlier work on CdTe NCs,^{98,99} developed a water-based synthesis approach for HgTe NCs.^{20,100} Mercury perchlorate was dissolved in water in the presence of thioglycerol as a ligand, and H₂Te was bubbled in this solution, acting as a Te²⁻ source. The reaction could be more effectively performed under alkaline conditions (pH = 9–12, typically). The obtained HgTe NCs

exhibited a tunable size from 3 to 6 nm, as shown in Figures 3a and 26a in terms of transmission electron microscopy (TEM) images and optical spectra, respectively. Synthesis was conducted at room temperature, and the NC emission reached the telecom range. Later, Kovalenko *et al.* introduced an additional Ostwald ripening step by heating the solution after growth at 80 °C.²⁷ Larger particles, 10–12 nm (Figure 3b,c), were obtained, and the PL signal reached 3.7 μm, as shown in Figure 26b. More recently, Rogach and co-workers expanded the concept of this synthesis to nonwater-based polar solvents by performing syntheses in DMSO¹⁰¹ and DMF.⁹² Using this approach, the PL spectra were pushed to 4.5 μm. This approach is also relevant for IR detection because NCs are readily available in polar solvents in which the particles are capped by short ligands compatible with transport. This approach has already been utilized for PbS nanocrystals by realizing NC formation without long insulating ligands to obtain an ink that can be spin-coated to form a conductive film of NCs with a low density of the surface traps.¹⁰²

2.1.2. Synthesis in an Organic Medium. A key limitation of aqueous syntheses is the use of H₂Te, which makes the synthetic setup considerably more complex than that for solid/liquid based syntheses. Alternative growth methods, typically conducted in organic media, have been developed. As for water-based syntheses, early evidence for mercury chalcogenides NC organic syntheses can be found. In the pioneering work by Murray, mostly dedicated to CdSe, mercury chalcogenides are also mentioned.⁶ In the earliest stage of the hot injection

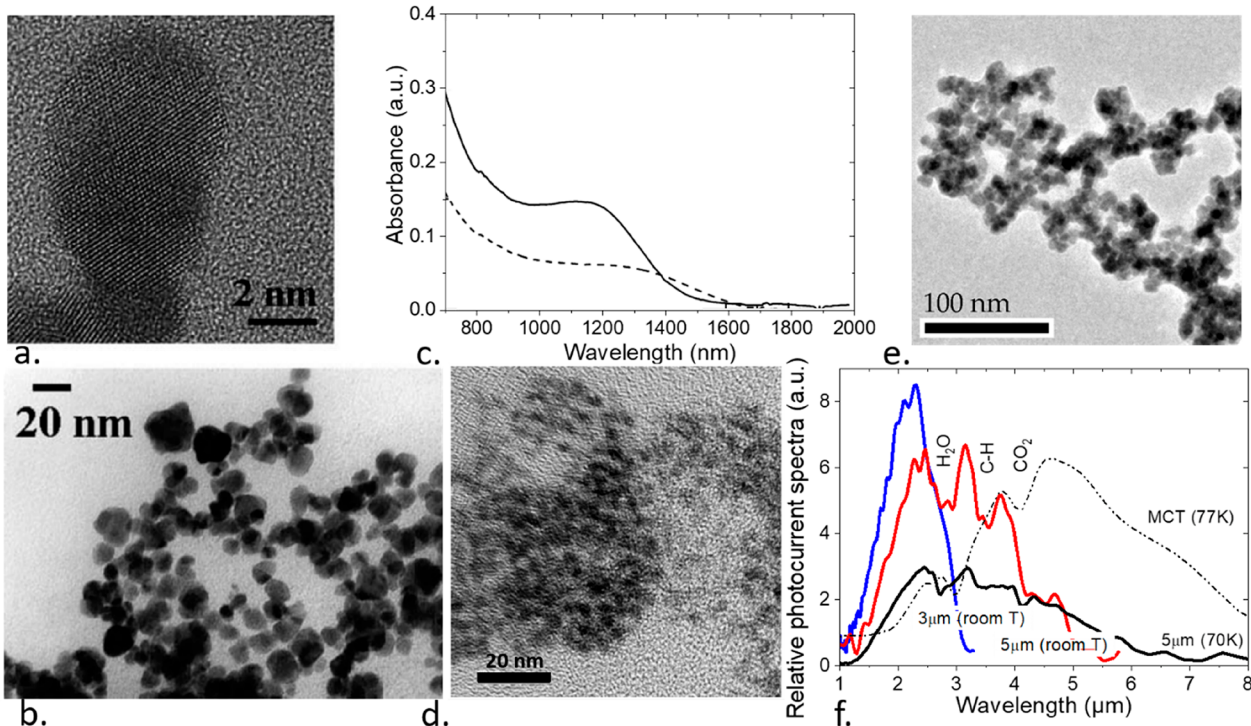


Figure 4. (a,b) TEM images of HgTe capped with octadecylamine and TOPO prepared at 70 °C. (a,b) Reproduced with permission from ref 93. Copyright 2003 Royal Society of Chemistry, permission conveyed through Copyright Clearance Center, Inc. (c) Absorption spectra of HgTe nanoparticles. (d) TEM image of HgTe QDs redissolved in toluene. (c,d) Adapted with permission from ref 94. Copyright 2006 American Chemical Society. (e) TEM image of HgTe CQDs with an absorption cutoff at 5 μm. (f) Fourier transforms of the interferograms measured from films of HgTe NCs with cutoff wavelengths at 3 and 5 μm. (e,f) Adapted from ref 3. Copyright 2011 Nature.

method, highly reactive precursors are used, such as metal alkyl. Diethyl cadmium is highly reactive and easily forms pyrophoric compounds; however, the toxicity of its mercury counterparts is even higher.¹⁰⁴ Consequently, additional approaches for synthesizing HgTe NCs based on alkyl mercury have not been proposed. Nevertheless, the use of phosphine as a capping ligand has been retained. Green *et al.* proposed a mild temperature (70 °C) synthesis in which a mercury halide reacts with the Te complex and trioctylphosphine (TOP:Te) in a mixture of a coordinating solvent (TOPO and long chain amines).^{93,105} This combination of the Hg²⁺ salt and TOP:Te in the presence of an amine has been the backbone of most syntheses approaches developed subsequently. The authors reported fairly large NCs (up to 17 nm), as shown in Figure 4a,b, with a featureless optical spectrum. Although the lack of excitonic features might correspond to polydispersity, such features likely appear at wavelengths larger than the reported data. Piepenbrock *et al.* enhanced this method to realize a less polydisperse synthesis of small NCs (3–4 nm), as shown in Figure 4d. This synthesis was based on the use of mercury acetate as the Hg precursor in the presence of long chain amines and ethanol as the solvent. The TOP:Te was slowly added while cooling the reaction to −78 °C (dry ice). The NCs exhibited an absorption shoulder at 1200–1600 nm, as shown in Figure 4c.

In 2011, Keuleyan *et al.* introduced the first synthesis technique to realize absorption and photoconduction up to 5 μm, as shown in Figure 4f.³ This approach represented a breakthrough from an application point of view because the material was sufficiently red to enable thermal imaging with NCs. However, the associated quality was low, with severe aggregation, as indicated in the TEM image shown in Figure 4e. In addition, this synthesis was relatively different from the

existing approaches. First, this approach was the only one based on elemental Te. While most syntheses rely on hot injection of the Te^{2−} anion, this procedure relies on the introduction of Te powder, pyridine, and butanol to the flask and injection of the cation salt (mercury acetate). Although the inherent aggregation of this synthesis technique was appealing for transport to achieve a high carrier mobility, the wide size dispersion of the NCs led to extremely broad optical features with only the tail of absorption actually reaching the 3–5 μm range. The synthesis led to larger particles (up to 12 nm); however, the monodispersity needed to be enhanced.

Around 2010, a series of conceptually similar procedures were demonstrated for the growth of HgTe NCs.^{22,95,106,107} Most of the concepts were based on the use of the conventional hot injection method (injection of an anion in a cation solution at a “high” temperature). A Hg²⁺ salt from HgCl₂ or mercury acetate was dissolved in the presence of an amine, a thiol, and possibly oleic acid.¹⁰⁷ A non-coordinating solvent could also be added.¹⁰⁶ The Te precursor was obtained from Te complexed with phosphine, as initially proposed by Green *et al.*⁹³ This approach is widely used for photoconduction devices.^{35,48,108} Thus, it is worth describing the role of each component. The reaction is conducted under mild temperatures (60–120 °C): a higher temperature corresponds to larger particles. Similarly, the reaction duration can be considered as a size control parameter. At a temperature of 60 °C and duration of 3 min, particles with a band edge at ≈6000 cm^{−1} (≈720 meV or 1.8 μm) can be obtained, corresponding to a particle size of approximately 6 nm; the conditions of 120 °C and 15 min yield a band edge at 2000 cm^{−1} (≈250 meV or 5 μm), leading to a particle size of approximately 12 nm, as shown in Figure 5a. To obtain even larger particles, a regrowth step in which the Te precursor is

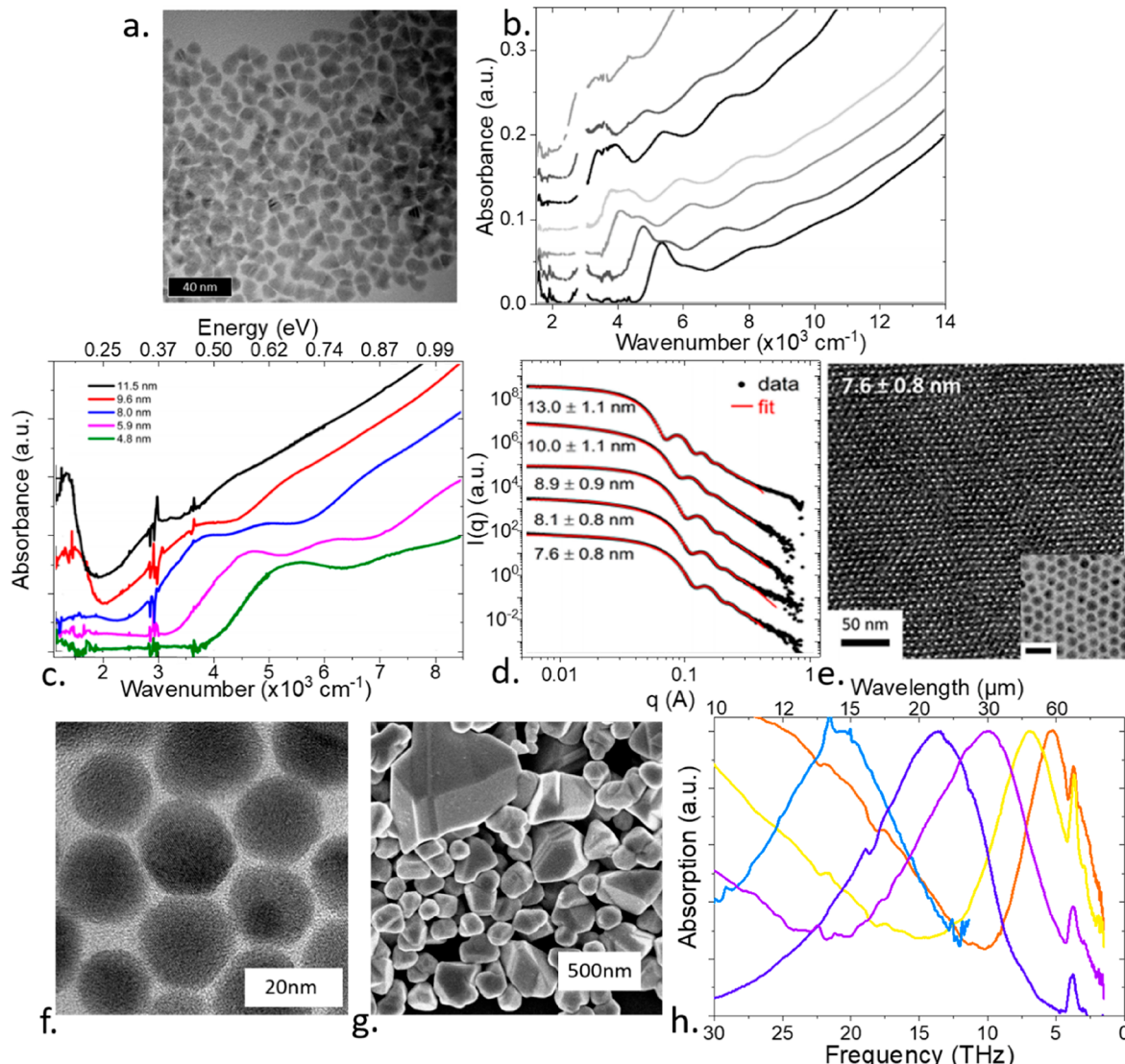


Figure 5. (a) TEM image of 9.1 nm HgTe NCs. (b) Absorption spectra of differently sized HgTe NCs showing multiple features. (a,b) Adapted with permission from ref 28. Copyright 2014 American Chemical Society. (c) Adapted with permission from ref 63. Copyright 2017 American Chemical Society. (d) Small-angle X-ray scattering signals and their corresponding fits for colloidal solutions of HgTe NCs exhibiting a reasonable size control ($\approx 10\%$ distribution) across a range of samples. (e) TEM image of a HgTe NC-based superlattice upon slow evaporation of the solvent. The inset image shows a monolayer of the same particles dropcast from a dilute solution (scale bar, 10 nm). (d,e) Adapted with permission from ref 61. Copyright 2018 American Chemical Society. (f,g) TEM images of large HgTe NCs with absorption above 10 μm . (h) Series of infrared spectra for extremely large HgTe NCs. (f–h) Adapted with permission from ref 30. Copyright 2018 American Chemical Society.

injected dropwise must be performed.²⁸ The Hg to Te ratio is generally set as approximately 1. An amine must be used to realize monodisperse syntheses,^{109,110} and a more dilute synthesis corresponds to a lower polydispersity.²⁸ Under the dilute condition, a NC solution with an absorption spectra involving up to six clearly defined optical transitions can be obtained,^{28,30} as shown in Figure 5b. In this case, the HgTe NCs in the IR range are as desirable as the Cd-based NCs in the visible range. The introduction of carboxylic acid (even using mercury oleate as the precursor¹⁰⁷) and thiol during the synthesis accelerates growth. Because the synthesis is conducted at mild temperatures, lowering the temperature to RT is not sufficient to stop growth. Thus, a long chain thiol (dodecanethiol) is generally introduced to render the NC surface inaccessible to the remaining precursors. Goubet *et al.* used the same approach; however, both the Hg and Te precursors were

injected into the hot solvent,³⁰ as earlier proposed to obtain highly monodisperse PbSe NCs.¹¹¹ In this case, the motivation was to avoid degradation of the Hg precursor. Using this strategy, the authors succeeded in realizing the growth of HgTe NCs from 60–300 °C and obtained an extremely broad size tunability from 5 to 1000 nm, as shown in Figures 5f–g.³⁰ Using such large particles, the authors reported absorption up to 60 μm for the peak and up to 200 μm for the cutoff,³⁰ as shown in Figure 5h. These NCs were the first reported structures larger than the HgTe Bohr radius¹¹² (40 nm) to present absorption in the THz range.

An interesting evolution of this method was the replacement of the phosphine-based Te precursor with bis(trimethylsilyl)-telluride (TMS)₂Te.⁶³ Bis(trimethylsilyl)sulfide has been widely used to realize the growth of CdS shells and synthesize PbS cores.⁸³ Bis(trimethylsilyl)chalcogenides are highly reactive

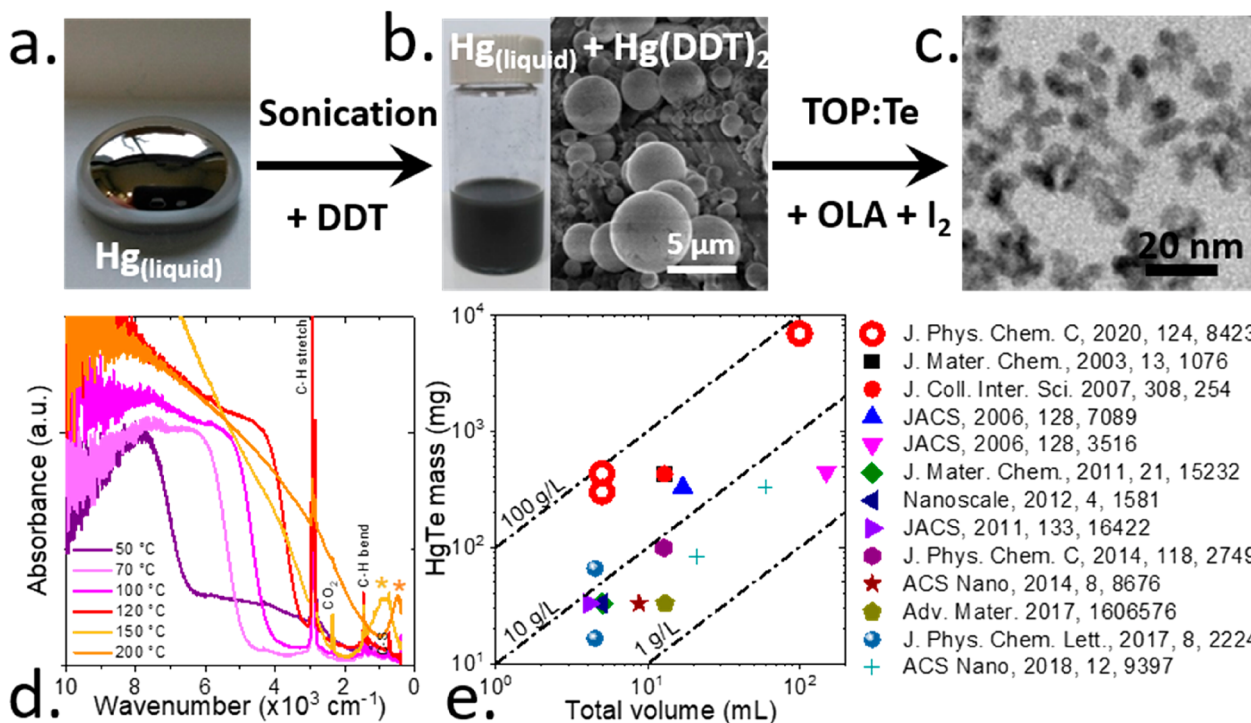


Figure 6. (a) Image of a liquid Hg drop. (b) Optical and SEM images of a solution of mercury dodecylthiolate prepared from liquid Hg. (c) TEM picture of HgTe QDs synthesized using liquid Hg as a precursor. (d) Absorbance spectra of HgTe QDs synthesized from 50 to 200 °C by using liquid Hg as the precursor. (e) Comparison of the mass yield obtained in this work to those reported previously, assuming the total conversion of the limiting reagent into a HgTe material. Adapted with permission from ref 96. Copyright 2020 American Chemical Society.

forms of chalcogenides that rapidly decompose and behave as reduced forms of anions (S^{2-} , Se^{2-} , and Te^{2-}). When $(TMS)_2Te$ is used to grow HgTe, more spherical shapes can be achieved; in contrast, the reaction based on TOP:Te leads to branched or faceted shapes. These spherical particles enable the formation of HgTe NC-based superlattices⁶¹ (see Figure 5d,e) and can be used to enhance the carrier mobility by allowing a higher packing density in the film.³⁶ However, this strategy is difficult to generalize owing to the high cost and small lifetime of $(TMS)_2Te$. Nevertheless, this synthesis is essential because it is the first reported intraband absorption for HgTe NCs, as described in the section dedicated to doping and Figure 5c.

Recently, Goubet *et al.*⁹⁶ addressed the scale-up of the synthesis. Current syntheses are significantly diluted (<10 g·L⁻¹ of NCs per volume of solution, as shown in Figure 6e). This issue is critical to scalability, as high dilution conditions require greater effort to obtain the same amount of NCs. Specifically, the setup and precursors are relatively inexpensive, and human effort is the main cost related to NC growth.¹¹³ In addition, a more concentrated solution generally reduces the amount of solvent (and associated cost) required in the NC cleaning steps. Therefore, it is necessary to develop more concentrated syntheses. As mentioned previously, most strategies to grow HgTe NCs rely on the use of Hg²⁺ salt. A key issue with the existing syntheses is the low solubility of Hg halide in organic solvents, especially at the temperatures used for HgTe NC growth (*i.e.*, below 100 °C). This aspect dramatically reduces the synthesis concentration. The use of an aprotic polar solvent (DMF, DMSO) can help increase the salt solubility and achieve a nanocrystal concentration of up to 60 g·L⁻¹.⁹² Alternatively, Goubet *et al.* proposed the use of Hg thiolate as a strongly soluble form of Hg in oleylamine (OLA) and demonstrated that Hg thiolate can be formed from liquid Hg sonicated with

DDT,⁹⁶ as shown in Figure 6a,b. This mixture reacts with TOP:Te, as shown in Figure 6c. Using this approach, the authors demonstrated the synthesis of HgTe NCs from NIR to longwave IR, as shown in Figure 6d. Up to 7 g of 2 μm cutoff HgTe NCs could be achieved from 150 mL of liquid (including the reaction and cleaning solvents),⁹⁶ as shown in Figure 6e.

Before completing this discussion related to material growth, the sizing curve must be mentioned as a valuable tool to empirically correlate the particle size to the absorption feature. For HgTe, two models have been proposed. The first model was reported by Kovalenko *et al.*,²⁷ which provided the energy of the interband transition (in eV) as a function of the particle (d in nm) size: $E = 0.3 + 0.2 d^{-1} + 5.7 d^{-2}$. The other model was proposed by Lhuillier *et al.*, which was obtained by mixing a two-band k·p model through empirical fitting.¹¹⁴ In this case, the band edge wavelength (in μm) was related to the particle radius (R) according to the following expression: $\lambda = \frac{30.5}{\sqrt{1 + \left(\frac{43}{R}\right)^2} - 1}$.

Nevertheless, these two expressions are old, and the size range that can be obtained has been considerably extended, as shown in Figure 2b. When considering only the interband transition (*i.e.*, excluding the feature resulting from the intraband and plasmonic absorption observed in the largest NCs), an empirical expression to fit the wavelength *vs* size (*i.e.*, effective diameter in nm) is $\lambda = 0.17d^{1.49}$, as shown in Figure 2b.

2.2. HgSe Nanocrystal Synthesis

In addition to telecom motivations,¹¹⁵ similar to those reported for HgTe, the initial motivation for HgSe NCs was to push the optical feature of CdSe toward larger wavelengths to attain an infrared transparency window in which bioimaging could be realized.¹¹⁶ Most of the initial papers on HgS(e) reported on its optical features in the red¹¹⁶ and near-infrared ranges¹¹⁵ as well

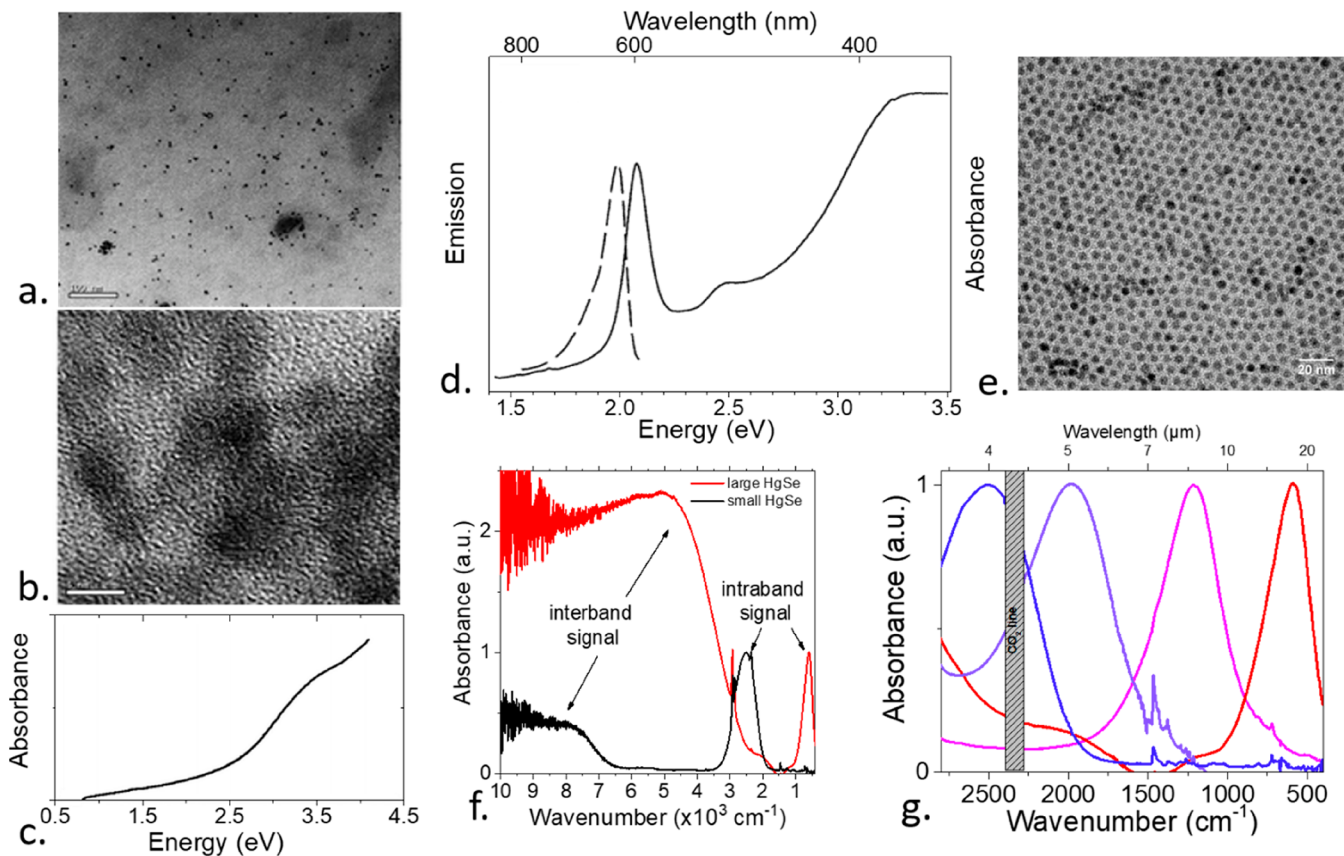


Figure 7. (a,b) TEM images of HgSe nanocrystals made with mercury acetate and TOP:Se in TOPO. (c) Corresponding visible to near IR absorption spectrum. (a–c) Reproduced with permission from ref 117. Copyright 2008 Royal Society of Chemistry, permission was conveyed through the Copyright Clearance Center, Inc. (d) Absorption (full line) and photoluminescence (dashed line) spectra of HgSe clusters prepared in the microemulsion. (d) Adapted with permission from ref 116. Copyright 2003 American Chemical Society. (e) TEM image of typical HgSe nanocrystals synthesized in oleic acid and oleylamine. (f) FTIR spectra of small and large HgSe nanocrystals. (g) Intraband absorption range of HgSe nanocrystal obtained from this synthesis. (e–g) Adapted with permission from ref 29. Copyright 2016 American Chemical Society.

as the UV range.¹¹⁷ The latter observation may appear surprising because the bulk bandgap of HgSe is smaller than that of CdSe and the optical band edge is expected to be found at a larger wavelength for HgSe NCs than for CdSe NCs at the same particle size. In 2014,³⁴ this puzzling observation was clarified by observing the intraband absorption in HgSe NCs. Similar to HgS,³³ HgSe NCs appear to be self-doped⁶⁸ (as described in the section dedicated to doping later in this chapter), indicating that doping is realized without the introduction of any impurities while conducting synthesis under the stoichiometric condition (the same molar amounts of Hg and Se). The main evidence for this doping is the presence of a narrow peak in the mid-infrared range, as shown in Figure 7f,g. This peak is the signature of a Fermi level located within the conduction band above the 1Se state, as shown in Figure 12f. Because of this degenerate doping, the interband transition is partly bleached, which considerably smooths the visible and even near-infrared absorption. This lack of clear features in the visible domain and the shifting of the absorption toward the mid-infrared range is likely why the intraband absorption was missed in early works on HgSe. Since this observation was made, we observe a complete shift of the motivation for HgSe NC development. The interest in bioapplications appears limited likely due to the presence of Hg, limiting the use of HgSe particles to *in vitro* applications. Nevertheless, many researchers have attempted to integrate HgSe NCs into devices by exploiting their infrared intraband absorption.^{29,34,39,47,118–120}

Table 2 summarizes the main synthetic processes used to grow HgSe NCs. The first synthesis of nanosized HgSe involved nearly cluster-sized materials. Similar to the first HgTe synthesis, these syntheses were performed in a polar solvent. Considering the case of HgTe nanocrystals,²⁰ Harrison *et al.*¹¹⁵ developed a synthesis technique for HgSe nanocrystals capped with thioglycerol in water. The chalcogenide precursor was obtained from H₂Se gas, generated *ex situ* in an adjacent inline flask. Eventually, small HgSe NCs with a PL at 1 μm were obtained.

Certain early synthesis approaches for HgSe were partly inspired by bioreactions: HgSe is a common byproduct of the Hg exposure of biological entities through the reaction of Se containing protein with Hg.⁵⁴ Thus, Kuno *et al.*¹¹⁶ realized the synthesis of a molecular cluster of binary and ternary mercury chalcogenides, including pure HgSe. Clusters were generated using a microemulsion with polyoxyethylene 4-lauryl ether (Brij30) and bis(2-ethylhexyl) sulfosuccinate (AOT) as a surfactant. The clusters were stabilized with a “magic” size and demonstrated optical absorption and photoluminescence at 595 and 628 nm, respectively, as shown in Figure 7d.

In 2008, the synthesis approach for HgTe proposed by Green *et al.*⁹³ was adapted for HgSe NCs.¹¹⁷ TOP:Se was used as a selenium precursor that reacted with mercury acetate in a mixture of TOPO and ethanol. Surprisingly, the 4.9 nm nanocrystals exhibited a band edge at 2.5 eV without any clear excitonic peaks, as shown in Figure 7a–c, indicating that the IR absorption of this material was likely missed in the early work.

Table 2. Conditions Related to the Selection of the Relevant Synthesis Techniques for HgSe NCs

Hg precursors	Se precursors	ligands	solvents/temperature	method	size (nm)	optical properties	year/ref
Hg(ClO ₄) ₂	H ₂ Se	thioglycerol	H ₂ O, thioglycerol T_{room}			1 μm 1 μm (PL)	2000 ¹¹⁵
Hg(acetate) ₂	(TMS) ₂ Se	thioglycerol	H ₂ O, thioglycerol, hexane, TOP, C ₁₂ E ₄ , AOT/ T_{room}	microemulsion	2–3	595 nm 620 nm (PL)	2003 ¹¹⁶
Hg(acetate) ₂	TOP:Se		TOPO, ethanol/ T_{room}		3.9–6	350 nm	2008 ¹¹⁷
HgCl ₂	Se=C(NH ₂) ₂	DDT	OLA/110 °C	hot injection	5.5–6.2	4 μm	2014 ³⁴
Hg(acetate) ₂	Se-ODE		OA, ODE/100 °C, T_{room} and −78 °C		15–100	800–1000 nm 1–1.3 μm (PL)	2014 ¹²³
Hg(acetate) ₂	cation exchange on CdSe	OA, OLA, DDT	hexane, oleylamine, trietylamine, OLA/ T_{room}	cation exchange	2–3 atomic ML 100 × 200	700–800 nm	2016 ³¹
Hg(acetate) ₂	TOP:Se SeS ₂	DDT	OA, OLA/60–120 °C	hot injection	6–23	4–20 μm	2016 ²⁹
HgCl ₂	cation exchange on CdSe	TOPO	ODE, OLA/150 °C	cation exchange	11.3	5 μm	2017 ¹²⁴
HgCl ₂ , HgI ₂	Se	DDT	OLA/250 °C	injection of both cation and anion in hot solvent	20–100	9–18 μm	2018 ³⁰

The use of a long chain and coordinating amine, such as oleylamine, is critical to synthesize well-defined HgSe nanocrystals. Deng *et al.*³⁴ described the synthesis of HgSe nanocrystals with a low size dispersion through the reaction of mercury chloride with seleneourea in oleylamine. The typical reaction temperature was approximately 100 °C (between 60 and 150 °C). Temperature was the main parameter driving the final particle size. At this stage, the surface chemistry of the particle consisted of oleylammonium chloride,¹⁰⁹ which could be replaced by thiol either through quenching of the reaction or by a postsynthesis treatment. Moreover, the reaction was generally conducted under stoichiometric conditions or with a slight Hg excess to favor the doping level and carrier mobility.¹²¹ The size of the nanocrystals (5.5–6.2 nm) was similar to that of the nanocrystals obtained through TOPO with TOP:Se.¹¹⁷ However, Deng *et al.* demonstrated that an intraband transition occurred in the mid-IR range (4 μm). It is likely that the particles obtained using Howes' method (mercury acetate/TOPO/ethanol reacting with TOP:Se) presented intraband absorption and that this aspect was missed by the authors. The process used by Deng *et al.*³⁴ was clearly derived from Keuleyan's procedure for HgTe.⁹⁵ In the latter procedure, the HgCl₂ in oleylamine was used as the reacting medium. Because TOP:Se is less reactive than TOP:Te, the authors replaced the Se precursor with a more reactive species, such as seleneourea. Lhuillier *et al.*²⁹ used the opposite approach; instead of increasing the reactivity of the Se precursor, TOP:Se was used as the precursor, and a more reactive form of mercury, such as mercury oleate, was employed. The latter can be easily prepared by mixing mercury acetate in the presence of oleate. This compound however degrades at high temperatures (more than or equal to 150 °C). Under such conditions, in the presence of an amine, Hg^{II} is easily reduced to metal Hg. The temperature and duration of the reaction were the two main parameters to tune the particle size. However, degradation of the Hg precursor likely hindered the generation

of large NCs. To overcome this difficulty, the authors recommended the use of a phosphine free synthesis, in which the Se precursor was derived from SeS₂ diluted in oleylamine. Owing to the lack of phosphine, a larger amount of the nanoparticle surface was available, which facilitated growth, leading to particle sizes of up to 40 nm. This large size tunability allowed the tuning of the energy of the intraband feature from 3 to 25 μm , as shown in Figure 7g.^{29,122}

The shape of the HgSe NCs is spherical (Figure 7e), in contrast to that of HgTe, for which most of the synthesis approaches lead to faceted objects. The resulting NCs appear to be relatively monodispersed and easily self-organize on TEM grids to form a close packed lattice. Nevertheless, recent transport measurements have indicated that the remaining size distribution limits mobility in HgSe NC arrays,¹²¹ requiring the development of more effective synthesis techniques.

2.3. HgS Nanocrystal Synthesis

HgS exhibits two key differences against HgTe and HgSe: (i) the metacinnabar (β -HgS) zinc blende structure of mercury sulfide is a narrow bandgap⁶⁸ (0.65 eV) semiconductor, while HgTe and HgSe are semimetals. Despite this bandgap, the optical properties of this phase are highly similar to those of HgSe, exhibiting intraband absorption (Figure 8c) due to self-doping. In addition, (ii) in the bulk and nanoscale, mercury sulfide is stable in two crystal structures. In the second structure, the cinnabar phase (α -HgS) exhibits a bulk bandgap of 2.1 eV and is known as vermillion due to its red color.

2.3.1. β -HgS Nanocrystal Synthesis. Table 3 summarizes the main synthetic processes used to grow β -HgS NCs. As in the case of HgSe, HgS nanocrystals were first synthesized for the visible–near-infrared ranges. The first synthesis^{125–127} of HgS nanosized materials was in the form of CdS/HgS/CdS heterostructured quantum dots. The core of CdS was coated with a few layers of HgS and capped with CdS, thereby achieving a kind of spherical quantum well heterostructure. The CdS

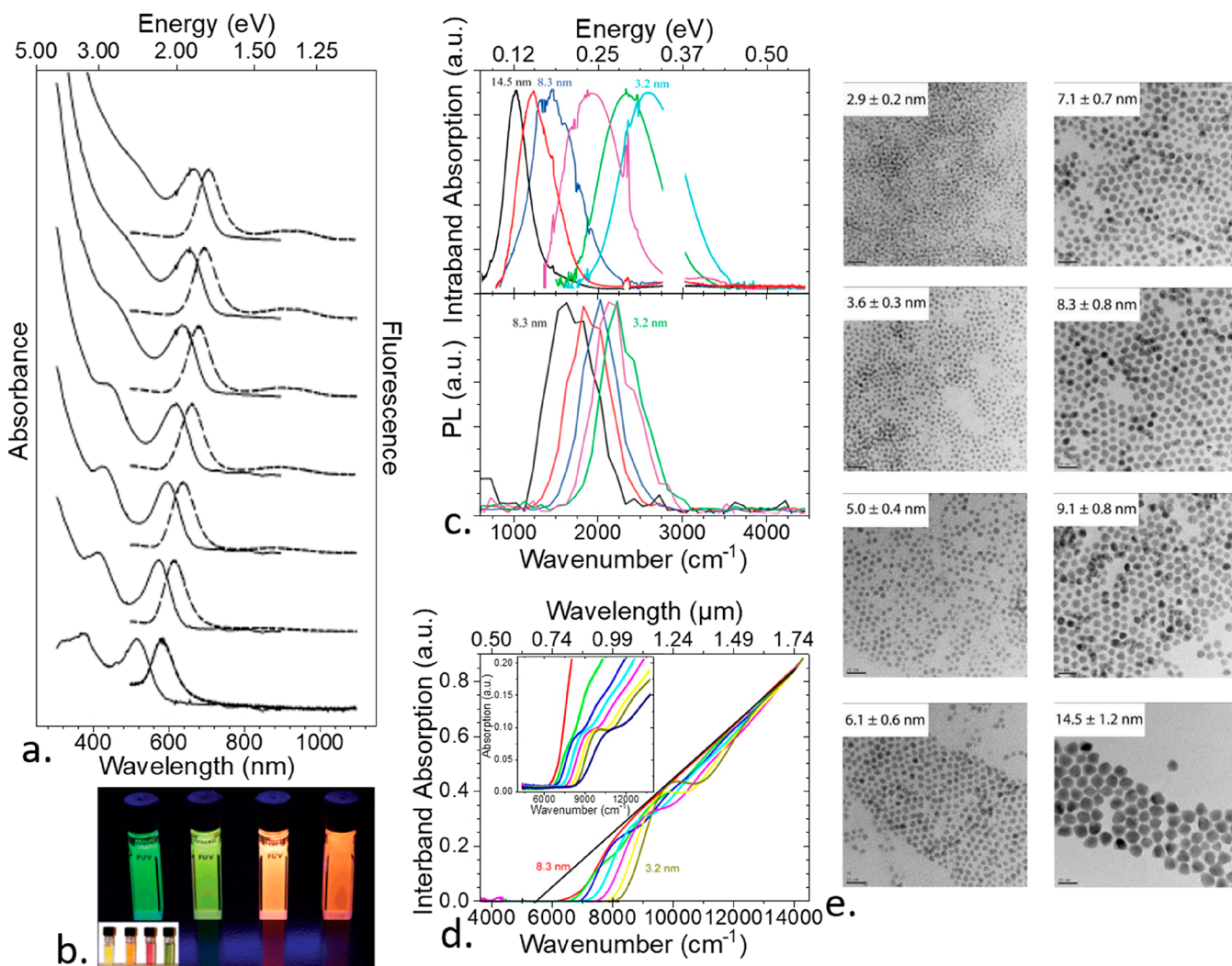


Figure 8. (a) Photoluminescence and absorption spectra of β -HgS nanocrystals during their formation. (b) β -HgS nanocrystals under UV illumination and normal illumination (inset). (a,b) Adapted with permission from ref 129. Copyright 2002 American Chemical Society. (c) Intraband absorption and photoluminescence of β -HgS nanocrystals with different sizes. (d) Interband absorption of β -HgS nanocrystals with various sizes. (e) TEM images of β -HgS nanocrystals with various sizes. (c–e) Adapted with permission from ref 136. Copyright 2016 American Chemical Society.

quantum dots were formed in the presence of sodium polyphosphate by reacting H_2S gas with $\text{Cd}(\text{ClO}_4)_2$ in water. The HgS quantum well was grown with $\text{Hg}(\text{ClO}_4)_2$ with the same sulfur precursor. Next, the reaction with $\text{Cd}(\text{ClO}_4)_2$ was repeated for the last layer. The number of HgS layers (from 1 to 4) sandwiched between CdS could be increased by increasing the deposition cycles. The particle band edge was red-shifted from 470 to 800 nm for larger heterogeneous materials. Later, Dorfs *et al.*¹²⁸ enhanced the complexity of this heterogeneous structure by proposing the synthesis of multilayered spherical CdS/HgS nanocrystals. The key concept was to increase the thickness of CdS materials before growing the HgS shell to reach partial substitution of the CdS layer by HgS. In 2002, Higginson *et al.*¹²⁹ developed a synthesis technique for small HgS nanocrystals. The use of a strongly binding ligand, such as thioglycerol for Hg^{2+} , and compartmentalization of the Hg and S precursors were crucial to realize controlled growth of HgS nanocrystals. HgS growth was stopped by adding a cadmium or zinc precursor, which increased the photoluminescence of the materials (QY of 5–6%) compared to that of the materials passivated using aqueous precursors (QY < 1%). The size range of the resulting materials was from 1 to 5 nm. As shown in Figure

8a, the corresponding absorption spectra exhibited resolved excitonic features, and band edge absorption appeared between 500 and 800 nm. Owing to the Cd and Zn passivation, the HgS nanocrystals exhibited clear photoluminescence in the visible range, as shown in Figure 8b. An alternative approach to synthesize aggregated HgS nanocrystals was proposed by Yang *et al.*,¹³⁰ who exploited the fact that ordered two-dimensional arrays of aggregates could be obtained at the interface between water and chloroform. HgS nanocrystals were formed through the interfacial reaction between $\text{Hg}(\text{acetate})_2$ and H_2S gas from the upper aqueous phase. The Langmuir monolayer of arachidic acid was used to create aggregates between this surfactant and the HgS nanocrystals. The aggregate size was approximately 260 nm, with the HgS nanocrystals having a size of 8 nm.

The first synthesis using TOP:S as a chalcogenide precursor was proposed by Wichiansee *et al.*¹³¹ in 2011. To overcome the low reactivity of TOP:S compared to TOP:Te and TOP:Se for TOPO-capped HgTe and HgSe syntheses, HgS nanocrystals were synthesized at 120 °C instead of near room temperature. The resulting nanocrystal solution was polydisperse, with a mean value of 3.9 nm. This polydispersity led to a broad band edge at 910 nm with a weak and broad photoluminescence from

Table 3. Conditions Related to the Selection of the Relevant Synthesis Technique for β -HgS NCs

Hg precursor	S precursors	ligands	solvents/temperature	method	size (nm)	optical properties	year/ref
Hg(ClO ₄) ₂	H ₂ S	CdS	water/ <i>T</i> _{room}	cation exchange	1–3 monolayers	500–800 nm 713–961 nm (PL)	1994 ¹³⁷
Hg(acetate) ₂	(TMS) ₂ S	thioglycerol, thiophenol, dodecylamine, hexadecanethiol	cyclohexane, hexane, H ₂ O, thioglycerol/ <i>T</i> _{room}		1–5	500–800 nm 600–700 nm (PL)	2002 ¹²⁹
Hg(acetate) ₂	TOP:S	TOP/TOPO	TOPO/150 °C	hot injection	4	broad 900 nm broad 1127 nm (PL)	2011 ¹³¹
Hg(ClO ₄) ₂	Na ₂ S	glutathione	H ₂ O, glutathione/ <i>T</i> _{room}		2.87	700–800 nm 775–1041 nm (PL)	2012 ¹³²
HgCl ₂	thioacetamide	lysozyme	H ₂ O, lysozyme/37 °C	lysozyme template	14–27	400 nm 470 nm (PL)	2012 ¹³³
Hg(NO ₃) ₂	Na ₂ S	BSA	H ₂ O, NaOH, BSA/ <i>T</i> _{room}	biomineralization	3.8	480–600 nm 680–800 nm (PL)	2012 ¹³⁴
HgCl ₂	thioacetamide, (TMS) ₂ S, (NH ₄) ₂ S	DDT	OLA/30 °C	injection	3.5–14	5 μm 5 μm	2014 ³³
Hg(NO ₃) ₂	S	DDA, OLA, or OA	toluene, OLA, OA, DDA/ <i>T</i> _{room} or 80 °C	phase transfer	12–100		2014 ¹³⁵
HgCl ₂	(NH ₄) ₂ S	OLA, TOP, or DDT	OLA, TCE, TOP/ <i>T</i> _{room}		3–15	4–10 μm 3.6–6.7 μm (PL)	2016 ¹³⁶
HgCl ₂ , HgI ₂	S	DDT	OLA/250 °C	injection of both cation and anion in hot solvent	50–150	9 and 18 μm	2018 ³⁰

900 to 1600 nm. Other near-infrared photoluminescence materials were obtained with glutathione-coated HgS nanocrystals.¹³² In this synthesis, the mercury precursor (Hg(ClO₄)₂) was dissolved with glutathione in water. HgS was obtained via the slow injection of Na₂S, and the obtained material exhibited an excitonic shoulder at 781 nm with photoluminescence at 915 nm. The photoluminescence was maximum at pH 9.5, which corresponds to the deprotonated form of glutathione; in this state, most of the thiol groups efficiently bind the Hg on the nanocrystal surface, reducing the impact of the surface traps.

Two synthesis approaches using proteins were proposed for HgS. First, Zhang *et al.*¹³³ used lysozyme as ligands for HgS nanocrystals: HgCl₂ dissolved with lysozyme in water reacted with the S²⁻ provided by thioacetamide at 37 °C. The second approach was proposed by Goswami *et al.*¹³⁴ In this synthesis, HgS nanocrystals were obtained using the biomineralization process. In the first step, HgO nanocrystals were obtained using mercury nitrate and bovine serum albumin (BSA) in water. Next, Na₂S was used for the sulfidation of HgO to HgS nanocrystals. The resulting HgS nanocrystals exhibited a clear excitation peak from 480 to 600 nm, with photoluminescence from 680 to 780 nm. The nanocrystal size was controlled through the Hg/S ratio. Later, Han *et al.*¹³⁵ adapted the phase

transfer synthesis from gold to HgS nanocrystals. Mercury was transferred from the aqueous phase to the organic phase by dissolving mercury nitrate and dodecylamine in water/ethanol. The ions were transferred by mixing the aqueous phase with toluene, oleylamine or oleic acid. Sulfidation was achieved by adding elemental sulfur to the organic phase for 3–4 h. The size of the HgS nanocrystals ranged from 12.5 to 56 nm depending of the sulfidation temperature and organic phase.

The first study that reported the synthesis of HgS nanocrystals with intraband transition was that of Jeong *et al.*³³ This synthesis was adapted from that of HgTe in oleylamine. However, because TOP:S exhibited an extremely low reactivity with HgCl₂, more reactive thioacetamide ((TMS)₂S, or (NH₄)₂S) chalcogenide precursors were used. The reaction was quenched using dodecanethiol (DDT). The obtained HgS nanocrystals had a size of 5 nm, and a clear intraband absorption peak was observed in the mid-infrared range (5 μm). Later, the size ranges of HgS nanocrystals with intraband transition were expanded in the study of Shen *et al.*¹³⁶ The HgCl₂ precursor was reacted with (NH₄)₂S, and the Hg/S ratio allowed size control from 2.9 to 14.5 nm, as shown in Figure 8e. As shown in Figure 8d, the interband transition was slightly modulated by the nanocrystal size compared to the intraband absorption that covered the mid-infrared range from 4 to 10 μm with corresponding photo-

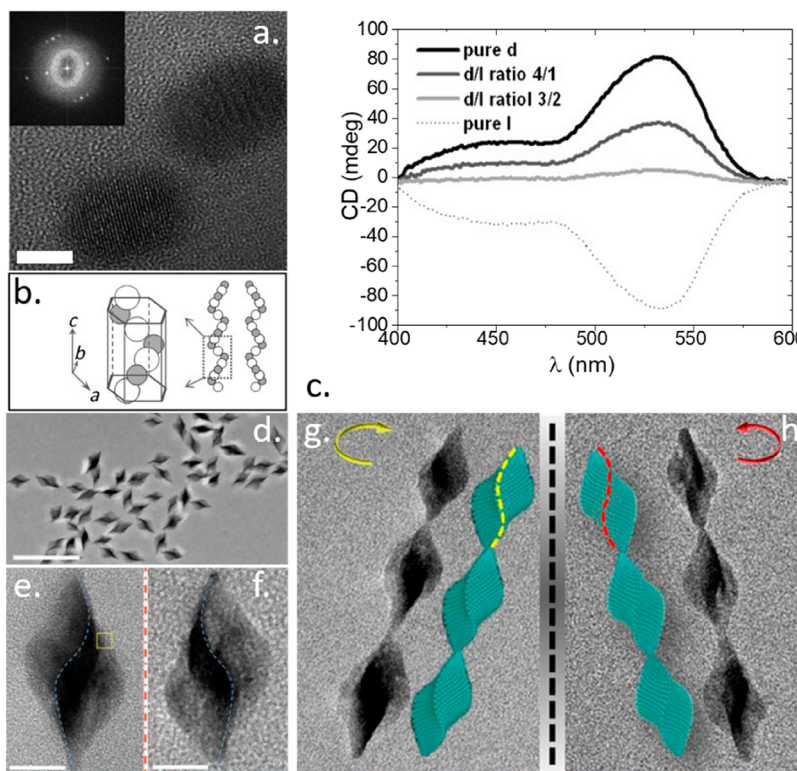


Figure 9. (a) High resolution TEM image of α -HgS nanocrystals. (b) Atomic model of the cinnabar HgS lattice. (c) Circular dichroism (CD) spectra of α -HgS nanocrystals with different D/L ratios of penicillamine. (a–c) Adapted with permission from ref 138. Copyright 2013 Wiley VCH. (d) TEM image of α -HgS nanocrystals synthesized with successful growth. (e,f) TEM image of two helicoidal enantiomorphs of α -HgS. (d–f) Adapted with permission from ref 139. Copyright 2017 Nature. (g,h) TEM images of left- and right-handed oriented attachment of α -HgS nanocrystals. (g,h) Adapted with permission from ref 142. Copyright 2017 American Chemical Society.

luminescence from 3.6 to 6.7 μm . Later, Goubet *et al.*³⁰ expanded the size control of several tens of nanometer nanocrystals from 50 to 150 nm by injecting a mixture of mercury halide and elementary sulfur in hot oleylamine. This last synthesis expanded the intraband absorption up to 18 μm .

2.3.2. α -HgS Nanocrystal Synthesis. Cinnabar crystallizes in the trigonal system with a noncentrosymmetric structure. Specifically, there are two enantiomorphic space groups ($P3_121$ and $P3_21$) in this material (see Figure 9b). Ben-Moshe *et al.*¹³⁸

were pioneers of the use of this structure and proposed the synthesis of intrinsically chiral HgS nanocrystals. In the first step, β -HgS nanocrystals were obtained by reacting mercury nitrate with thioacetamide in the presence of penicillamine in water. By using enantiomeric penicillamine, HgS growth could be directed to one space group of α -HgS after a few hours of reaction. The structural transition from β to α could be observed as the color changed from black to red, the color of the vermillion. As shown in Figure 9a, the long and short axes of the elongated nanocrystals were sized 15 and 10 nm, respectively. The optical activity of α -HgS was characterized by circular dichroism (see Figure 9c). At 550 nm, the band edge exhibited clear optical activity depending on the enantiomeric mixture of the capping penicillamine. The resulting dissymmetry factor (10^{-2}) was 1 order of magnitude larger than that of the achiral nanocrystals coated with chiral ligands. Later, Wang *et al.*¹³⁹ proposed a synthesis of pure HgS enantiomorphs. A pronounced helicoidal chiral shape was obtained after successive growth of α -HgS (see Figure 9d–f). Seeds are grown by reacting mercury nitrate and thioacetamide in the presence of penicillamine (either D or L

enantiomers). A second epitaxial growth was realized with the injection of the same precursors through a syringe pump.

Parts g and h of Figure 9 show the oriented attachment of the left- and right-handed helicoidal α -HgS nanocrystals, respectively. Wang *et al.*¹³⁹ demonstrated that the alignment of the chiral nanocrystals enhanced the optical activity of the material compared to that of the individual nanocrystals. Several studies^{140,141} demonstrated that a slight modification of the chemical structure but not the stereochemistry induced reversed optical activity of the α -HgS nanocrystals.

2.4. Shape Control and 2D Materials

For cadmium^{7,143,144} and lead chalcogenides,¹⁴⁵ a broad variety of NC shapes can be synthesized; for mercury chalcogenides, more effort has been focused on size control to extend the absorption toward larger wavelengths^{3,27–30} than on realizing shape control. The shape of the particles is thus dictated by the targeted bandgap. For example, by using the process recommended by Keuleyan *et al.*,⁹⁵ the shape can be changed from small aspect ratio rods to branched particles to tetrahedrons, with bandgaps of 6000, 4000, and 2000 cm^{-1} , respectively. Among various efforts, Shen *et al.* attempted to control the particle shape of the HgX core nanocrystals.⁶³ The authors used $(\text{TMS})_2\text{Te}$ as a telluride precursor and obtained more spherical particles. Although the initial objective was to reduce material aggregation,⁶³ the shape of the NCs likely considerably enhanced the effective carrier mobility, especially for films of spherical NCs compared to branched nanoparticles with similar surface chemistries.^{36,146}

The most notable efforts to control the shape of HgX NCs were driven by Ithurria and co-workers. This team pioneered the

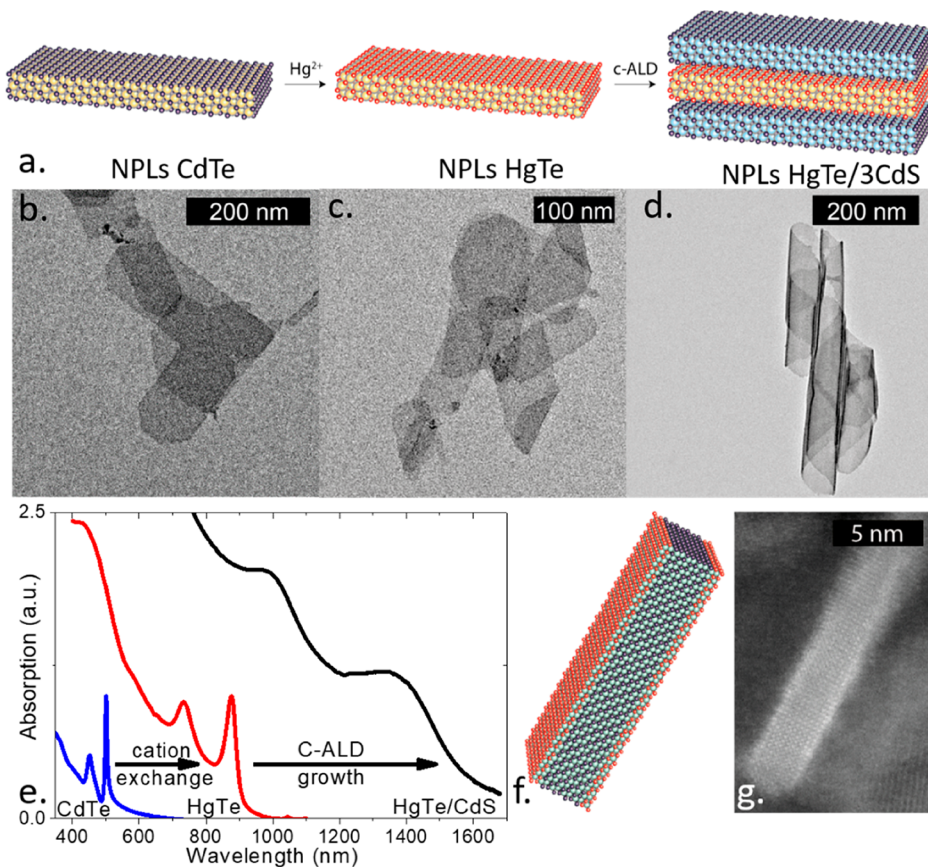


Figure 10. (a) Scheme of the growth mechanism used to achieve HgTe/CdS core–shell NPLs. (b) TEM image of CdTe NPLs. (c) TEM image of HgTe NPLs. (d) TEM image of HgTe/3CdS NPLs. (e) Absorption spectra of CdTe, HgTe, and HgTe/CdS (1 layer of CdS) NPLs. (a–e) Reproduced with permission from ref 165. Copyright 2019 Royal Society of Chemistry. (f) Scheme of a 10 ML HgSe/CdSe/HgSe heterostructure. (g) High resolution TEM in the dark field mode of a 10 ML HgSe/CdSe/HgSe heterostructure. (f,g) Adapted with permission from ref 166. Copyright 2018 American Chemical Society.

field of 2D cadmium chalcogenide nanoplatelets (NPLs).^{8,144} Compared to spherical nanocrystals, the specific growth mechanism of NPLs^{147,148} prevented roughness from occurring along the only confined direction. Consequently, there is no inhomogeneous broadening, and the single particle spectrum overlapped with the ensemble spectrum.¹⁴⁹ This phenomenon led to a narrower PL signal than that for spherical quantum dots, for which the line width was limited by the particle size distribution. The 2D shape also led to a higher exciton binding energy, which increased the radiative rate. The NPLs were thus successfully integrated as active materials to realize stimulated emission,^{150–152} LEDs,^{153–156} and other optoelectronic devices¹⁵⁷ such as field effect transistors and phototransistors.^{41,158,159} The 2D-shaped particles based on lead chalcogenides were reported with optical absorption in the NIR and SWIR.¹⁶⁰ However, the corresponding growth mechanism was different (by clustering preformed nanocrystals) from that of the CdX NPLs, and narrow optical features were not observed in these PbX NPLs. Consequently, researchers attempted to develop HgX materials with a clearly defined structure (clearly defined facet) and optimized optical properties (narrower optical feature, faster PL).

The growth method of CdX NPLs relies on the formation of NC seeds from cadmium oleate and selenium in a non-coordinating solvent (octadecene). When the seeds reach a critical size, a short ligand is introduced to break the symmetry of the seeds, inducing 2D growth.¹⁴⁷ Applying this method while

replacing Cd with Hg has been unsuccessful to date, and no direct synthesis of HgX NPLs has been reported. Izquierdo *et al.*¹⁶¹ proposed an indirect method to obtain HgX NPLs through cation exchange, as shown in Figure 10a. The authors first synthesized CdSe and CdTe NPLs¹⁶¹ and later exposed these NPLs to a Hg precursor. The critical condition to preserve the 2D shape of the NPLs without introducing surface roughness is to ensure slow exchange to avoid destabilizing the chalcogenide lattice.^{161,162} If the cation exchange is incomplete, then puddles of Hg form and dual color emission may occur: the first PL signal comes from CdSe, while the other signal involves Hg trap states.¹⁶³ In contrast, when sufficient Hg is introduced in the Hg oleate form, complete (confirmed through EDX analysis) cation exchange can be realized while preserving the 2D shape of the particles (see Figures 10b,c). During the process, absorption occurs owing to the CdX NPLs first disappearing and later reappearing in a strongly red-shifted manner (see Figure 10e). Because bulk HgTe is a gapless semiconductor, all the band edge energy of the NPL (≈ 1.5 eV) can be attributed to the quantum confinement. Consequently, HgTe NPLs are one of the most confined semiconductor NCs. This aspect is of particular interest to explore the dispersion relation far from the Brillouin zone center. The specific signature of this strong confinement in the temperature dependence of the NPL bandgap is discussed later in the section dedicated to the electronic structure. Because of this strong confinement, the exact band edge energy is strongly dependent on the surface chemistry, which reflects the

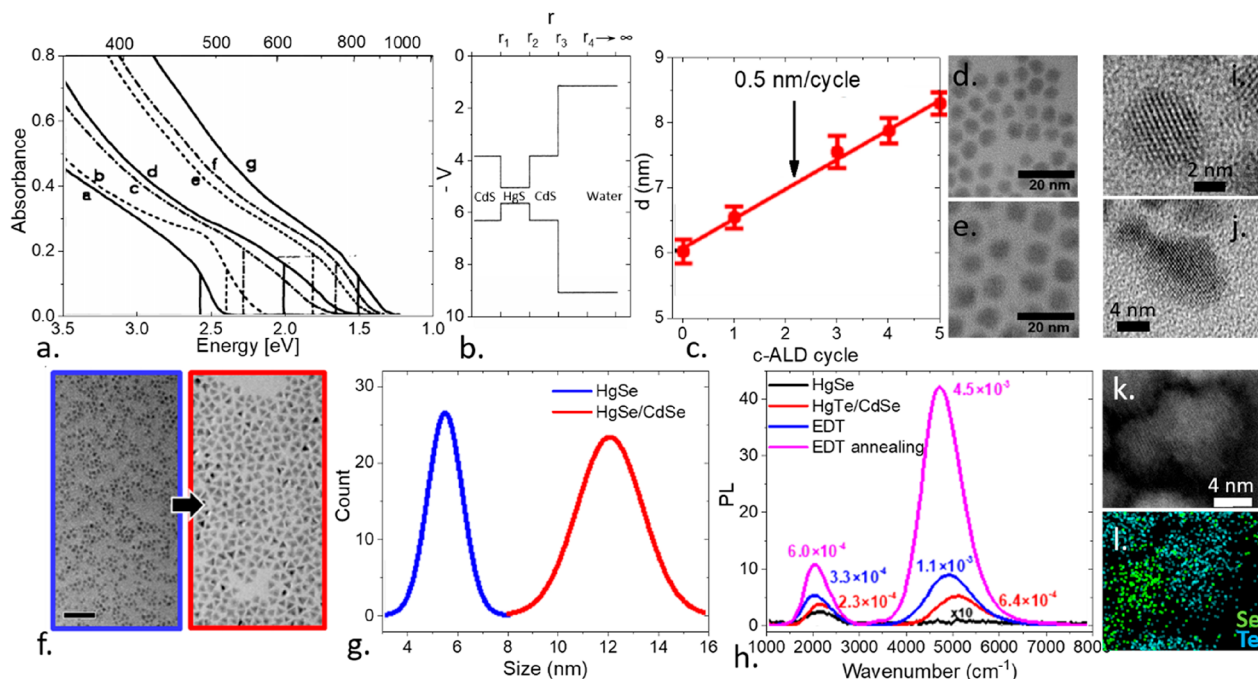


Figure 11. (a) Comparison of the absorption spectra of CdS/HgS/CdS heterostructures with various compositions with the theoretical results. (b) Potential diagram of the CdS/HgS/CdS heterostructure in water. (a,b) Reproduced with permission from ref 125. Copyright 1994 by the American Physical Society. (c) Average nanocrystal diameter as a function of the colloidal ALD cycle. The vertical error bars represent the standard deviation of the QD distribution. Bright field TEM image of 6.0 nm HgSe QDs (d) and HgSe/CdSe QDs (e) obtained after three c-ALD cycles. (c–e) Adapted with permission from ref 185. Copyright 2017 American Chemical Society. (f) TEM of the (blue) HgSe core and (red) thick HgSe/CdSe core/shell (scale bar: 50 nm). (g) Normal size distributions of the (blue) HgSe core and (red) thick HgSe/CdSe core/shell. (h) PL spectra of HgSe/CdSe films with different treatments measured in air. (f–h) Adapted with permission from ref 187. Copyright 2019 American Chemical Society. High resolution TEM images of (i) the HgSe core and (j) HgSe/HgTe heterostructure nanocrystals. (k) HAADF STEM image of a HgSe/HgTe heterostructure and (l) localization of the Se and Te, as obtained through EDX mapping. (i–l) Adapted with permission from ref 120. Copyright 2018 American Chemical Society.

leakage of the wave function outside of the HgX core. A clear signature of the well-preserved roughness-free NPL shape is observed from the PL linewidth. The PL line width (full width at half-maximum) is below 60 meV for an emission at 880 nm, narrower than that of a single PbS NC (typically 100 meV).¹⁶⁴ In addition, the emission is considerably faster than that reported for other NCs emitting in the near IR, with PL decay times of approximately 50 ns, in contrast to the corresponding value of several microseconds for PbS NCs.

While complete cation exchange occurs for thin NPLs (three monolayers), the exchange is not complete for thicker NPLs as only the two surface planes of Cd are exchanged (see Figure 10fg). This observation fully agrees with previous attempts to conduct cation exchange with mercury on nanocrystals, which exhibited a clear self-limiting process.^{15,124,167} This aspect prevents NPL absorption in the mid-infrared region. Nevertheless, an alternative strategy to push the absorption of the NPLs above the telecom range (>1.55 μm) is to realize additional postsynthesis growth of CdS with c-ALD, as reported by Gréboval *et al.*¹⁶⁵ This procedure strongly red-shifts the absorption feature of the HgTe NPLs, as shown in Figures 10a,d,e. The method is described in more detail in the next section dedicated to heterostructures. Note that because of the lattice mismatch between the two materials (>10%), the NPLs twist (see Figure 10d). Regrettably, the introduction of a shell with a type I band alignment does not improve the PL, but the presence of the shell can extend the wave function delocalization and lead to photoconductivity in the SWIR region.

2.5. Heterostructures

Even though the efforts to grow mercury chalcogenide NCs are not as notable as those dedicated to Cd-based materials, significant progress has been made. For CdSe, access to the display mass market has been enabled through the growth of heterostructures with enhanced photoluminescence. Similarly, HgX-based heterostructures appear to be promising candidates to realize high IR luminescence and more stable IR absorbers.

The investigation of colloidal mercury chalcogenides started from heterostructures when Eychmüller *et al.*¹⁵ proposed growing a HgS quantum well on top of CdS NCs, as discussed later in this section. There are multiple motivations for HgX-based heterostructures. First, owing to the low interband and intraband^{33,169} PL efficiency of IR NCs,¹⁷⁰ it is desirable to grow type I heterostructures to boost the luminescence yield. The narrow bandgap of HgX leads to a type I band alignment with most semiconductors. Moreover, the number of non-radiative decay paths, which impede detection owing to the introduction of additional noise,¹²⁵ can be reduced by increasing the quantum yield. In addition, the design of type II band alignment heterostructures is desirable to enhance the exciton dissociation at the nanoparticle level. Another key motivation for HgX-based heterostructures arises from their tendency to sinter even under mild annealing conditions. HgX NCs are mostly grown at low temperatures (60–120 °C); consequently, when such structures are exposed to similar temperatures, HgX nanocrystals aggregate. HgTe appear as strongly confined (*i.e.*, exhibit a large Bohr radius of 40 nm compared to that of 7 nm for CdSe) gapless material. In the case of a stronger

confinement, the electronic structure is more sensitive to size changes. Moreover, in contrast to other semiconductors, the gapless nature of this compound enables the reduction of the NC bandgap to zero. In other words, annealing changes the optical bandgap, which red-shifts the cutoff wavelength; moreover, annealing increases the number of thermally activated carriers and thereby the dark current of the device. Consequently, 60 °C corresponds to the upper temperature limit for the operation of HgX-based optoelectronic devices. Moreover, owing to the low temperature sintering, the diverse fabrication steps typically employed for the encapsulation and protection of nanomaterials or the processing of the resist (*i.e.*, baking step) during the microfabrication procedure cannot be applied in this case. For example, atomic layer deposition, a commonly used protection technique, is generally conducted at 150 °C for several hours. Overall, the growth of HgX heterostructures while ensuring both tunable IR optical features and temperature stability appears to be a bottleneck to integrate these IR materials for mass market devices,¹⁶⁹ beyond the presence of heavy metals.

From a practical perspective, the growth of heterostructures should have benefited from methods developed for other II–VI compounds. However, the growth of HgX-based heterostructures appears to be highly challenging. This difficulty is related to the low temperature growth of HgX compounds, owing to which the core of the HgX NCs is incompatible with the high temperature growth of shells, which occurs at approximately 250 °C and above 300 °C for CdX materials and zinc-based semiconductors, respectively. Several precursors used for HgX growth, such as Hg oleate, also decompose at temperatures above 150 °C. Nevertheless, only a few studies have examined the growth of colloidal HgX heterostructures.

As mentioned previously, the development of HgX nanocrystals was initiated by Weller and co-workers.^{15,125,171–173} The authors performed cation exchange on CdS nanocrystals with Hg²⁺ ions,^{17–19} thereby effectively replacing some of the surface Cd²⁺ with Hg cations. In this manner, a HgS shell was generated, which behaved as a HgS spherical quantum well due to the type I band alignment with CdS,¹²⁷ as shown in Figure 11b. The presence of this narrow bandgap material led to a red-shift of the optical features, and the absorption reached the red and near IR,^{127,174} considerably below the bulk bandgap of CdS (2.4 eV); see Figure 11a. Later, this process was extended to other chalcogenides to obtain CdTe/HgTe^{32,167,175,176} and CdSe/HgSe¹⁶⁶ heterostructures. Despite this early success of cation exchange to grow HgX heterostructures, the reverse strategy in which the HgX core is exposed to Cd atoms has not been extensively investigated. In contrast, in the case of lead chalcogenides, the heating of PbS(e) in the presence of cadmium oleate is the most common strategy to obtain core–shell structures^{177–180} likely because of the relatively large size of Hg atoms, owing to which the introduction or removal of Hg atoms is more difficult than the removal of smaller atoms.

At present, colloidal atomic layer deposition (c-ALD),^{181,182} as a shell growth method, is the most effective approach to realize CdX shell growth on HgX cores. The method was initially developed by Talapin and co-workers to grow shells on 2D nanoplatelets^{67,157,183} (NPLs) of cadmium chalcogenides. At that time, attempts to grow shells using the high temperature method developed for spherical quantum dots on CdSe NPLs were inconclusive. The NPL shape was considerably affected by the process, owing to which the surface roughness increased and the narrow optical features of the NPLs were lost. c-ALD is a

room temperature process and leads to gentler particle surface reorganization. In the case of successive ion layer adsorption and reaction (SILAR),¹⁸⁴ precursors of cations and anions are introduced successively; however, unlike SILAR, the process is self-limited, similar to conventional ALD. Because the surface of II–VI NCs is generally cation rich, the method starts with the exposure of the nanoparticles to anions. This process is similar to the phase transfer applied for ligand exchange to prepare conductive inks²⁴ and is generally conducted in polar solvents even though shell growth can be realized in the nonpolar phase.¹⁸¹ Next, the cation precursor is introduced, generally from cadmium halide, for the growth of the CdX shell. These two steps are repeated as many times as necessary to obtain the targeted shell thickness; see Figure 11c–e. In practice, saturation of the shell growth occurs after 4/5 layers. To grow thicker shells, additional washing steps to remove excess precursors are required.

Because of the broader range of the available sulfide precursors and the possibility of conducting the reaction in air, the method was first applied to the growth of CdS shells on HgTe,³³ HgS,¹³⁶ and HgSe^{185,186} cores. In general, the sulfide precursor was Na₂S or (NH₄)₂S. The approach was later extended to other chalcogenides, CdSe¹⁸⁵ and CdTe shells.¹⁸⁷ For Se and especially for Te, air-free conditions are required. In the case of selenide (or telluride), the precursor is generally obtained from elemental Se (or Te) reduced by NaBH₄,^{188,189} although the use of Na₂Se and Na₂Te has also been reported.¹⁸⁷

Methods other than c-ALD have been developed to grow CdX shells on HgX cores to overcome the limitations of c-ALD. In the case of CdSe NPLs, it has been observed that materials with c-ALD-grown shells present lower PL efficiencies than those obtained from high temperature shell growth.^{153,155,156,190} This loss of performance can be attributed to the presence of defects in material grown by c-ALD.^{188,191} To date, alternative shell growth has only been reported using HgSe cores.^{120,169,187} To avoid the exposure of HgX nanoparticles to high temperatures, Deng *et al.* employed mild conditions to grow CdS.¹⁶⁹ Shell growth was conducted at 100 °C, and a reactive form of sulfide (H₂S) was used to balance the small growth kinetics induced by the low temperature. Precursors of Cd (cadmium oleate) and sulfide were introduced successively for SILAR growth. The growth of a CdSe shell using a high temperature method was reported by Shen *et al.*¹⁸⁷ Specifically, shell growth was conducted at 200 °C, while a mixture of cadmium oleate and TOP:Se was slowly injected (see Figure 11f,g). The overall introduction of CdX shells on HgX core NCs led to no or marginal improvements in the PL signal,^{33,136,187} as shown in Figure 11h. The reason for this lack of gain in PL efficiency remains unclear.

To date, most efforts have been focused on the growth of a CdX shell to reduce the number of nonradiative recombination pathways. Goubet *et al.* explored the growth of a narrow bandgap layer of HgTe on HgSe cores;¹²⁰ see Figure 11i–l. In this case, shell growth was achieved from the slow injection of mercury iodide and TOP:Te onto HgSe cores at low temperatures (60–100 °C). The objective was to uncouple the optical absorption and transport properties. The authors designed a heterostructure in which intraband absorption was enabled by the HgSe cores and the HgTe shell facilitated transport. As discussed in the section dedicated to intraband detection, the obtained heterostructure spectrum combined intraband absorption in the HgSe and a type II interband transition with the hole located in HgTe (see Figure 40b). This heterostructure led to an

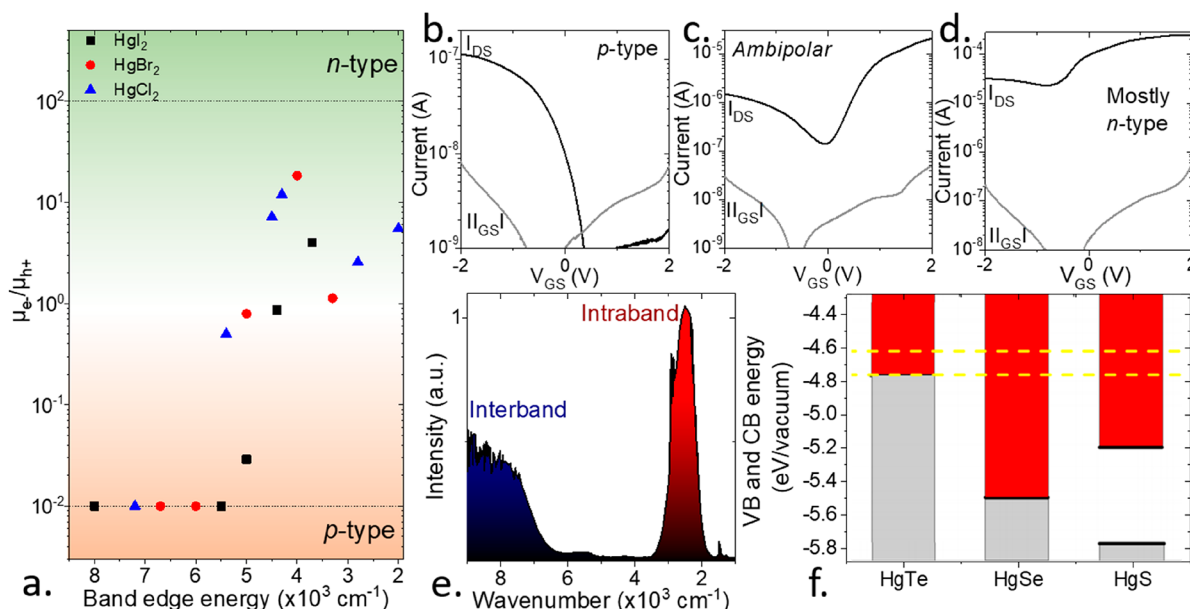


Figure 12. (a) Ratio of the electron and hole mobilities (μ_{e-}/μ_{h+}) for thin films of HgTe with different band edges. When the material presents only p-type conduction, the ratio μ_{e-}/μ_{h+} is set as 10^{-2} . (b,c,d) Transfer curves for electrolyte-gated thin films of HgTe CQDs with band edges at 7000, 4000, and 700 cm^{-1} , respectively. (a–d) Adapted with permission from ref 192. Copyright 2018 American Chemical Society. (e) Infrared spectrum of HgSe NCs. (f) Energies of the valence band (gray volume) and conduction band (red volume) for HgTe, HgSe, and HgS. The yellow dashed line shows the range of the Fermi level measured through electrochemical approaches. (f) Reproduced with permission from ref 68. Copyright 2017 American Chemical Society.

enhanced detection performance of MWIR devices (reduced dark current, increased thermal activation energy, and shorter response time) compared with that achieved using the HgSe core only material (see Figure 40c,d).

2.6. Doping

The electronic structure of mercury chalcogenides is often defined in terms of the inverted band ordering and gapless nature of the bulk material. The influence of the narrow bandgap of mercury chalcogenides on the carrier density and, consequently, on the optical and transport properties of such materials has not been extensively discussed. The two latter aspects are discussed later in this review. However, it is critical to understand the origin of the large carrier density in HgX nanocrystals and the mechanism to realize tuning of the doping magnitude.

Specifically, the high carrier density in HgX NCs cannot be attributed to the introduction of extrinsic impurities. The high carrier density results from two processes. The first process is self-doping (*i.e.*, doping that occurs without the introduction of extrinsic impurities), which can lead to degenerate doping. The Fermi level lies within the conduction band, resulting in one or more free carriers per NC. This doping mechanism is specific for narrow bandgap materials, as discussed further in the next section. The second process is observed in small (diameter less than 12 nm) HgTe NCs and corresponds to thermal activation. For example, assuming a degeneracy of two for the conduction band 1S state and assuming a spherical particle with random close packing (film compactness of 0.64), the population of the

1S state is given by $n_{1S} = \frac{2 \times 0.64}{4 / 3 \pi R^3} \exp\left(-\frac{E_G}{2k_B T}\right)$, where $k_B T$ is the thermal energy (25 meV at room temperature). This expression assumes no doping, with the Fermi level positioned in the middle of the bandgap. At $R = 6\text{ nm}$, $10^{16}\text{ carriers} \cdot \text{cm}^{-3}$ (or 1 carrier per 100 NCs) are obtained for HgTe with a 250 meV (5

μm or 2000 cm^{-1}) bandgap, while for CdSe, under the same particle size and same degeneracy, $6\text{ carriers} \cdot \text{cm}^{-3}$ (or 9×10^{18} per NC) are obtained. This aspect is a major difference in the two materials: even if a perfectly intrinsic material is assumed, the carrier density of HgTe is 10^{16} times higher than that of a conventional wide bandgap material with optical features in the visible region.

In addition, thermal activation originates from material nonstoichiometry. When using various synthetic procedures and precursors,^{30,96,192} small HgTe NCs present a p-type characteristic; see the transistor measurement exhibiting only hole conduction in Figure 12b. When the size is increased, a switch toward ambipolar conduction occurs, and finally, n-type characteristics occur for the largest particles (see Figure 12c,d). The threshold of this transition (see Figure 12a) depends on the growth process.⁹⁶ The exact mechanism of this transition remains unclear; however, the mechanism can likely be correlated with the change in shape observed for HgTe with size. The difference in the exposed facets might induce a change in the amount of excess Hg cation surface with the particle size.

In HgX NCs, a second doping process occurs that leads to degenerate doping. This process is defined as self-doping because high carrier densities (>1 carrier per NC) can be achieved without introducing external impurities. Self-doping is a specific behavior limited to narrow bandgap materials. Self-doping plays a key role because it leads to intraband absorption (Figure 12e) in the steady state.^{65,66} This phenomenon allows the design of IR detectors for wavelengths larger than $3\text{ }\mu\text{m}$.^{34,47,120} The intraband absorption signature is a relatively narrow peak, typically in the mid-infrared region (see Figure 12e). This peak results from the optical transition within the conduction band between the $1S_e$ and $1P_e$ states and was first observed for HgS NCs.^{33,193,194} This intraband absorption, which is the signature of degenerate doping, was likely present in even older syntheses of HgS and HgSe NCs but was missed by

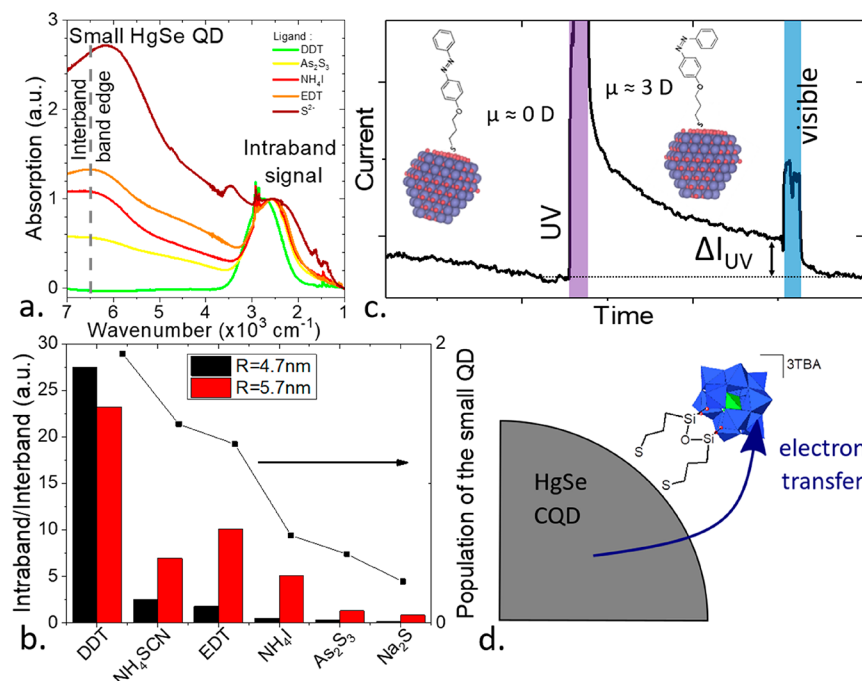


Figure 13. (a) Absorption spectra of small (4.7 nm) HgSe nanocrystals capped with different ligands. (b) Plot of the ratio of the intraband signal divided by the fitted magnitude of the interband signal for two sizes of HgSe nanocrystals. The related population of the CQDs is given on the right scale for small CQDs (diameter of 4.7 nm). (a,b) Adapted with permission from ref 186. Copyright 2016 American Chemical Society. (c) Current in an azobenzene-capped HgTe film as a function of time. UV ($\lambda = 365 \text{ nm}$) or blue light ($\lambda = 460 \text{ nm}$) is projected on the film during the intervals highlighted with purple and blue rectangles, respectively. ΔI_{UV} is the change in the current as the azobenzene switches the configuration. (c) Adapted with permission from ref 197. Copyright 2019 American Chemical Society. (d) Scheme of HgSe capped with POM-SH ligands. The arrows correspond to the electron transfer from doped HgSe QDs to the electron attractor POM-SH ligand. (d) Adapted with permission from ref 204. Copyright 2018 American Chemical Society.

researchers, who focused on visible and near IR absorption. HgSe exhibits a similar intraband absorption.^{29,34,118,124} Such intraband absorption was also later reported for HgTe;^{30,61,63,96} however, fairly large HgTe NCs are required to observe this effect (>10 nm typically). The band alignment of the bulk HgX material, observed through electrochemical approaches,⁶⁸ as shown in Figure 12f, allows the rationalization of these observations. HgX exhibits a fairly large work function (>5 eV for the bulk material), with a value of $4.6 \pm 0.1 \text{ eV}$, as determined through electrochemistry- and photoemission-based methods for short thiol-capped HgX NCs.^{49,122,192} Consequently, the bottom of the conduction band in this material is below the Fermi level. Thus, even in the presence of quantum confinement, the 1S_e state can be found below the Fermi level, and it becomes populated. The stable form of the QDs is negatively charged. In this sense, self-doping can be considered to be a reduction of the NCs by their environment.¹⁸⁶ The large band offset between HgSe and HgTe can explain the difference in behavior between the two materials. The conduction band of HgSe lies comparatively lower than the Fermi level, and only strongly confined HgSe NCs (*i.e.*, size less than 3 nm) exhibit the 1S_e state above the Fermi level (*i.e.*, no intraband absorption feature exists). For HgTe, the conduction band is relatively close to the Fermi level, and thus, only the weakly confined particles can exhibit intraband absorption. HgS NCs exhibit an intermediate behavior because of the presence of a nonzero bulk bandgap.

It was clarified that this inherent doping could be used to control and tune the magnitude of the carrier density postsynthesis. Similar to what has been observed for PbS nanocrystals,^{195,196} the NC surface chemistry affects not only

the interparticle coupling but also the electronic spectrum in terms of the dipole μ associated with the surface ligands, leading to a surface gating effect. The induced band bending is defined as $\Delta E_{\text{vac}} = -N \frac{\mu_{\text{ligand}}}{\epsilon_0 \epsilon_{\text{ligand}}}$, where N is the surface density of the

dipoles and ϵ_{ligand} is the dielectric constant of the ligands. This induced shift is not related to the bandgap of the nanocrystal. Consequently, the induced relative ligand shift of the Fermi level with respect to the bands is considerably more dramatic for narrow bandgap materials. A series of ligands compatible with transport (*i.e.*, typically short thiol-based ligands that strongly bond to the surface Hg) were investigated for HgTe and HgSe (see Figures 23c and 13a–b, respectively). For small HgTe NCs (which have a band edge energy of approximately 720 meV), as depicted in Figure 23c, ligand exchange allows the tuning of the Fermi level energy within the bandgap of the material. Furthermore, this shift is sufficiently strong to induce a change in the majority carriers. Amine and thiocyanate lead to *n*-type NCs, while ethanedithiol or chloride lead to *p*-type NCs. For HgSe, as shown in Figure 13a, the effect is even more dramatic. While the absorption spectrum of PbS¹⁹⁵ or HgTe⁴⁹ is not considerably influenced by the ligand exchange process (generally, only a small red-shift and band edge broadening are observed), the optical properties of HgSe are notably affected.¹⁸⁶ HgSe NC absorption is strongly modulated, as the Fermi level is displaced from within the conduction band to the bandgap. In this context, ligand exchange is a successful strategy to tune the doping level, typically over 1 order of magnitude, as shown in Figure 13b, by using only commercially available ligands that are compatible with charge transport.

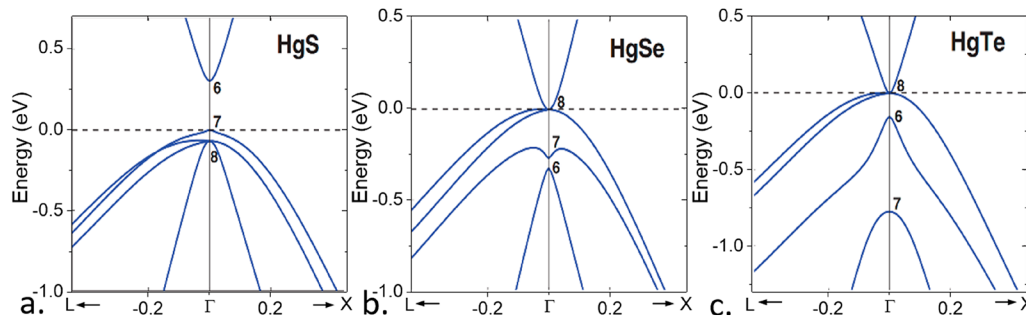


Figure 14. (a) Band structure of HgS near the Γ point from a hybrid QSGW approach. The unit along the x axis is $2\pi/a$, with $a = 0.584$ nm. (b) Band structure of HgSe near the Γ point from a hybrid QSGW approach. The unit along the x axis is $2\pi/a$, with $a = 0.608$ nm. (c) Band structure of HgTe near the Γ point from a hybrid QSGW approach. The unit along the x axis is $2\pi/a$, with $a = 0.647$ nm. Reproduced with permission from ref 70. Copyright 2011 by the American Physical Society.

However, the dipole-induced tunability of the ligands is insufficient. To overcome this issue, Martinez *et al.* proposed using azobenzene (AZB) as a ligand.¹⁹⁷ AZB is a small organic molecule with two conformations (*cis* and *trans*) of the aromatic group around the N=N bond.^{198–200} Under UV and visible light illumination, the molecule can switch its conformation (from *trans* to *cis*, and vice versa, under UV light and visible light or heating, respectively). This change in conformation is commonly implemented to design molecular motors.^{201,202} Instead of the mechanical aspect, Martinez *et al.* exploited the change in dipole resulting from the conformation shift. Typically, the dipole of the *cis* configuration is larger than that of the *trans* conformation.²⁰³ To graft AZBs onto HgX NCs, a short thiol chain was added, whose role was to bond to the Hg surface, as shown in the inset of Figure 13c. Using this approach, the authors demonstrated that the dark conduction and photoresponse magnitude in the IR range can be controlled via UV and blue light stimulation (see Figure 13c).¹⁹⁷ Note that despite the length of the thiol-functionalized AZBs, the HgX NC film functionalized by AZB is still conductive.

To further change the carrier density of the HgX NCs, Martinez *et al.*²⁰⁴ proposed grafting redox groups onto the surface. In practice, the authors used a highly oxidized form of tungsten stabilized in a polyoxometalate (POM). POMs are strong electron attractors, and by coupling these attractors to electron-rich NCs, a redox reaction occurs between the two groups. The authors used the same strategy as that for AZBs to graft the POMs onto the NCs,²⁰⁵ functionalizing the POMs with a short thiol chain used as a linker,²⁰⁴ as shown in Figure 13d. Electron transfer from the HgX NCs to the POMs was evidenced by the reduction of the intraband absorption of the NCs, and the reduced form of W (*i.e.*, with an oxidation state lower than + VI) was detected through photoemission measurements. POMs are strong electron attractors, and up to five electrons per particle can be transferred with POM functionalization.

To ensure the completeness of the discussion on doping, although very few researchers have focused on the introduction of external impurities in the case of HgX NCs, it must be mentioned that Liu *et al.* demonstrated Mn doping of HgS nanoparticles.²⁰⁶ The authors introduced Mn (up to 15%) into the HgS matrix and observed an enhanced magnetoresistance with a higher Mn content.

Despite the notable progress that has been made to enhance the knowledge of doping and its tunability, the amount of work dedicated to doping in HgX NCs remains inadequate. This lack of information is detrimental for the design of devices with a

vertical geometry. Better control of the carrier density can facilitate the design of p-n junctions with higher built-in electric fields and enhanced charge extraction. Diverse routes can be explored, including the introduction of impurities with +I or +III redox states within the HgX atomic lattice. The introduction of such impurities might be eased by the recent development of liquid Hg-based synthesis⁹⁶ from which many metals can easily be solubilized. In addition, doping at the NC array level through the introduction of donor and acceptor dopant nanoparticles is a promising approach and has already led to several successes^{207,208} in the case of lead chalcogenides. This pathway might be worth investigating in the case of HgX NCs.

3. ELECTRONIC STRUCTURE AND OPTICAL PROPERTIES

The electronic structure of HgX NCs is unique and responsible for the many attractive properties of these materials. The spectral tunability of HgTe (Figure 2a), without a competing example, is directly connected to the semimetal nature of the bulk material. While this unique electronic structure leads to several opportunities, it is associated with several challenges. In the case of IV and III–V semiconductor groups, considerable research has been dedicated to the study of the electronic structure of the materials, and the associated results are highly accurate. This aspect is not true for II–VI semiconductors, for which, beyond the bandgap value, large discrepancies in the parameters describing the electronic structure can be observed in different studies. The state of HgTe is not the most inferior owing to the effort dedicated to its integration in IR sensors and topological insulator designs. Nevertheless, current knowledge on HgX materials is incomplete. This statement is more relevant when the material is in the nanocrystal form. In addition to the unique bulk properties, quantum confinement and surface chemistry affect the electrical spectrum as well. This aspect prevents the textbook design of colloidal heterostructures and devices, including films of HgX NCs. Despite these difficulties, substantial progress has been made over the past decades owing to the combination of various theoretical ($k\cdot p$, DFT, tight binding) and experimental methods.

This section is organized as follows. We first discuss the electronic spectrum of bulk HgX materials and their associated characteristics. Next, we describe the confinement and systematically discuss the efforts dedicated to model the electronic structure of HgX NCs. A brief section is focused on the optical properties of HgX NCs. Subsequently, we describe the effects of temperature and pressure on the spectroscopic properties of HgX NCs. A section is also dedicated to the experimental

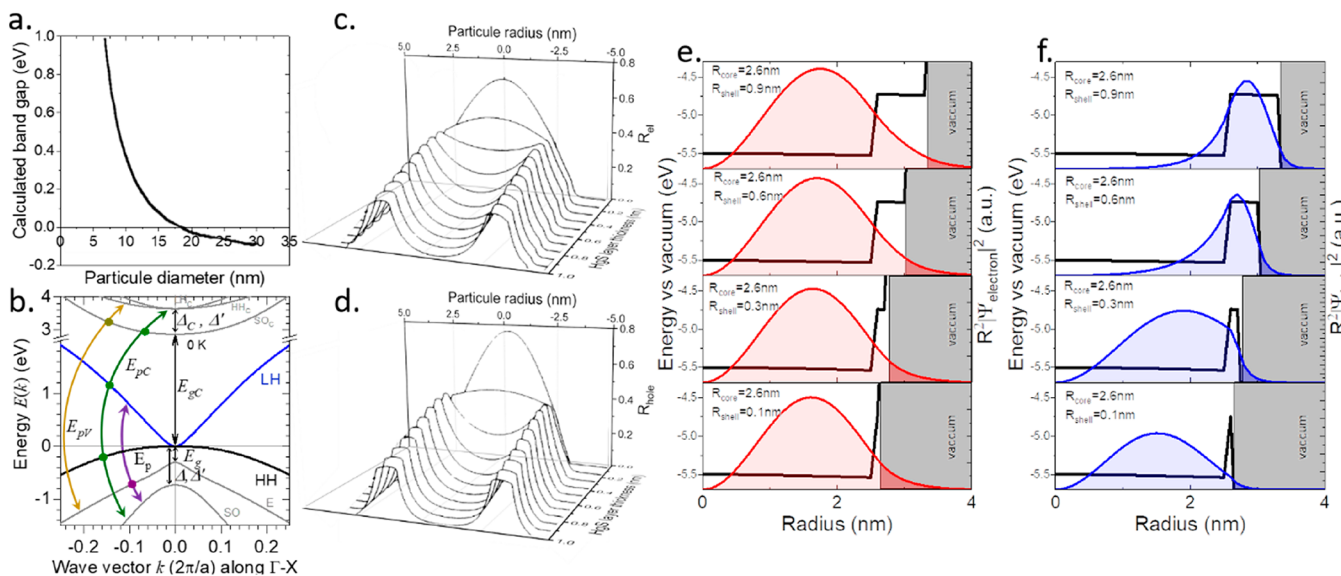


Figure 15. (a) Prediction of the bandgap from the HgTe particle size using the effective mass approximation. (a) Reproduced with permission from ref 93. Copyright 2003 Royal Society of Chemistry, from permission conveyed through the Copyright Clearance Center. (b) Simulated dispersion $E_i(k)$ for the 14 bands of the $k\text{-}p$ model. (b) Adapted with permission from ref 62. Copyright 2020 American Chemical Society. Radial wave functions of the electrons (c) and holes (d) in CdS/HgS/CdS heterostructures with various HgS thicknesses. (c,d) Reproduced with permission from ref 125. Copyright 1994 by the American Physical Society. Conduction band (e) and valence band profiles (f) and electron/hole density distribution in HgSe/HgTe heterostructures for various thicknesses of the HgTe shell. (e,f) Adapted with permission from ref 120. Copyright 2018 American Chemical Society.

determination of the absolute band alignment of HgX NCs, considering the design of heterostructures and devices. Finally, the last section is dedicated to the dynamic aspects of the electronic structure, and we review the experimental methods used to examine carrier relaxation.

3.1. Bulk Material and Confinement

3.1.1. From Bulk HgX to NCs. Conventional wide bandgap II–VI semiconductors, such as CdSe, exhibit a band gap in the visible region. The valence band states (Γ_8 band) are mostly composed of the p orbitals from the chalcogen. This band is strongly degenerated (8-fold) and split into light and heavy hole bands far from the Γ point. The valence band exhibits weak dispersion, corresponding to a large hole effective mass. The conduction band (Γ_6 band) is generally composed of the s orbital of the metal. This band is 2-fold degenerated and is more strongly dispersive than the valence band. Specifically, the associated effective mass is lower. The spin-orbit band is located in the valence band and is responsible for increasing the splitting between the light and heavy hole bands. A general trend is an increase in the spin-orbit coupling as the mass of the chalcogenides increases.

While switching from Cd-based to Hg-based II–VI materials, the band ordering is considerably affected (see Figure 14). HgTe is the most investigated material among the series of HgX compounds because of its use for IR sensing. This material exhibits an inverted band ordering, as observed for gray tin.^{209,210} The band with Γ_6 symmetry appears below the Γ_8 bands (see Figure 14c). This inversion is generally formulated considering a negative bandgap value that describes the relative position of the Γ_6 and Γ_8 bands. This value corresponds to a Γ_6 value below the Fermi level. In HgTe, this negative bandgap is typically approximately -0.3 eV at 0 K,²⁰⁹ and it is reduced by approximately -0.15 eV at room temperature.⁷² Because of the band ordering inversion, the conduction band in HgTe corresponds to the band with light hole symmetry in cadmium

chalcogenides. This band is strongly dispersive and associated with a weak electron effective mass (typically 0.01 – $0.03 m_0$).^{62,72} This low value of the effective mass leads to the large carrier mobility ($>1000 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ and $>10^5 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ at room temperature and low temperature,²¹¹ respectively) observed in bulk HgTe. In contrast, the valence band is almost flat, and the heavy hole mass is larger than $0.3 m_0$.⁷² The Fermi level for the intrinsic HgTe lies between these two bands, which share the same energy at the Γ point. This aspect is the second key characteristic of the bulk HgTe band structure: the degeneracy of the valence and conduction bands at the Γ point causes HgTe to exhibit zero bandgap semiconductor characteristics, corresponding to a semimetal. This lack of a bulk bandgap is the reason for the ultrabroad tunability of the optical spectrum of HgTe NCs (see Figure 2a) compared to that of the other semiconductors, for which the spectral tunability is limited by the material bulk bandgap. In HgTe, the band edge energy can be designed to be arbitrarily narrow, thereby expanding the absorption of this material toward extremely large wavelengths in the infrared region. The optical absorption in HgTe occurs between the less dispersive valence band and conduction band. In this sense, the optical bandgap is always positive and should not be erroneously considered as the negative value mentioned previously, which only describes the band ordering. At the Γ point, the two bands with Γ_8 symmetry are mostly composed of p Te orbitals. Nevertheless, the strong band mixing in HgTe causes the wave functions to acquire the s nature from the Hg atoms as soon as the wave vector is moved away (few % of π/R) from the Γ point.

HgSe presents a band structure similar to that of HgTe, although, quantitatively, the negative bandgap value is lower. HgSe can also be described as a zero bandgap semiconductor with an inverted band ordering. Nevertheless, the lower mass of Se compared to Te reduces the spin-orbit coupling, and the Γ_7 band lies above the Γ_6 band (see Figure 14b). Among HgX

materials, HgS appears to be an exception as a narrow bandgap semiconductor.²¹² The exact value of the bandgap is typically a few hundred meV:^{68,70,213} a value of approximately 0.65 eV matches that reported in spectroscopy experiments with HgS NCs.⁶⁸

Once HgX materials are grown in the NC form, quantum confinement occurs, leading to the following two phenomena. First, the NCs are 0D objects, and the density of the states becomes discrete. Second, the finite size of the particles makes the center part of the Brillouin zone (*i.e.*, close to the Γ point) no longer accessible, and the band edge energy increases (*i.e.*, a bandgap opens). In HgTe and HgSe, due to the lack of a bulk bandgap, the NC band edge energy is purely due to confinement, as shown in Figure 12f. Because of the strong asymmetry of the hole and electron effective masses ($m_h^*/m_e^* > 10$), confinement mostly occurs in the conduction band, owing to which the density of the states in the conduction band is lower than that of the states in the valence band.

3.1.2. Effective Mass Approximation and Multiband k·p Model. The effective mass approximation (or two band k·p model) is the simplest strategy to model the electronic structure of a NC and accounts for the confinement. As a primitive approach, confinement is considered through the evaluation of the wave vector $k = \pi/R$, where R is the particle radius. Next, the energy of this wave vector is evaluated. In this simple model, two assumptions are considered. First, spherical particles are considered, which is often not the case for HgTe; see part a or c of Figure 5. Particles with other shapes may present a different relation between k and the particle size, which is likely not analytic. Second, infinite confinement is considered, which, *a posteriori*, is a reasonable approximation. Such models, have been used to estimate the sizing curve (energy of the optical transition *vs* size),^{93,114} as shown in Figure 15a.

Owing to the ease of parametrization, the effective mass approximation can easily be applied to heterostructures. The time independent Schrodinger equation in a spherical geometry can be expressed as
$$\left[-\frac{\hbar^2}{2m^*} \left(\frac{2}{r} \frac{\partial}{\partial r} + \frac{\partial^2}{\partial r^2} \right) + V(r) \right] \psi(r) = E\psi(r)$$
, where \hbar is the reduced Planck constant, $m^*(r)$ is the effective mass profile, $V(r)$ is the energy profile for the valence and conduction bands, and $\psi(r)$ is the wave function. The material parameters are injected through the mass and energy profiles (bandgap and band alignment) of the bulk. This method is typically used to observe the wave function localization in heterostructures. Moreover, this approach has been applied to the spherical quantum well obtained from the CdS/HgS(/Cds) heterostructure,¹²⁵ as shown in Figure 15c,d, to observe the electron and hole localizations within HgS resulting from a type I band alignment between the two materials. A similar approach has been applied to HgSe/HgTe heterostructures,¹²⁰ as shown in Figure 15e,f. In this case, the two bulk materials exhibit a type II band alignment according to Figure 12f. As a HgTe shell is grown around a HgSe core, the electron wave function remains localized in HgSe. Consequently, the intraband transition resulting from the doping of HgSe is not considerably affected by the presence of the HgTe shell, as shown in Figure 15e. In contrast, the hole wave function is promptly localized in the HgTe shell as it is grown, as shown in Figure 15f.

However, the validity of the effective mass approximation is extremely limited. The 14-band k·p model indicates, as shown in Figure 15b, that the bands in HgTe are only parabolic in a small portion of the Brillouin zone, typically less than 1%. Consequently, the sizing curves based on the effective mass

approximation are only valid for a small range of sizes; see Figure 15a.

When the description of the electronic structure requires the consideration of a broader range of energy, more bands need to be included. A recommendation is to use a 2 + 1-band model,^{114,193} in which the light hole and conduction bands are described by a two-band k·p model and coupled through a Kane parameter, while the heavy hole band is treated as being uncoupled due to its large effective mass. The spin-orbit band, approximately 1 eV below the heavy hole band, is neglected. When a detailed description of the valence band is required, an 8 band k·p model describing the Γ_6 , Γ_7 , and Γ_8 bands and their degeneracies is used.^{72,214} It has been estimated that the 8-band k·p model correctly describes the first 8% of the Brillouin zone.⁷² Although the effective mass approximation only describes the bottom parabolic part of the light hole (Γ_8) band, the inclusion of an additional valence band is critical to capture the linear dispersion occurring at higher wave vectors.

When highly confined forms of HgTe are considered, as in the case for NPLs, it is necessary to account for higher conduction bands (Figure 15b) because the conduction band is responsible for most of the confinement due to the mass asymmetry. In this case, an accurate description of the bands close to the Fermi level but away from the Γ point can be obtained using the 14 band k·p approach.⁶² The inclusion of 14 bands correctly describes $\approx 15\%$ of the Brillouin zone.⁶² The additional upper conduction bands are responsible for the deviation from linearity of the light hole band observed at large wave vectors (Figure 15b), which, as described later in this chapter, is responsible for the inversion of the temperature induced band edge shift with the particle size (see Figure 18).

3.1.3. Tight Binding Method. Tight binding methods are reasonably effective for describing HgTe NCs. The input parameters are less straightforward to determine compared to k·p modeling. However, the method is less demanding in terms of calculation than DFT, and it can be applied to larger NCs and even superlattices.²¹⁵ Allan and Delerue used an $sp^3d^5s^*$ basis to reproduce the electronic spectrum of bulk HgTe and nanocrystals (see Figure 16a,b).²¹⁶ The authors observed that a deviation from the spherical form had a limited impact on the spectrum. Moreover, the authors concluded that Coulombic corrections exerted only marginal contributions to the optical spectrum. Although the excitonic effects are weak at the band edge, HgTe remains an interesting platform to observe excitonic and multiexcitonic effects. The narrow bandgap nature of HgX NCs enables the realization of multiexcitonic effects at relatively low photon energies. Taking into account both energy and momentum conservation, the threshold voltage for the Auger multiplication effect²¹⁷ to occur is $E_{th} = \left(1 + \frac{2 + m_c^* / m_h^*}{1 + m_c^* / m_h^*} \right) E_G$, where E_G is the bandgap and m_c^* and m_h^* are the electron and hole effective masses, respectively. In HgTe, the strong effective mass asymmetry almost suppresses the excess energy due to the momentum conservation, leading to $E_{th} \approx 2E_G$. Experimentally, the threshold for multiexciton generation is slightly higher, approximately three times the bandgap, as obtained using both optical²¹⁸ and transport measurements.²¹⁹

Tight binding was later used to explain the line width of the intraband absorption of HgTe.⁶¹ In HgSe and HgS, the intraband transition appears as a single peak (Figure 12e); for HgTe, the peak corresponds to more than one contribution (see Figure 16c). A naive interpretation may have been to attribute this second contribution to polydispersity. However, this effect

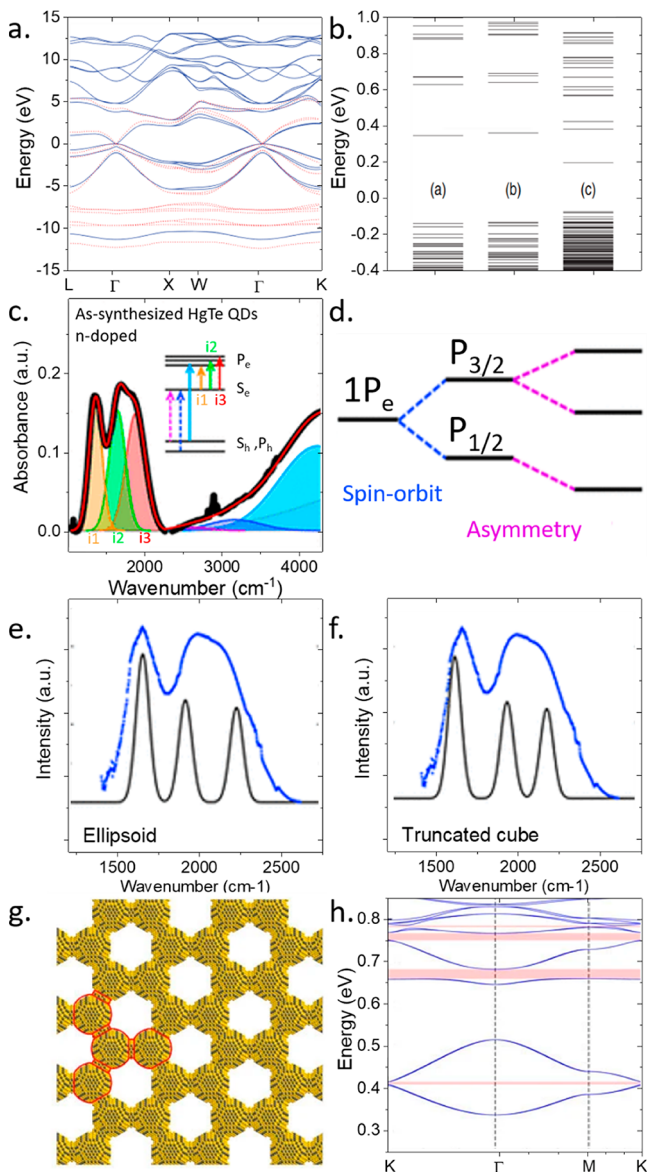


Figure 16. (a) Band structure of HgTe at 0 K calculated in tight binding. The red dots indicate the values obtained using a hybrid GW approach. Zero energy corresponds to the top of the valence band. (b) Calculated energy levels in HgTe QDs at 300 K [(a) sphere, $d = 5.98$ nm; (b) octahedron, $d = 5.83$ nm; (c) sphere, $d = 8.99$ nm]. Zero energy corresponds to the top of the bulk valence band. (a,b) Reproduced with permission from ref 216. Copyright 2012 American Physical Society. (c) Absorption spectrum (black line = experiment) of large doped HgTe NCs. The red line is a fit obtained from a sum of Gaussians and indicates that doping suppresses the first two excitonic transitions and induces intraband absorbance ($i_1 - i_3$). (d) Schematic demonstrating the splitting of the P_e state by both spin–orbit coupling and structural asymmetry. (e,f) Predicted intraband absorbance spectra (black lines) for several shapes of HgTe QDs with a spherical volume with a radius of 4 nm, and comparison with experimental data for a solution of 7.6 nm diameter QDs (blue points). (c–f) Adapted with permission from ref 61. Copyright 2018 American Chemical Society. (g) Top view of an assembly of spheres connected by cylinders that form a honeycomb lattice of HgTe. The Hg and Te atoms are presented in yellow and gray, respectively. (h) Conduction band dispersions resulting from the atomistic tight binding calculation. Nontrivial gaps are indicated by pink shaded regions. (g,h) Adapted with permission from ref 215. Copyright 2015 Nature.

is only observed with an enhancement of the monodispersity of HgTe NC synthesis based on the $(\text{TMS})_2\text{Te}$ precursor.^{63,64} Tight binding simulations clarify that the large spin–orbit coupling in HgTe increases the degeneracy of the $1P_e$ state (see Figure 16d). Because HgSe is lighter, the spin–orbit coupling is weaker, and thus, the $1P_e$ state remains degenerate. A second lifting of the degeneracy also results from crystal field splitting (see Figure 16e,f). Contrary to the interband, which appears to be weakly influenced by the shape, $1P_e$ splitting depends on the exact nanoparticle shape. The power of tight binding modeling is more notable when more complex structures are considered. In 2013, Evers *et al.* reported connected superlattices of PbSe NCs, in which the NCs were necked and arranged into squares and honeycomb lattices.^{220–222} Such self-assemblies provide an opportunity to merge the field of nanocrystal superlattices with that of superlattice transport. For the first time, it becomes possible to combine a high degree of organization with strong coupling, which may facilitate the design of high mobility films.^{223,224} Through development of this material, the growth of materials with a fractal dimensionality of $2 - x$ can be examined, for which the electronic properties can be engineered by tuning both the composition and coupling. Beugeling *et al.* examined the electronic structure resulting from a similar ordered honeycomb lattice made of necked HgTe NCs;²¹⁵ see Figure 16g. The direct synthesis of the structure is likely not be attributed to the HgTe NC solution because the necking process observed for PbSe benefits from the truncated cube shape due to the rock salt structure of PbSe. In addition, such a lattice may be obtained by cation exchange from PbTe preformed superlattices. This artificial graphene is especially promising because, contrary to its carbon-based counterpart, it exhibits a large spin–orbit coupling, which is of considerable interest for spintronic applications. Beugeling *et al.*²¹⁵ observed the formation of a Dirac cone and topological bandgaps as large as 30 meV in such a superlattice (see Figure 16h). Although this work is theoretical, it likely paves the way toward more complex optoelectronic properties beyond the simple tunability of the bandgap with confinement. This work suggests that ordering NCs at the supracrystal level might be a promising strategy to achieve high carrier mobility in NC arrays.

3.1.4. DFT Calculations. Density functional theory (DFT) remains marginally used to describe the electronic structure of HgX NCs. This lack of results mostly reflects the demanding computing power required for this method to be applied to large NCs. Geiregat *et al.*²² used DFT to clarify the origin of the low stimulated emission observed in small HgTe NCs; see the section dedicated to lasing and Figure 27a,b. The authors observed, in particular, long-lived (several μs) PL signals and correlated this observation to the surface trap states (Figure 17b) that can occur in undercoordinated HgTe NCs. DFT shows (Figure 17a) that the resulting gap states are close to the maximum valence band state. The presence of these states eases the population inversion in these NCs by forming a four level system (see Figure 27b).²²

3.2. Optical Properties

One of the most striking properties of HgX NCs is their widely tunable absorption (see Figure 2a). The first optical feature of HgTe NCs can be tuned from 11000 cm^{-1} (880 nm or 1.4 eV) to 150 cm^{-1} (60 μm or 20 meV). The missing information is that the nature of the transition changes as well. The observed optical feature undergoes a change of origin, from interband (smallest NCs) to intraband^{29,33,65} and finally to plasmonic¹³⁶ (for sizes

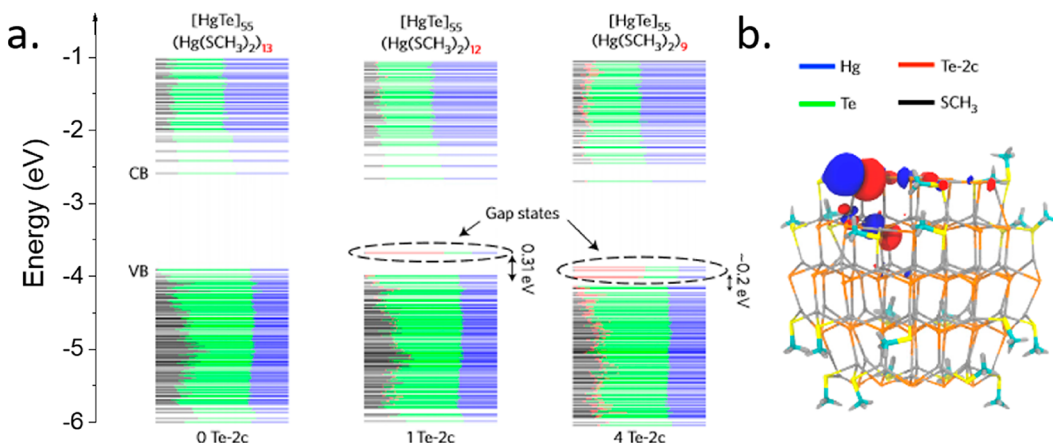


Figure 17. (a) Energy level spectra of (left) a $[HgTe]_{55}(Hg(SCH_3)_2)_{13}$ model nanocrystal, (center) the same nanocrystal after the removal of one $Hg(SCH_3)_2$ complex, and (right) four $Hg(SCH_3)_2$ complexes, calculated using density functional theory. The colors indicate the projection of each state on (blue) Hg atoms, (green) Te atoms, (black) methanethiolate, and (red) dicoordinated Te on different atom/fragment types. The successive removal of one $Hg(SCH_3)_2$ complex results in a band of gap states separated by approximately 100 meV from the highest occupied molecular orbital (HOMO). (b) Wave function of the gap state. This state largely corresponds to the 5p orbital of a two-coordinated tellurium surface atom that is orthogonal to the neighboring Hg 6s orbitals. Reproduced with permission from ref 22. Copyright 2018 Nature.

larger than the Bohr radius) transitions. Note that this process is no longer limited to mercury chalcogenides and has been recently observed for PbS(e) NCs.^{225–227} Because the confinement drives absorption spectrum, the largest particles have the reddest absorption. The doping magnitude is also controlled by the confinement through the relative position of the conduction band with respect to the Fermi level, the highest doping is thus achieved for the largest particles. Following a Drude model, the free carrier density n can be linked to the surface plasma frequency ω_{LSPR} through the simple expression $\omega_{LSPR} \propto \sqrt{n}$. Thus, in conventionally doped semiconductors, higher doping leads to a blue-shift of the plasmonic absorption. This phenomenon is typically observed for ITO nanocrystals (see bottom of Figure 2a) as the tin content increases in the In₂O₃ matrix. In self-doped NCs, the opposite trend is observed and this stress the strong role of confinement in HgX NCs.

A key emerging direction is the control of light matter coupling within films of infrared NCs. As a film of NCs is coupled to a photonic structure, enhanced PL²¹ and absorption^{40,38} can be achieved. Using this approach, the spectral response can be shaped. The absorption spectrum is no longer driven by the material but can be strongly affected by its coupling to light resonators. More details regarding such devices with controlled light–matter coupling are presented in the chapter dedicated to light emission and photodetection. However, a common aspect in these approaches is the requirement for a high level of electromagnetic design. The inputs of these simulations are the geometrical parameters and complex optical indexes. To date, determination of the optical index in the infrared region has not been performed for HgX NC films, which impedes the coupling of NC films to complex photonic structures. This lack of data relates to the low availability of infrared ellipsometers with operations above 1.7 μm . Bejaoui *et al.* performed one of the only investigations related to the measurement of the complex optical index in HgCdTe NC films.²²⁸ However, the NCs are small and include Cd, and the band edge is not in the infrared region. Current simulations⁴⁰ typically assume a constant real part for the NC optical index equal to 2. The real part of the dielectric constant of bulk HgTe at room temperature is approximately ≈ 7 , and this value increases to above 10 as cryogenic temperatures are

reached.^{229,230} A value of 2 for the NC film is consequently lower than $\sqrt{\epsilon}$, which accounts for the presence of remaining organics within the film. This value seems convenient for preliminary design; however, more accurate determination, including spectral dependence, is necessary.

In terms of absorption, the cross-section far from the band edge is a key tool to estimate the amount of crystalline material independently of the size and shape of the particles. For HgTe, the cross-section at 415 nm is $\sigma_{415}(Hg) = 2.6 \pm 0.4 \times 10^{-17} \text{ cm}^2/\text{Hg atom}$.¹¹⁴ The absorption coefficient at the band edge is $3 \times 10^3 \text{ cm}^{-1}$ at 2.6 μm (3900 cm^{-1}) and decreases as larger particles are grown. The value is estimated to be $1.5 \times 10^3 \text{ cm}^{-1}$ at 3.5 μm .¹¹⁴ This value is similar to the absorption observed for bulk HgTe.⁷² This aspect is a major difference from epitaxially grown quantum dots. While epitaxially grown QDs obtained considerable success in terms of quantum engineering^{231,232} and as single photon emitters,^{233,234} films of epitaxially grown quantum dots exhibit weak absorption due to their extremely low density ($10^{11} \text{ QD} \cdot \text{cm}^{-2}$ per plane). Consequently, epitaxially grown quantum dots exhibit considerably weaker absorption than the bulk material. Using NCs, dense films can be fabricated (only limited by the NC packing), and thus, the absorption can be as large as that in bulk. Nevertheless, the absorption coefficient being below 10^4 cm^{-1} , several micrometers of film are required to absorb most of the incident light. At present, such thicknesses are 1 order of magnitude larger than the carrier diffusion length and remain difficult to achieve while maintaining high quality (*i.e.*, smooth and crack free) films. This aspect is the reason for the criticality of the control of light–matter coupling; more details are presented in the section dedicated to this aspect in the last chapter.

In terms of the intraband transitions, the absorption coefficient is between 2 and $3 \times 10^3 \text{ cm}^{-1}$ for 5–6 nm large HgSe NCs,²⁹ corresponding to intraband peaks in the 3–5 μm range. This large absorption is one of the benefits of intraband-based detectors compared to those relying on interband absorption.

3.3. Temperature and Pressure Dependence

The temperature and pressure dependence dramatically affect the spectral response of HgX NCs. The influence of temperature

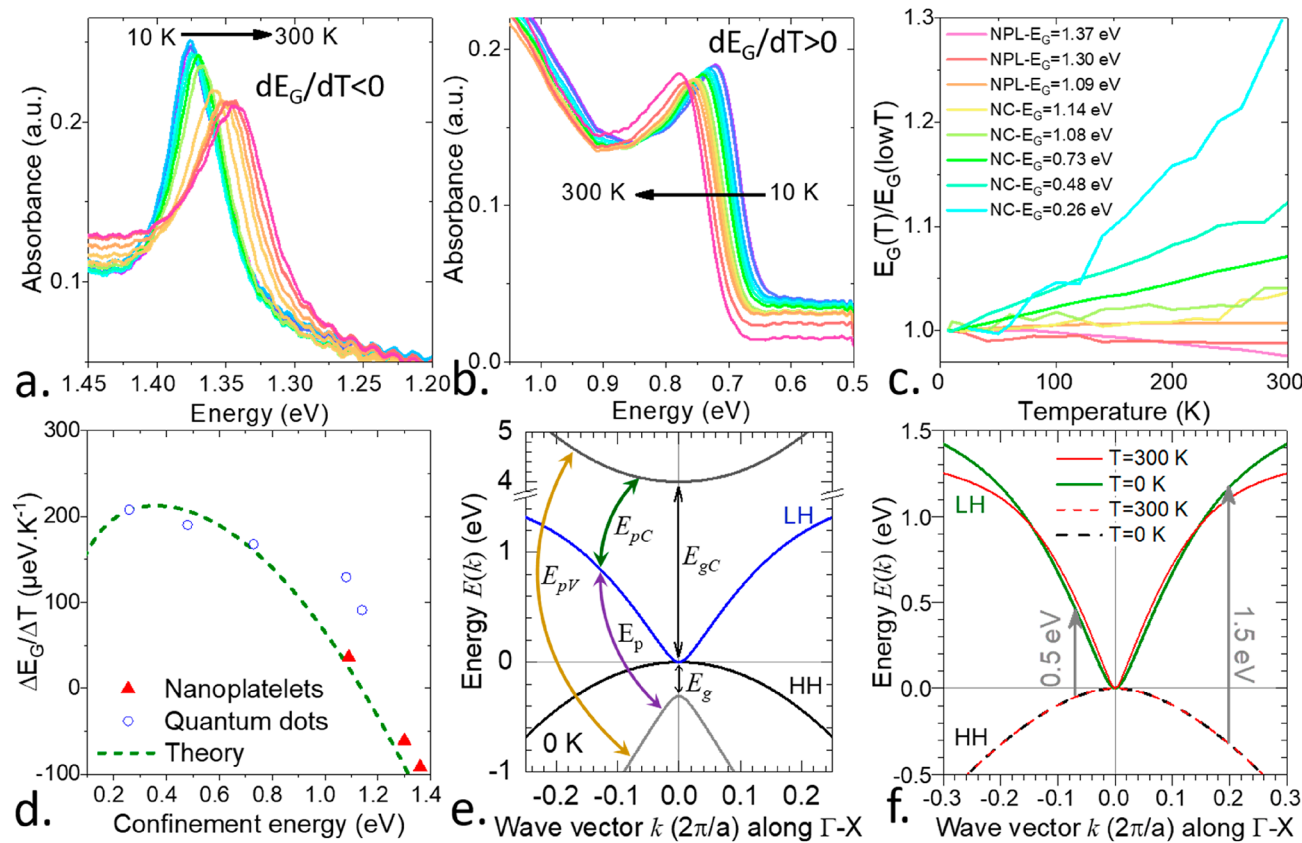


Figure 18. (a) Infrared absorption spectra of HgTe NPLs at various temperatures between 10 and 300 K under zero pressure. The excitonic feature blue-shifts as the temperature is lowered. (b) Absorption spectra of HgTe NCs with a band edge at 6000 cm^{-1} for various temperatures between 10 and 300 K under zero pressure. (c) Experimental shift of the bandgap as a function of temperature for different sizes of HgTe NCs and 3 ML HgTe NPLs with different surface chemistries. (d) dE_G/dT as a function of the confinement energy for the same set of samples. The simulated value of dE_G/dT extracted from the pseudo k-p modeling is plotted as the green dashed line. (e) Dispersion relation $E(k)$ of the conduction band of bulk HgTe extracted from the 3 band pseudo k-p model. (f) Effect of temperature on $E(k)$ when the two bandgaps change with temperature. The gray arrows define the typical confinement energies. The dashed curves are the temperature and pressure independent valence bands assumed in the confinement model. Adapted with permission from ref 62. Copyright 2020 American Chemical Society.

effects on the electronic spectrum is of interest because cryogenic operation is usually implemented for infrared devices. As the bandgap is affected by temperature, the cutoff wavelength can be modified. The effect of pressure can be considered as a nonconventional strategy to tune the NC spectrum. The motivation to examine the pressure dependence originates from the design of colloidal heterostructures, in which the growth of two nonmatched lattice semiconductors generates strain. An independent investigation of the pressure effect allows us to disentangle the changes in the electronic spectrum resulting from strain and that derived from charge delocalization.

3.3.1. Effect of Temperature. In conventional semiconductors (CdTe, CdSe), the trend is to observe a red-shift as the temperature increases. Bulk HgTe is an exception to this rule, exhibiting a positive value for dE_G/dT .^{235–237} HgTe NCs present a similar trend, as reported by Lhuillier *et al.*¹¹⁴ However, HgTe NCs are not the only NCs that present such an inverted temperature dependence, as it has also been reported for lead chalcogenides²³⁸ (PbS) and, more recently, for lead halide perovskites²³⁹ (CsPbBr_3). The first naïve effect of temperature is thermal dilatation. Upon heating, the lattice may expand to affect the particle size and thus the confinement. However, this effect appears to be quantitatively marginal.^{114,238}

This temperature dependence depends on the size of the quantum dots. While HgTe NCs present a blue-shift of the band edge upon heating (see Figure 18b), HgTe NPLs exhibit an inverted trend (see Figure 18a). Nevertheless, this sign inversion in dE_G/dT is not driven by the shape of NPLs but by their confinement. In other words, NCs and NPLs exhibit similar dE_G/dT values for similar band edge energies, as shown in Figure 18c,d. As the confinement increases in HgTe, the value of dE_G/dT decreases, until the sign changes as well (the threshold corresponds to a band edge energy of 1.1 eV). A similar trend is also identified for PbS NCs, for which dE_G/dT decreases and changes in size are observed. This effect is attributed to the size dependence of the electron–phonon coupling.²³⁸

To clarify the mechanism of the coupling of confinement and temperature dependence, Moghaddam *et al.*⁶² used a 14 band k-p model and a simplified 3+1-band model depicted in Figure 18e. The obtained band structure clearly demonstrates the nonparabolicity of the conduction band. The effective mass approximation only holds over <1% of the Brillouin zone. However, for strongly confined NPLs (≈ 1.5 eV of confinement energy), the dispersion relation must be correctly described for large wave vectors. In particular, this model includes the second conduction band, which is >4 eV above the heavy hole band. This band plays a key role. As the temperature is increased, the bandgap E_{gC} shrinks, causing the second conduction band to

approach the first one. To avoid an intersection, the first band bends downward. Because of this temperature dependent curvature, the effect of the temperature on the band edge is different for small and large wave vectors (*i.e.*, for large and small NCs, respectively), as shown in Figure 18f. It can be concluded that the temperature dependence reflects the dispersion relation and that dE_G/dT is directly related to the measurement of the band curvature.

To complete the discussion regarding the temperature dependence on the spectroscopic properties of HgX NCs, it is essential to mention that the intraband transition blue-shifts upon cooling (Figure 19).^{34,122} This shift is also dependent on

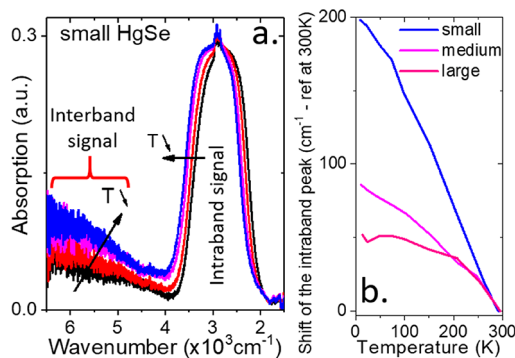


Figure 19. (a) Absorption spectra of small CQDs at temperatures decreasing from 300 to 4 K. (b) Shift of the intraband/plasmonic peak as a function of temperature for three sizes of HgSe CQDs. Adapted with permission from ref 122. Copyright 2017 American Chemical Society.

the NC size, as the shift is larger ($\approx 80 \mu\text{eV}\cdot\text{K}^{-1}$) for smaller NCs. This size dependence is related to the change in the nature of the transition as the size increases. In HgSe, small NCs are less doped and present a true intraband transition,²⁹ while the largest NCs are more degenerately doped and the transition is more plasmonic.¹³⁶

3.3.2. Effect of Pressure. As stated above, the interest in pressure studies is driven by the growth of heterostructures. Over recent years, significant effort has been dedicated to the growth of CdTe/HgTe quantum wells and superlattices to realize topological insulators.^{241,242} Pressure modulation has been identified as a strategy to finely tune the energy landscape of such heterostructures.^{243,244} Similar progress has been made

for the growth of colloidal heterostructures based on HgX NCs; see the section dedicated to heterostructures. This aspect includes shelling with a CdX layer to design type I^{136,185,187} and type II band alignments to enhance the PL efficiency and to realize transport applications¹⁶⁶ and enhanced charge dissociation in HgSe/HgTe intraband detectors, respectively.¹²⁰

The first question raised by the application of pressure is the domain of existence of the zinc blende phase in which HgX NCs crystallize at atmospheric pressure. Bulk HgTe undergoes a transition from zinc blende to cinnabar^{245,246} phase at 1.4 GPa and to the rock salt phase at approximately 8 GPa.²⁴⁷ In HgSe, similar phase changes are observed, albeit with different pressure thresholds.²⁴⁸ The value of 1.4 GPa typically falls in the range of pressures that are induced by the growth of a shell on a core if the two material lattices are not matched.²⁴⁹ Livache *et al.*²⁴⁰ estimated that the growth of a CdS shell on a HgTe core (10% lattice mismatch) should lead to a pressure of 2.6 GPa on the core, enough for the bulk to induce a structural phase transformation. Thus, by applying pressure on HgX NCs, it can be determined whether a change in the spectroscopic properties is the result of a phase change. Livache *et al.* investigated the pressure effect on a series of HgTe and HgSe NCs on both inter- and intraband transitions.²⁴⁰ X-ray diffraction under pressure indicated that the zinc blende phase exists at least up to 3.7 GPa, as shown in Figure 20a. This delay in the phase transition compared to the bulk is attributed to a nucleation barrier.²⁵⁰ Additionally, a similar behavior (within measurement resolution) has been observed for NCs and NPLs.^{62,240} At 4 GPa, the two phases coexist, although it is unclear whether this coexistence occurs within a single NC or if various NCs undergo phase changes at slightly different pressures. By fitting the shift of the diffraction peak in the zinc blende phase (Figure 20b,c), the change in the lattice parameter can be estimated to be 10 pm/GPa, corresponding to a bulk modulus for HgTe NCs of 33 GPa. The same value holds for HgTe NPLs⁶² and corresponds to the bulk value,²⁵¹ suggesting that the ligands do not considerably influence the mechanical properties of the NC film.

To apply pressure, the sample is loaded in a diamond anvil cell in a pressure transmission medium (alcohol or infrared transparent salt for diffraction and infrared spectroscopy, respectively), whose role is to ensure pressure transmission to the sample. In addition, a ruby is inserted within the cell, whose luminescence wavelength is used as a pressure gauge; see the scheme of the setup in Figure 21a. As pressure is applied, the

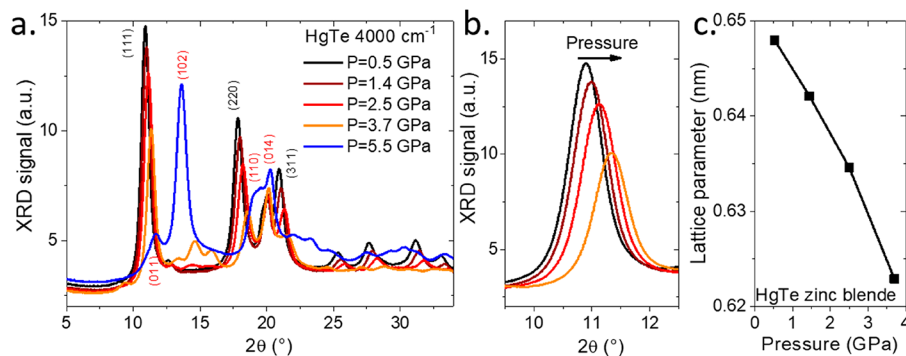


Figure 20. (a) Diffraction pattern of HgTe CQDs (with the band edge at zero pressure being 4000 cm^{-1}) under different pressures. (b) Zoomed-in view of the (111) Bragg reflection of HgTe CQDs (with the band edge at zero pressure being 4000 cm^{-1}) under different pressures while the material remains in the zinc blende phase (*i.e.*, $P < 4$ GPa). (c) Lattice parameter of the zinc blende phase as a function of pressure. Adapted with permission from ref 240. Copyright 2019 American Chemical Society.

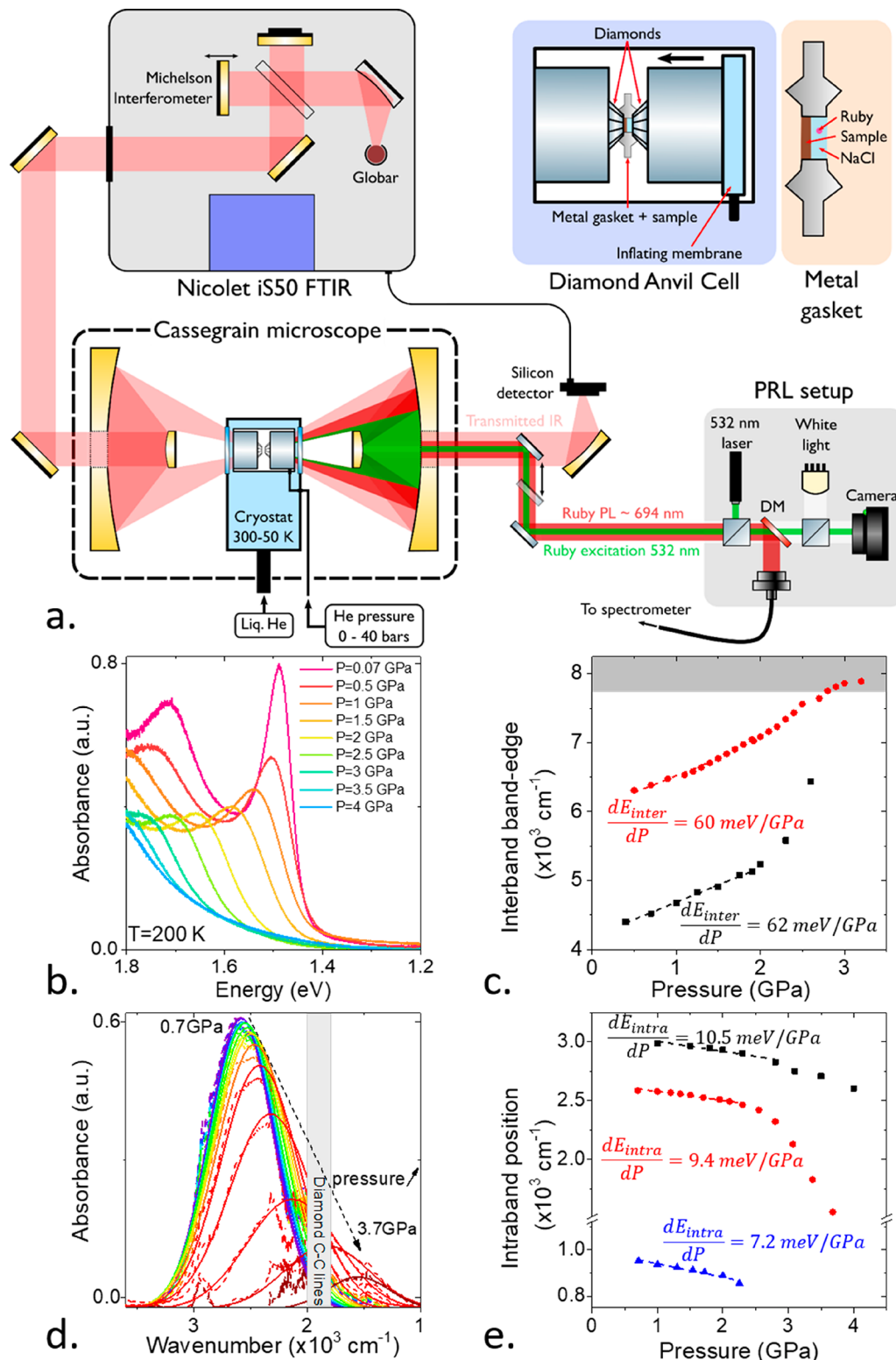


Figure 21. (a) Scheme of the experimental setup used to examine the infrared spectrum of HgTe NPLs under pressure. PRL: pressure according to ruby luminescence. DM: dichroic mirror. (b) Infrared spectrum of HgTe NPLs at 200 K under various pressures ranging from 0 to 4 GPa. (c,e) Adapted from ref 62. Copyright 2020 American Chemical Society. (c) Interband band edge energy as a function of the applied pressure for two populations of HgTe CQDs with band edges of approximately 4000 cm^{-1} (in black) and 6000 cm^{-1} (in red). (d) Infrared spectrum of HgSe CQDs with a zero pressure intraband feature at 2500 cm^{-1} under various pressures. (e) Intraband band edge energy as a function of the applied pressure for three populations of HgSe CQDs (3000 cm^{-1} in black, 2500 cm^{-1} in red, and 1000 cm^{-1} in blue). (c–e) Adapted with permission from ref 240. Copyright 2019 American Chemical Society.

infrared spectrum of HgTe NCs is dramatically affected (see Figure 21b,d). A shift as large as 300 meV can be observed under

4 GPa of pressure. In this sense, the application of pressure is more dramatic than the temperature effect. For the sake of a

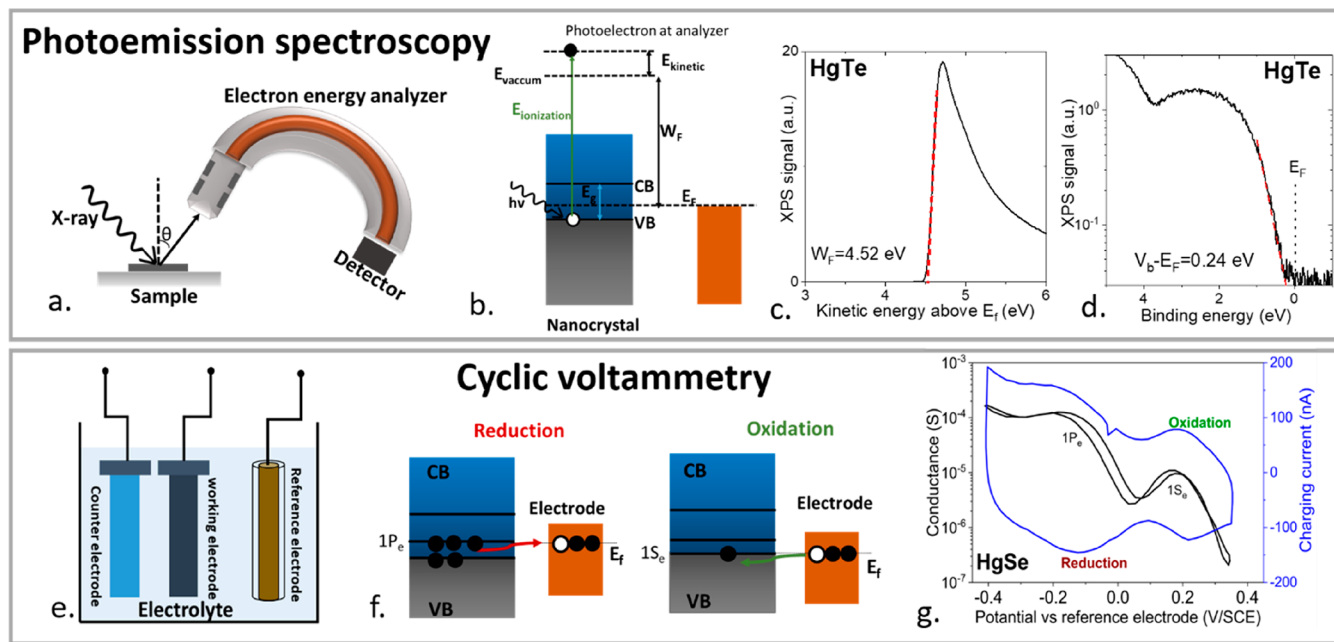


Figure 22. (a) Schematic of the X-ray photoemission experimental technique. (b) Representation of the energy level diagram demonstrating the principle of X-ray photoemission. X-ray photoemission spectrum relative to the cutoff of secondary electrons (c) and those from the valence band (d) for a HgTe nanocrystal film. The band energies are determined from the intersection of an extrapolation from the signal to the baseline. (e) Schematic of the cyclic voltammetry experimental setup. (f) Scheme of the energy level diagram demonstrating the reduction and oxidation mechanism employing cyclic voltammetry. (g) Cyclic voltammetry of a HgSe nanocrystal film. (g) Adapted with permission from ref 68. Copyright 2017 American Chemical Society.

comparison, it can be considered that cooling a HgTe sample by 300 K shifts the band edge by less than 60 meV (see Figure 18b,d). Applying pressure induces a blue-shift of the interband transition (see Figures 21b,c). This shift toward high energy occurs for any confinement value, although the magnitude of the shift is slightly dependent on the confinement energy: 100 meV/GPa for NPLs as opposed to 60 meV/GPa for 8 nm NCs. Using the same (3+1) band k-p model used for temperature, Moghaddam *et al.*⁶² demonstrated that the pressure effect is mostly driven by the bulk Γ_8 - Γ_6 bandgap (E_G in Figure 18e), which starts from a negative value at zero pressure and finally opens up for pressures above 2.6 GPa as the band ordering recovers the normal order (*i.e.*, Γ_6 above the Γ_8 band). Additionally, note that above 4 GPa (in the cinnabar phase), the excitonic transition fully disappears. However, this process is reversible with a strong hysteresis for pressures up to 10 GPa.

Moreover, the intraband transition is influenced by the pressure application; see Figure 21d,e. The shift is the opposite to that observed for the interband transition (*i.e.*, a red-shift upon pressure application). The absolute magnitude of this shift is smaller (<10 meV/GPa). This shift can be attributed to the increased effective mass in the conduction band.²⁴⁰

Distinct behaviors such as the pressure effect, decrease in the magnitude of the transition, and disappearance of the transition induced by phase transformation must be considered for the future design of HgX-based heterostructures.

3.4. Experimental Determination of the Band Alignment

Infrared optical spectroscopy is the most convenient strategy to measure the band edge energy and possibly higher transition energies when sufficiently monodisperse syntheses are employed (see Figure 5b). However, such techniques are differential methods and do not provide insights into the density of the states of each band or absolute band alignment

(*i.e.*, relative position of the band *vs* the vacuum or Fermi level). The latter information is critical to understand the charge delocalization in heterostructures and to design vertical geometry devices. For example, reducing the Schottky barrier at the metal–semiconductor interface is critical to avoid losses. Another example is the measurement of the dipole-induced shift resulting from ligand exchange, which is typically used for the design of p–n junctions. Consequently, additional experimental methods must be developed. Tunnel spectroscopy^{252–255} is one such approach, which can be used to correlate spectroscopic data with structural information. The method has been applied to a variety of nanocrystals, including lead chalcogenides²⁵⁶ and II–VI²⁵⁷ and III–V semiconductors.²⁵⁸ However, to date, tunnel spectroscopy remains mostly unused in the case of HgX NCs. The only report related to on-chip tunnel spectroscopy is discussed in terms of a single particle field effect transistor in the chapter dedicated to transport,²⁵⁹ as shown in Figure 33.

Two methods appear to be of key interest to measure the band alignment of NC materials: photoemission and electrochemistry (see Figure 22). Photoemission is the conventional approach to measure the band alignment of bulk semiconductors.²⁶⁰ Electrochemistry is a more chemistry-oriented method²⁶¹ that exploits the large porosity of NC films. Both methods have been applied to NCs and, in particular, to HgX NCs.^{68,192,122} However, both techniques exhibit certain limitations. The ultrahigh vacuum used for photoemission is not representative of the operating conditions of the device, even though this aspect is more notable for solar cells than for a HgTe NC-based IR sensor that is likely operated under secondary vacuum in a cryostat. Similarly, in the case of electrochemistry, the use of an electrolyte is not representative of the final NC environment in the device. A systematic comparison of the two methods might be useful but has not been reported yet.

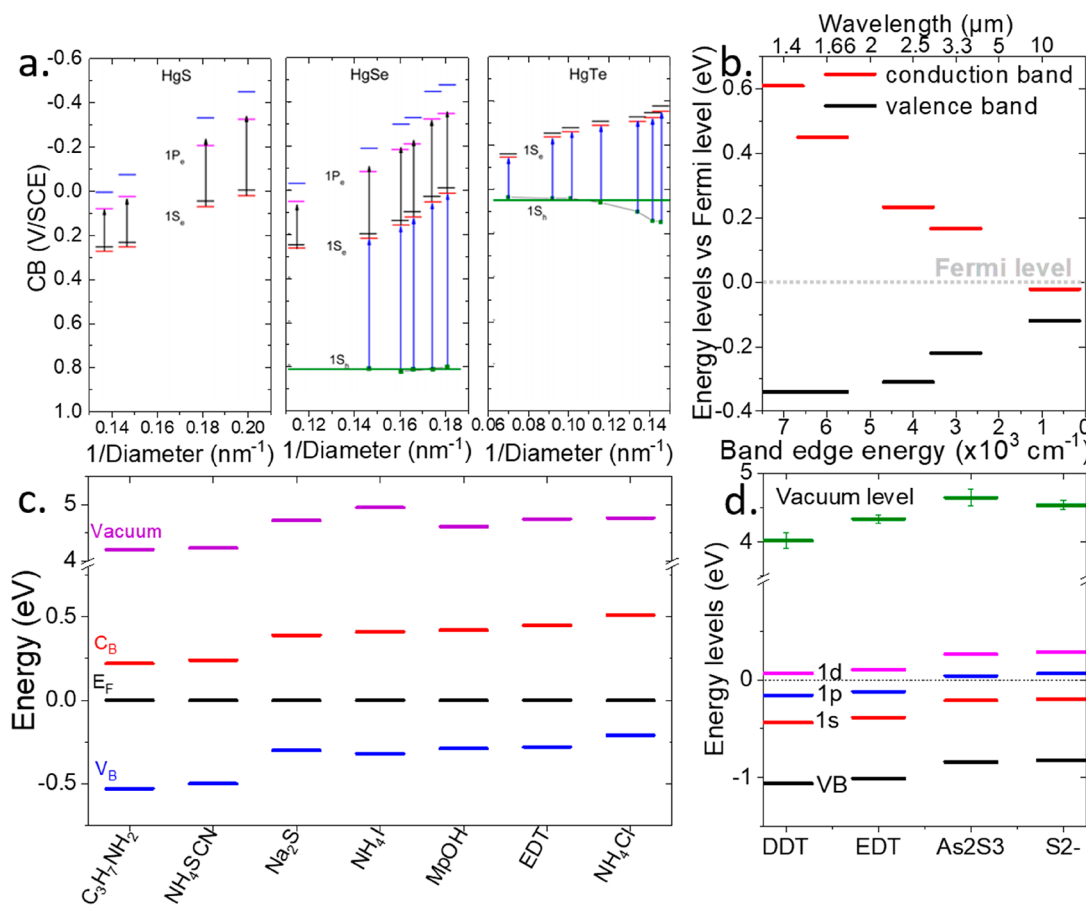


Figure 23. (a) Electronic spectra of the HgX quantum states measured by electrochemistry before (black and blue represent 1S_e and 1P_e, respectively) and after correction by the charging energy (red and pink represent 1S_e and 1P_e, respectively). Intraband 1S_e–1P_e (the length of the black arrow) and interband 1S_h–1S_e (the length of the blue arrow) transitions measured through spectroscopy and corrected for the exciton binding energies are also indicated. (a) Adapted with permission from ref 68. Copyright 2017 American Chemical Society. (b) Adapted with permission from ref 192. Copyright 2018 American Chemical Society. (c) Energy of the valence band, conduction band, and vacuum level for HgTe nanocrystals with a 1.7 μm cutoff wavelength and different capping ligands. The Fermi level is set as 0 eV. (c) Adapted with permission from ref 49. Copyright 2019 American Chemical Society. (d) Electronic spectrum (black, red, blue, pink, and green correspond to the valence band, 1S_e state, 1P_e state, 1D_e state, and vacuum level, respectively) for medium HgSe CQDs capped with four different ligands. Zero on the energy scale corresponds to the Fermi level. (c) Adapted with permission from ref 122. Copyright 2017 American Chemical Society.

3.4.1. Photoemission. To conduct photoemission measurements, a UV or X-ray photon is projected on the sample, with a typical energy higher than the material work function (see Figure 22b). Consequently, a photoelectron is extracted from the sample, and an electromagnetic lens focuses this electron into an analyzer used to determine its kinetic energy (E_K). Owing to the short escape depth of the electrons in the material (a few nm), this experiment must be conducted under ultrahigh vacuum. A scheme of the setup is given in Figure 22a. According to the energy conservation law, $h\nu = E_{\text{BE}} + E_K + \text{WF}_{\text{analyzer}}$, where $h\nu$ is the photon energy and $\text{WF}_{\text{analyzer}}$ is the work function of the analyzer. This equation is used to convert the experimental scale E_K into a material scale given in binding energy E_{BE} .

Large photon energies (1486 eV from the Al K α line for the laboratory setup) are employed to analyze the core levels. X-ray photoemission spectroscopy (XPS) is used to analyze the chemical composition of the samples and determine the oxidation states of each atom. In the case of HgX NCs, the XPS analysis consistently reveals two contributions for the Hg 4f state, resulting from the core and surface Hg excess.^{122,262} Thiol is generally used for the surface chemistry technique, and S is

more electronegative than other chalcogenides. The surface contribution appears smaller and occurs at a higher energy. The XPS signal associated with the chalcogenides generally appears as a single contribution.^{122,262} By tuning the photon energy using synchrotron radiation, the contributions of the bulk and surface can be tuned.

To analyze the valence band, a lower photon energy is generally used, as obtained from a He lamp ($h\nu = 21.2$ eV) in a laboratory or from low photon energy in a synchrotron ($h\nu < 100$ eV typically). This condition is typically employed to determine the work function and position of the valence band with respect to the Fermi level. In photoemission, the Fermi level is assigned as the zero binding energy. In practice, a clean metallic substrate is used to calibrate the energy scale. The low binding energy part of the photoemission spectra is fitted with a Fermi Dirac distribution. If necessary, a correction is introduced to set the Fermi energy to zero, and this correction is later applied to the whole XPS spectra. Alternative calibration methods may also involve a core level of a well-defined binding energy (C 1s or Au 4f are typically used). In a semiconductor such as HgTe, the signal associated with the valence band

appears to shift from the Fermi level (see Figure 22d). One can determine $V_B - E_F$ (*i.e.*, the energy of the valence band with respect to the Fermi level) from a linear extrapolation of the photoemission signal to the baseline. It is critical to conduct this extrapolation, while the photoemission signal is on a log scale; otherwise, the value of $V_B - E_F$ will be systematically overestimated, which makes the semiconductor appear more *n*-type than it is. The other valuable information that is extracted from this spectrum is the work function. The work function is associated with the electrons with the lowest kinetic energies, owing to which an additional external bias must be applied to allow these low kinetic energy electrons to move up to the analyzer. The kinetic energy associated with the cutoff of the secondary electrons is measured (see Figure 22c), and this value is related to the material work function. For HgTe and HgSe NCs, all the reported data^{49,122,192,262} match $\text{WF} = 4.6 \pm 0.1 \text{ eV}$ for EDT-capped NCs.

A critical aspect to obtain reliable measurements is to avoid the charging of the material. As photoelectrons escape from the sample, they leave positive charges behind, inducing an electrostatic shift. To allow rapid charge compensation and avoid this charging effect, NC films are generally deposited on a conductive metallic substrate (typically, gold). Moreover, the thickness of the film is a critical aspect: if the film is not sufficiently thick, then it will be discontinuous and a signal from the substrate will be observed. This aspect is generally not detrimental for core levels but can be problematic when determining the valence band. However, if the film is excessively thick and not sufficiently conductive, then the top part will not be connected to the conductive substrate and will be susceptible to charging. This aspect may lead to the observation of two contributions for each core level: a shifted one from the charged parts of the sample and a nonshifted one for the parts in which the sample is more electrically connected. The sample resistance thus appears to be a major concern. It is critical to process the nanocrystals as if they were used for transport (*i.e.*, to conduct ligand exchange). In this approach, not only is the measurement representative of the actual environment within the device but the charging effect is also strongly reduced. Nevertheless, these concerns must be mitigated for HgX NCs, for which the film resistance is considerably smaller than that for wider bandgap materials and for which charging is not systematic even with weakly electrically coupled NCs.

Once the $V_B - E_F$ and WF values have been determined and the interband band edge energy E_G is known, the energy landscape of a NC population (see Figure 23b–d) can be reconstructed. The valence band and vacuum level are localized with respect to the Fermi level, while the conduction band is localized with respect to the valence band by using the measured optical bandgap. In this case, it is assumed that the Coulombic correction of the optical spectrum is small and negligible, as verified by tight binding simulations.²¹⁶

3.4.2. Electrochemistry. Electrochemistry has long been used as a tool to determine the quantum states of NCs^{263–265} and was first applied to HgTe NCs by Liu *et al.*²⁶⁶ who demonstrated the ambipolar nature of MWIR (5 μm cutoff) HgTe NCs. To conduct electrochemistry, the film of NCs is deposited on a conductive substrate and dipped into an electrolyte, while two additional electrodes control the current and applied potential (see Figure 22e). As the current flows, the NCs become reduced or oxidized, thereby tuning the filling of the surface and quantum states of the NCs (see Figure 22f). A typical cyclic voltammetry diagram is plotted in Figure 22g. The

bumps in the cyclic voltammetry curve are associated with the filling/unfilling of the quantum states. In this measurement, the energy reference is the reference electrode, whose potential must be constant and is later rescaled *vs* vacuum. In this experiment, the Fermi level is the rest potential of the film in the open circuit condition. If two electrodes are present on the substrate of the sample, then it is possible to measure the conductance of the sample and carrier mobility, thereby providing additional information compared to what can be obtained from photoemission. A typical refinement of the cyclic voltammetry experiment is to couple the experiment with spectroscopy to conduct spectroelectrochemistry. In this case, it is possible to observe the bleach of the interband as the intraband peak appears^{193,266} and obtain a more accurate estimation of the quantum state filling as opposed to any other states that might be electrochemically filled and contribute to the electrochemical current.

In regard to HgX NCs, systematic studies have been conducted as a function of the material composition,⁶⁸ size,^{68,192,122} and surface chemistry,^{49,122} as shown in Figure 23. In the event of marginal quantitative disagreements between the electrochemistry and photoemission techniques, the following quantitative trends can be considered for future discussion:

- **Band alignment:** As chalcogenides become heavier (atomic mass), the maximum valence moves toward the vacuum level. Using the electrochemistry approach,⁶⁸ the energies of the top of the valence band are determined to be 5.85 (HgS), 5.5 (HgSe), and 4.77 eV (HgTe) below the vacuum level. The values obtained from photoemission are 5.3 and 4.85 eV for HgSe¹²² and HgTe,¹⁹² respectively, corresponding to a slightly reduced band offset. The trend of the conduction band is slightly more complicated due to the presence of a bandgap in HgS, leading to the bottom of the conduction band energy being 5.2 (HgS), 5.5 (HgSe), and 4.77 eV (HgTe) below the vacuum level.⁶⁸ As mentioned previously, there is reasonable agreement regarding the work function values, with values of 4.7 eV and 4.4–4.6 eV obtained using electrochemistry⁶⁸ and photoemission for both HgSe and HgTe for the same capping ligands (EDT), respectively.
- **Size effect:** Size, through confinement, influences the relative position of the Fermi level and bands. In HgS and HgSe, for sizes above 3 nm, the Fermi level is in the conduction band. This phenomenon corresponds to degenerate doping, which is consistent with the observation of an intraband feature in the infrared spectra of these materials (see Figure 23d). Nevertheless, the Fermi level can be found to be lower in this band depending on the size, barely filling the 1S_e level for the smallest particles, and it is above the 1D_e state for the largest particles,¹²² as shown in Figure 29b. This shift is responsible for a metal–semiconductor transition, as later discussed in the transport part of this review. In the case of HgTe, the Fermi level is almost resonant with the conduction band, and the size effect can be even more dramatic (see Figure 23b). For small confined particles, the Fermi level is within the bandgap, closer to the valence band (*i.e.*, p-type). As the particle size increases, the Fermi level reaches the upper part of the bandgap (n-type) and finally reaches the conduction band as the cutoff

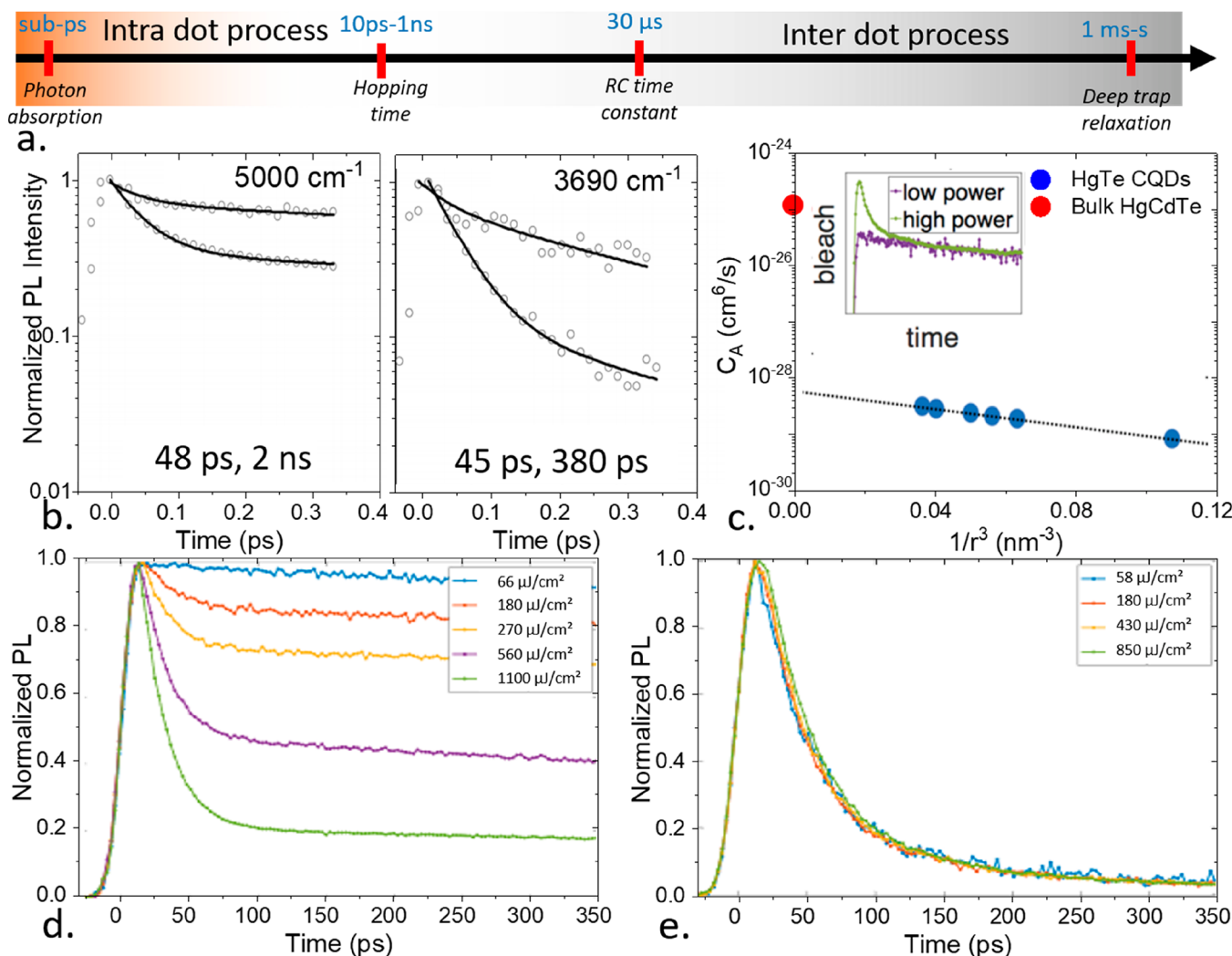


Figure 24. (a) Time scale of the different carrier relaxation times in a film of HgTe NPLs. (b) PL decays for HgTe CQDs emitting at 2 μm (left) and 2.7 μm (right). The open circles denote the measured intensity; the solid lines represent the biexponential fits with the time constants presented as labels. The samples were excited with 1.2 mW (high power) and 0.027 mW (low power). (b) Adapted with permission from ref 168. Copyright 2014 American Chemical Society. (c) Magnitude of the Auger coefficient as a function of the inverse of the particle volume for HgTe NCs. (c) Adapted with permission from ref 268. Copyright 2018 American Chemical Society. (d,e) Interband fluorescence decay at 5600 cm^{-1} for self-doped HgSe NCs. (d,e) Adapted with permission from ref 269. Copyright 2019 American Chemical Society.

wavelength exceeds 5–10 μm (12–20 nm) (see Figure 23b).

- **Surface chemistry effect:** As observed for PbS NCs, the band alignment is affected by the change in the capping ligands; see the section dedicated to doping. The library of ligands that has been tested remains smaller than that reported for PbS^{195,196,267} and is mostly focused on short thiols or short surface ligands compatible with transport (see Figure 29c,d). The shift of the bands induced by the change in the surface chemistry can reach 0.3 eV, which is, given the narrow bandgap nature of the material, of the order of $E_g/2$. A value of 0.3 eV is sufficient to switch HgTe from the n- to p-type (see Figure 29c) or to switch from degenerate doping to a Fermi level within the bandgap in the case of HgSe (see Figure 13a,b).

3.5. Carrier Dynamics

The highly specific size-dependent electronic structures of mercury chalcogenide NCs as well as the semimetal nature of the bulk material are expected to impact carrier relaxation, especially

compared to that of wider bandgap nanocrystals. The stronger density of states is expected to allow efficient phonon-assisted relaxation, and the higher doping level may accelerate Auger relaxation. Carrier relaxation is of interest for both light emission and detection efficiency, and consequently, this is a key aspect to examine in HgX NCs. Time-resolved methods require a setup in which both the source and detection are designed to operate in the infrared region. Time-resolved optical methods, such as time-resolved photoluminescence (TRPL), appear to be well suited for examining processes that occur at short time scales (a few ps to 100 ns), corresponding mostly to intra-NC relaxation (see Figure 24a). Note that time-resolved transient absorption (TA) is mostly unexplored for HgX NCs.²⁷⁰ This aspect is paradoxical from a material viewpoint because the absorption signal does not degrade with increasing wavelength, as opposed to PL. To achieve larger time scales (μs to s), typically to understand the relaxation at the NC array level, alternative methods must be used. In the case of HgX NCs, a transient

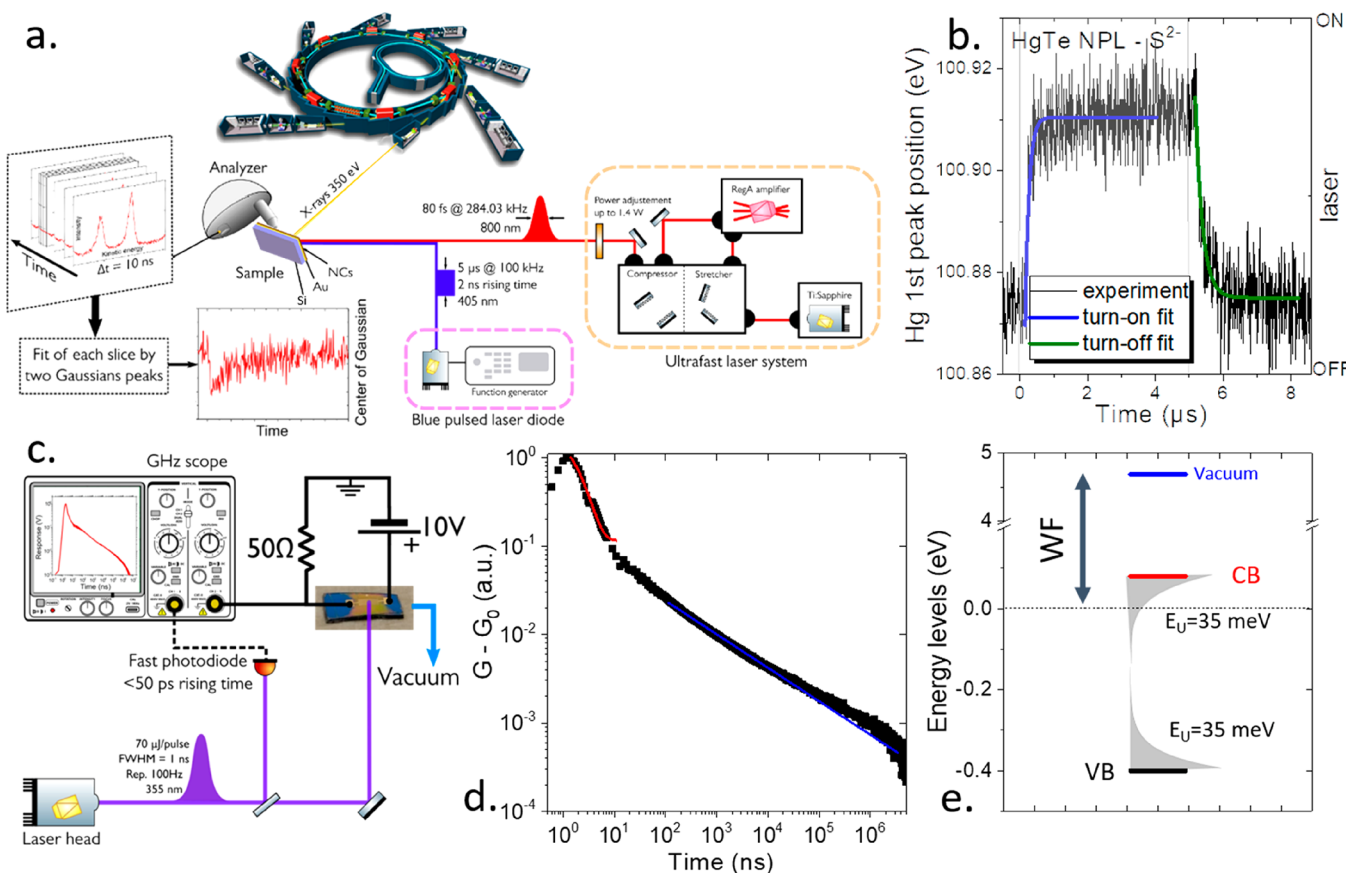


Figure 25. (a) Scheme of the pump probe photoemission setup. Two laser sources can be used: an 80 fs amplified Ti sapphire laser operated at 800 nm or a 2 ns rise time blue (405 nm) laser diode. Light is projected on the sample under ultrahigh vacuum. X-rays at 350 eV sourced from the Soleil synchrotron also illuminate the sample on the same spot. Every 10 ns, a photoemission spectrum is acquired. The time resolution of the experiment is 30 ns, limited by the analyzer. These spectra are fitted by two Gaussians, and the center of the low binding energy Gaussian is plotted as a function of time. (b) The energy of the Hg $4f_{7/2}$ core level fitted position as a function of time, while the light (blue –405 nm laser) is turned on and off for a HgTe NPL film capped with sulfide. (a,b) Adapted with permission from ref 262. Copyright 2017 American Chemical Society. (c) Transient photocurrent setup. (d) Evolution of conductance as a function of time after a 1 ns long pulse of light at 355 nm. (e) Reconstructed energy spectrum including the trap distribution. (c–e) Adapted with permission from ref 278. Copyright 2017 American Chemical Society.

photocurrent and pump probe photoemission have led to interesting results.

3.5.1. Time-Resolved Photoluminescence. TRPL is the most common strategy to measure carrier relaxation in NCs. The success of this approach strongly relies on the ability to achieve single particle measurements. Such time-resolved measurements with a limited number of photons are enabled by fast single photon detectors, such as avalanche photodetectors. However, pushing this experiment in the infrared domain and, in particular, above $1.7 \mu\text{m}$ is highly challenging, and only a few researchers have attempted this expansion.^{21,168} Consequently, there is a lack of consistent data, with PL lifetimes spanning from 50 ps¹⁶⁸ up to several tens of μs .²² The limited amount of available data may lead to unfair comparison of the carrier relaxation times. Moreover, the experimental conditions differ from one study to another with different particle sizes (by at least a factor of 2 based on the diameter) or different excitation pulse durations (ps¹⁶⁸ vs ns²² pulses). At short time scales, a fast component (50 ps) exists, as shown in Figure 24b, whose magnitude strongly depends on the excitation power. This decay time also correlates to the fast component observed in TA (Figure 24d,e) and is attributable to the biexciton lifetime.²¹⁸ At long time scales, an extremely long decay constant has been observed and attributed to trap states, according to

Geiregat *et al.*²² The authors used extremely small HgTe NCs, which increased the relative contribution of the surface. Moreover, the authors proposed using these trap states to design a three-level system, leading to a low excitation power stimulated emission.²² In the intermediate range (100 ps to 100 ns), the contribution from the exciton was identified. However, this contribution decreased promptly with NC size, from 500 ns for a band edge at $1.2 \mu\text{m}$,²² to 10–20 ns for a band edge at $1.7 \mu\text{m}$,²¹ to 2 ns at $2 \mu\text{m}$ and to 400 ps for emission at $2.7 \mu\text{m}$.¹⁶⁸ This rapid drop in the exciton lifetime strongly correlates with the decrease in the PL efficiency with increasing emission wavelengths. This reduction in the exciton lifetime is the result of a higher coupling to the ligands, especially as the PL signal starts to overlap with the absorption of the C–H bonds, enabling efficient near-field energy transfer.^{21,168,271}

3.5.2. Transient PL to Examine the Auger Relaxation. The Auger relaxation of the interband transition in HgTe²⁶⁸ and HgSe²⁶⁹ presents a decay time of approximately 20–30 ps (see Figure 24d). This lifetime follows the volume scaling observed in nanocrystals.^{272,273} However, the relative magnitude of the Auger relaxation quantized using the expression $C_A = \frac{V^2}{8\pi r_B}$, where V is the particle volume and r_B is the biexciton lifetime, is strongly reduced compared to that in the bulk.²⁶⁸ This aspects

can help reducing the number of nonradiative decay paths, which may lead to brighter HgTe NCs and a reduction in the noise magnitude in IR detectors.⁵⁸ A key feature is related to the relaxation of the intraband in HgSe. While the interband decay appears to be similar for HgTe (undoped: $\ll 1$ carrier per NC) and HgSe (doped: ≈ 2 carriers per NC), despite their different carrier densities, the intraband feature displays no or extremely slow Auger relaxation. In addition, the decay constant is independent of the change in the fluence; see Figure 24e.²⁶⁹ This result is counterintuitive, as the Auger value is favored by high carrier densities; however, in this case, the dynamics are retarded for a transition that is induced by the high carrier density of the material. This slowing of the Auger decay is interpreted as the consequence of the density of states in the material. Future challenges in this direction are related to the investigation of the extremely early time scale (<50 ps), which remains unaddressed for HgX NCs to date.

3.5.3. Pump Probe Photoemission. Time-resolved photoemission is a pump–probed technique in which optical photons from a pulsed source are used to excite NCs, and X-ray photons, typically from a synchrotron source, are used as probes.²⁷⁴ A typical scheme of the experimental setup is shown in Figure 25a. In practice, the probe is used to acquire a core level photoemission spectrum at different times of the pump. The obtained core level spectra are then fitted with Gaussians, and the peak binding energy is plotted as a function of time. Because of this fitting procedure, a super-resolution of ≈ 5 meV is obtained, which is an enhanced resolution compared to that (30–50 meV) obtained by static X-ray photoemission measurements. This method can be seen as a time-resolved transient photovoltage experiment that examines the surface band bending at the interface.²⁷⁵ This approach represents a powerful *in situ* method to probe the band alignment in optoelectronic devices.^{276,277} Under dark conditions, a nonstoichiometric semiconductor presents surface band bending. Under illumination, electron–hole pairs are generated, and consequently, minority carriers flow toward the surface due to band bending. The band bending is continuously reduced, and under a sufficiently large irradiance, flat band conditions can be obtained. Once the semiconductor returns to dark conditions, the majority carriers from the bulk flow toward the surface and recombine with the minority carriers accumulated on the surface, thereby restoring the initial band bending. This method provides two key pieces of information. (i) First, the nature of the majority carriers can be confirmed. The sign of the photoinduced shift is related to the nature of the majority carriers. An increase in the binding energy under illumination, as observed in Figure 25b, corresponds to the electrons acting as majority carriers.²⁶² In this sense, the method is correlated with field effect transistor (FET) measurements without requiring any electrodes. The second (ii) key piece of information involves the carrier dynamics. The turn on and turn off time is related to the diffusion time of the minority and majority carriers, respectively.¹⁶⁵

Pump probe photoemission has been mostly applied to HgTe NPLs. Livache *et al.* used this approach as a strategy to clarify whether the response time of their device was limited by the material or device geometry.²⁶² Gréboval *et al.* demonstrated that HgTe NCs and NPLs exhibited similar diffusion times for the majority carriers and concluded that the transport was mostly driven by confinement instead of the particle shape.¹⁶⁵

3.5.4. Transient Photocurrent to Examine the Trap Distribution. As NCs exhibit low luminescence in the infrared

region and on targeting time scales longer than a few nanoseconds, alternatives to TRPL and TA must be developed. The use of the transient photocurrent eliminates the constraints regarding the material luminescence and exploits the photoconductivity of HgX NC arrays. Investigation of the transient photocurrent was driven by the determination of the response time for photodetectors.²⁷⁹ Most HgTe NC arrays exhibit a reasonably small response time. In general, responses of microseconds or less are commonly obtained for both photoconductive and photovoltaic devices based on HgTe (see Tables 6 and 7, respectively). Intraband materials, however, present a slow response (millisecond–second range), which likely reflects the bolometric origin of the response for arrays of pristine HgSe NCs.

However, most short-pulsed sources are not available in the infrared region, and excitation occurs far above the band edge energy,²⁷⁹ which may result in a hot electron effect.^{280,281} Livache *et al.* demonstrated that multiexciton generation (MEG) occurs in HgTe illuminated by high photon energies.²¹⁹ Typically, the threshold for MEG is between 3 and 5 times the band edge energy. To correctly examine the dynamics at the band edge, the authors used a pulsed quantum cascade laser, whose emission was resonant with the HgTe NC absorption.²¹⁹ The authors demonstrated that a response time of as small as 20 ns (limited by the QCL) can be obtained from a conductive array of HgTe NCs.

Martinez *et al.* designed a specific setup to examine the transient photocurrent, as depicted in Figure 25c.²⁷⁸ Excitation was realized using a high power source enabling measurement of the current over several orders of magnitude. The excitation pulse was short to access fast dynamics; however, the repetition rate was maintained at a small value to examine the long time scale relaxation. This setup combined both broad temporal and signal dynamics. At short time scales, the authors observed a fast exponential decay of conductance followed by a long tail,²⁸² see Figure 25d. In most setups dedicated to the detector response time, owing to the low signal resolution (1 or 2 orders of magnitude), only the early decay could be accessed before the photocurrent reached its baseline. Clarifying the slow component of the photocurrent required expansion of the signal dynamics by combining a strong excitation with sensitive detection. The hopping mechanism in HgTe NC arrays is estimated to be approximately a few nanoseconds. As a result, no internanoparticle transport can occur, and the current observed is the displacement current.^{280,281} The drop in current reflects electron–hole pair recombination and trapping. At long time scales, the observed power law decay is the signature of multitrapping transport, as previously observed in the case of amorphous materials.^{283,284} As transport starts, the carriers are first trapped and then thermally released. The release time reflects the depth of the trap. Consequently, the power exponent (b) of the decay directly reflects the trap distribution through the relation $b = -1 + \frac{k_b T}{E_U}$, where $k_b T$ is the thermal energy and E_U is the Urbach energy describing the exponential decay of the trap states within the bandgap.²⁸⁵ The transient photocurrent is an alternative to spectroscopic methods to investigate subbandgap states.²⁸⁶ In HgTe, the Urbach energy ranges from 35 to 50 meV depending on the capping ligands.²⁷⁸ A low value for the Urbach energy indicates high surface passivation and can minimize the open circuit losses for photovoltaic devices.²⁸⁷ Combining this determination of the trap state distribution and photoemission measurements, Martinez *et al.* proposed a reconstructed

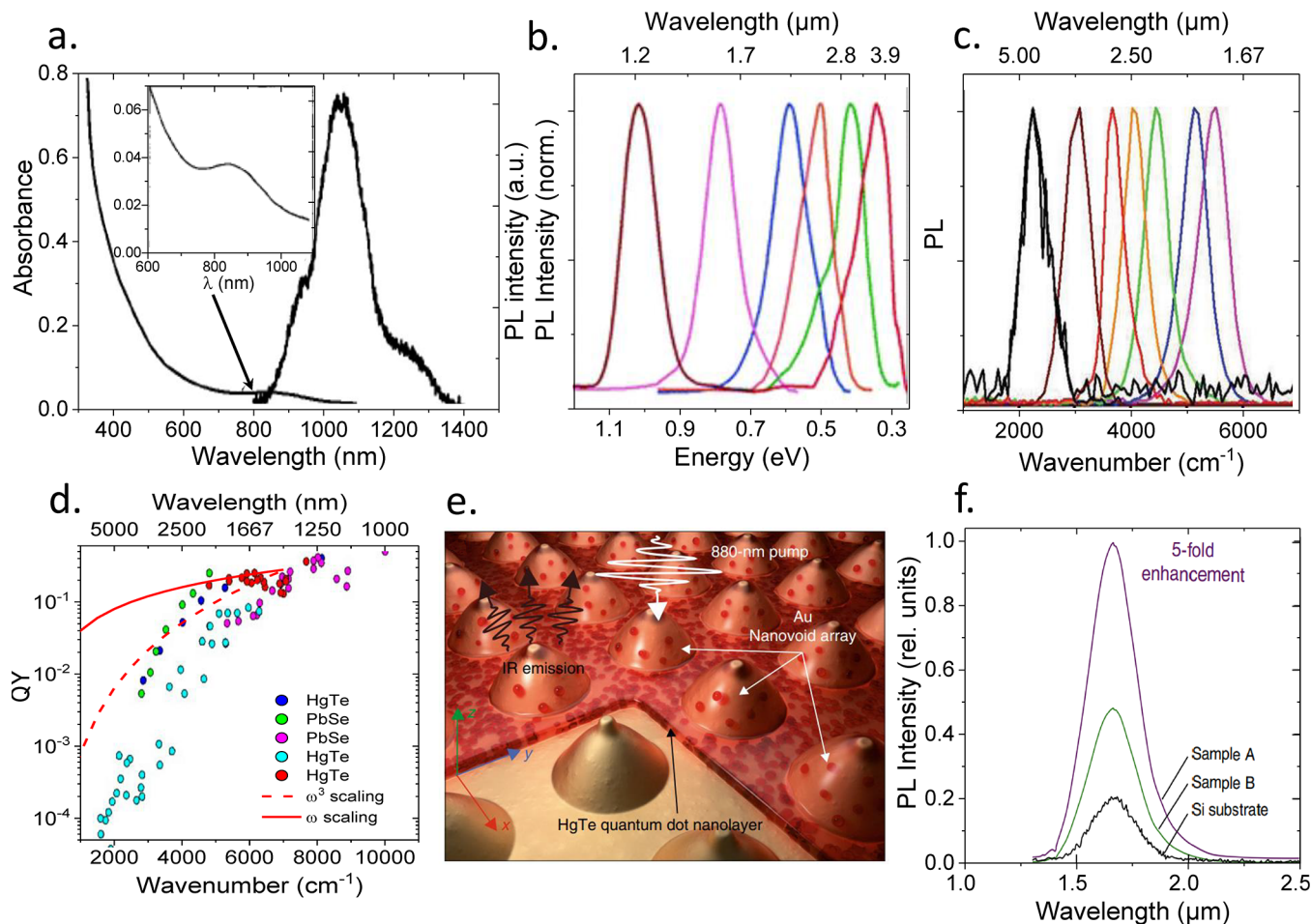


Figure 26. (a) Absorption and photoluminescence spectra of HgTe NCs reported by Rogach *et al.* Adapted with permission from ref 32. Copyright 2001 Wiley. (b) Room temperature PL spectra of HgTe NCs reported by Kovalenko *et al.* (b) Adapted with permission from ref 27. Copyright 2006 American Chemical Society. (c) PL spectra of HgTe NCs reported by Keuleyan *et al.* (c) Adapted with permission from ref 95. Copyright 2011 American Chemical Society. (d) Comparison of the PLQY between HgTe and PbSe with different growth methods. (d) Adapted with permission from ref 101. Copyright 2017 American Chemical Society. (e) Scheme of a HgTe film coated on a laser printed Au nanobump array. (f) Emission spectra of the HgTe NC layer on a Si substrate or on a laser printed Au nanobump array (e). Samples A (purple) and B (green) have different lattice resonances (1480 ± 30 nm for A and 1600 ± 30 nm for B). (e,f) Adapted with permission from ref 21. Copyright 2020 Nature.

spectrum for HgTe NCs, which included the trap state distribution (see Figure 25e).

4. LIGHT EMISSION

Because of their high luminescence in the UV and visible regions, Cd- and Zn-based chalcogenides were the first and most studied materials under NCs. Mercury chalcogenides were introduced later as promising II–VI narrow bandgap materials for infrared light emission.¹⁷⁰ The first findings were reported by Rogach *et al.*, who grew HgTe NCs for the first time,²⁰ as shown in Figure 26a. The authors achieved small HgTe NCs with a tail of absorption in the infrared region. This material was photoluminescent in the near-infrared region and up to the first telecom region ($1.3 \mu\text{m}$). The interest in such near-infrared emitters was, at first, motivated by bioimaging,^{116,129} because the near-infrared window overlaps with the low absorption spectral range of water (and consequently of biological tissues). Light emission was not extensively investigated at the device level for mercury chalcogenides.^{23,288,289} This drawback was not the result of a material limitation but rather due to the difficulty of introducing Hg-based materials for lighting due to their toxicity. A significant part of the work related to the HgX NCs'

PL relates to its use as a tool to examine nonradiative channels. This aspect is used to understand the fundamental limits of NCs occurring through Auger relaxation or coupling to the organic ligands.

This section is divided in two main parts. The first part is related to the photoluminescence in HgX NCs. Intraband luminescence and reports for stimulated emission are discussed in this section. The second part is dedicated to electroluminescence in HgX-based devices.

4.1. Photoluminescence

4.1.1. Interband Luminescence. Following the work of Rogach *et al.*,^{20,115,290} many researchers attempted to increase the emission wavelength. This effort was strongly correlated with synthetic developments. Among the key developments, Kovalenko *et al.* reported an emission up to $\approx 4 \mu\text{m}$ (see Figure 26b) that was extended to $5 \mu\text{m}$ (Figure 26c) and $8 \mu\text{m}$ by adding a regrowth step.²⁸ While further progress was made to strongly red-shift the absorption,^{29,30} $8 \mu\text{m}$ remains the reddest value ever reported for the PL of NCs.

The difficulty of measuring redder PL signals is 2-fold: the lack of an easily available commercial setup and high cost of detectors in the MWIR/LWIR ranges and the intrinsic issue limiting the

Table 4. Figures of Merit Relative to the PL from HgX NCs

material	PL peak (μm)	line width	PLQY (%)	life time	comment	year/ref
HgTe	1.3	$\approx 200 \text{ nm}$	48		synthesis based on thiol capping in aqueous medium	1999 ²⁰
CdHg _{1-x} Te _x	0.8–1.1	$\approx 150\text{--}350$	40		size tunability for NIR luminescence	2001 ³²
HgTe	0.9–2.0	$\approx 200\text{--}400$	50			
HgTe	1.2–3.5 @3.5	≈ 800	40	0.5–2	wide emission from the near to the mid-IR	2006 ²⁷
HgTe	1.3–5 @5	$\approx 1000 \text{ nm}$			narrow luminescence from near and mid-IR	2011 ⁹⁵
HgTe	1.5–5.7 @2	$\approx 300 \text{ nm}$	10–10 ⁻² 2	48 ps (Auger) 2 ns (exciton)	PLQY of mid-IR emission	2014 ¹⁶⁸
HgTe	1.3–2.3 @2.07	$\approx 700 \text{ nm}$	17		aprotic based synthesis	2017 ¹⁰¹
HgTe	1.5–2.7	200–700 nm				two step synthesis using a H ₂ Te gas precursor
HgS	5	$\approx 1150 \text{ nm}$	$10^{-1}\text{--}10^{-2}$		intraband PL	2014 ³³
HgSe/CdS	5.08	$\approx 1150 \text{ nm}$	$10^{-1}\text{--}10^{-2}$		intraband PL	2016 ¹⁶⁹

PLQY of infrared NCs. As depicted in Figure 26d and reported in Table 4, the PLQY rapidly decreases with wavelength. For example, a reasonable PLQY can still be obtained at 2 μm when performing aqueous synthesis. Rogach and co-workers pushed the PLQY up to 17% for an emission of approximately 2 μm .¹⁰¹ However, at 5 μm , the PL efficiency is only a few 0.01%. This drop is caused by the combination of two effects. If we neglect the non-emitting (dark) NCs,²⁹¹ then the PLQY can be expressed as $\text{QY} = \frac{\Gamma_r}{\Gamma_r + \Gamma_{nr}}$, where Γ_r is the radiative emission rate and Γ_{nr} is the nonradiative decay rate, accounting for all the possible loss mechanisms (Auger, phonon, energy transfer, etc.). The radiative lifetime can be expressed as²⁹² $\Gamma_r = \frac{\omega^3}{3\hbar c^3} \mu^2$, where ω is the frequency of emission, \hbar is the reduced Planck constant, c is the light velocity, and μ is the dipole associated with the transition. Van Driel *et al.* reported a different scaling for the radiative decay rate, with the emission frequency presenting a linear dependence.²⁹³ Nevertheless, both the modeling approaches for the radiative decay rate indicated a lower rate as the emission shifted toward longer wavelengths. Thus, if the nonradiative decay rate is considered to be a result of the surface chemistry, then the slowing of the radiative component in the infrared region is primarily responsible for the observed QY loss. Specifically, this phenomenon is responsible for one and a half orders of magnitude decay (in the ω^3 scaling hypothesis) out of the two and a half orders of magnitude decay of the PLQY observed as the emission wavelength increases from 2 to 5 μm .

This intrinsic decay cannot be overcome by loss reduction (*i.e.*, decrease in the nonradiative decay rate). To design faster radiative IR emitters, the dipole remains the only degree of freedom. Quantum engineering can facilitate the development of heterostructures in which the wave functions are designed to increase the dipolar moment of the transition. At present, this strategy remains unexplored for mercury chalcogenides. This approach can be implemented “artificially” by tuning the light matter coupling: recently, Sergeev *et al.* reported the coupling of a HgTe NC film to an array of nanobumps,²¹ as shown in Figure 26e. This array is used to introduce plasmonic resonances and

increase the effective electromagnetic field absorbed by the NCs. The obtained PL is enhanced by a factor of 5, see Figure 26f.

The second limitation to obtain bright IR emitters is also specific to infrared. IR is often seen as part of the electromagnetic spectrum in which chemical bonds present a fingerprint signal: infrared spectroscopy is commonly used as an analytical technique to clarify the chemical composition of a sample. The surface chemistry of NCs leads to a dramatic limitation in regard to the IR emission of NCs because of the strong IR absorption of ligands. At the end of synthesis, NCs are generally capped with ligands that include long alkyl chains. The ligands present a strong IR absorption at $\approx 2900 \text{ cm}^{-1}$ due to the C–H bond vibration. The magnitude of this absorption is often useful to determine whether ligand exchange is successful. Because of the proximity of ligands to the inorganic core, a highly efficient energy transfer occurs from the exciton to the organic shell. This process introduces nonradiative losses in the PL and can be attributed to a 1 order of magnitude drop in the PL efficiency below the radiative limit (see Figure 26d). A possible strategy to overcome this limitation is to use purely inorganic surface chemistry. On HgX NCs, As₂S₃ is, to date, the only fully inorganic ligand that has been grafted onto NC surfaces.³⁵ However, no report exists on the potentially beneficial effects of these inorganic ligands for PL, as these ligands were introduced for electron transport enhancement. The introduction of a core/shell with a type I band alignment is also a viable strategy to enhance the PLQY by preventing the wave functions from experiencing surface traps.^{9,16} As discussed in the section dedicated to heterostructure growth presented earlier in this review, the growth of HgX-based heterostructures is highly challenging because HgX cores are mostly incompatible with the high temperature growth methods required to obtain CdX shells. Among the few reports dedicated to the shelling on HgX cores, mostly toward CdX shells,^{33,136,185,187} the introduction of the shells does not lead to a significant PLQY enhancement.

4.1.2. Intraband Luminescence. While previous discussions have mostly focused on undoped HgX NCs, self-doped HgX NCs can also present intraband PL,^{33,169} as shown

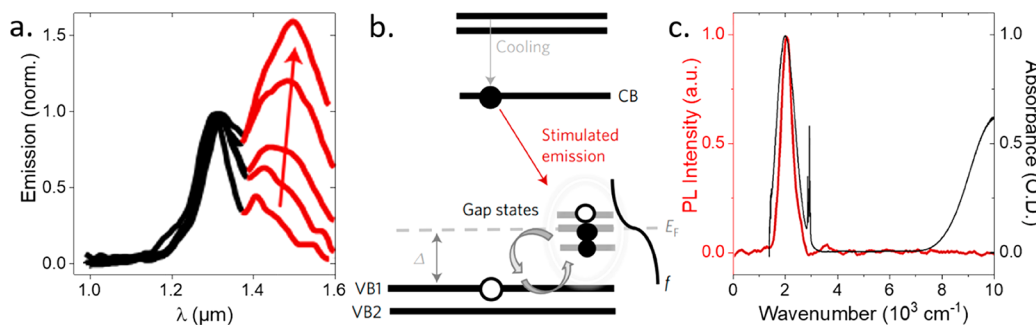


Figure 27. (a) Amplified spontaneous emission spectra of a free-standing HgTe thin film with increasing stripe length (red arrow). (b) A four-level scheme in which the stimulated emission involving an electron from the CB to the empty surface defect levels close to the valence bands ($\text{VB}_{1/2}$) does not have any counterpart in absorption, leading to a thresholdless gain. (a,b) Reproduced with permission from ref 22. Copyright 2018 Nature. (c) Intraband PL emission spectrum (red) from the 1S_e - 1P_e transition of ambient n-type HgS NCs after photoexcitation at 808 nm. The absorption spectrum (in black) overlaps with the PL emission spectrum. (c) Adapted with permission from ref 33. Copyright 2014 American Chemical Society.

in Figure 27c. In this case, the PL signal spectrally overlaps with the intraband absorption without the Stokes shift. Depending on the exact filling of the 1S_e state, which can be modulated with surface chemistry, dual color luminescence attributable to both the inter and intraband absorption can be obtained.¹⁶⁹ The PLQY of the intraband transition is similar to that from the interband transition at the same wavelength.

4.1.3. Lasing. In addition to the PL, lasing of NCs²⁹⁴ has been developed in recent years.^{295–298} To design a cavity, Shopova *et al.* used Si microspheres that couple to Hg(Cd)Te NCs.²⁹⁹ The authors obtained a lasing signal from 1240 to 1780 nm with a low pump threshold.³⁰⁰ In these early reports, short (fs and ps) pulsed excitations were considered to generate a transient gain at times smaller than the Auger lifetime. Nevertheless, the long-term goal is to demonstrate the steady state or quasi (ns) steady state gain to enable broad band light amplification at a telecommunication bandwidth (GHz and more). A step in this direction was reported recently by Geiregat *et al.*²² The authors realized stimulated emission with a low threshold ($40 \text{ mW}\cdot\text{cm}^{-2}$) in small HgTe NCs (see Figure 27a). Such a low level of excitation is an asset because it is compatible with sunlight excitation or low current density operation of LEDs ($20 \text{ mA}\cdot\text{cm}^{-2}$). The authors succeeded in demonstrating a gain using a nanosecond pulsed laser, which is a critical step toward developing electrically driven laser diodes. The authors attributed the origin of the long life gain to the presence of surface traps in the vicinity of the valence band (see Figure 27b). Such traps likely contribute to the formation of a four-level system enabling a long-lived population inversion. These results must be correlated with future developments of HgTe NC-based LEDs.

4.2. Light Emitting Diode

The efforts dedicated to achieving electroluminescence (EL) using mercury chalcogenides are extremely few compared to those for PbS NCs, which is likely correlated to the lower amount of work dedicated to the development of vertical geometry photovoltaic devices based on HgTe^{46,48,146,192,301–304} compared with PbS. In addition, HgS and HgSe are self-doped and present mid-infrared intraband transitions (see the previous section dedicated to intraband luminescence) owing to which the device integration is relatively challenging due to the low PL efficiency.^{33,169} A first structure was proposed by Kotysh *et al.*,²⁸⁸ which involved HgTe NCs mixed into a conjugated polymer to form a blend structure. The latter was surrounded by Al and ITO contacts (see Figure 28a).

EL at 1300 nm was observed; however, the turn on voltage was high, at approximately 10 V (*i.e.*, 10 times the bandgap energy, see Figure 28b), suggesting an unoptimized structure. Later, O’Connor *et al.*²⁸⁹ proposed a new structure based on an ITO/PEDOT:PSS/HgTe/Al stack. This ITO:PEDOT combination is extensively used as a hole injection layer for OLEDs and visible quantum dot-based LEDs. Compared to wider bandgap materials, the narrow bandgap of HgTe reduces the need for a second (or even third) hole transport layer to make the HOMO potential resonant with the emitting layer valence band. The operating bias is significantly reduced, and the turn on voltage is below 2 V when using this structure (see Figure 28c). The later remains far above the bandgap. Since this research, the electronic structure and doping of the materials have undergone considerable progress. HgTe NCs with emission in the near-IR region exhibit a strong p-type nature. The aluminum top contact is easily oxidized into insulating alumina, which leads to an insufficient electron injection. After this demonstration, there was a 15 year period without any report dedicated to EL based on HgTe NCs. Recently, Qu *et al.*²³ proposed a structure inspired by PbS NC solar cells.^{305,306} The authors used a ZnO layer and an EDT capped layer of PbS for the electron and hole transport layers, respectively (see Figure 28d). The emitting layer was made of a ZnO:HgTe mixture. The authors found that the optimal ZnO to HgTe ratio was approximately 1:1. This ratio facilitates the formation of an ambipolar bulk heterojunction while controlling the charge injection by modulating the amount of ZnO as a hole blocking layer. With this structure, a subband gap (0.6 V, see Figure 28e) turn on voltage is achieved. The authors demonstrated the development of a bright LED with a maximum EQE of 0.7%, stable emission over 30 h of continuous operation in air without any LED encapsulation, and spectral tunability from 1250 and 1600 nm (Figure 28g). The authors finally used the bright LED as an SWIR light source for active imaging applications (see Figure 28f). The authors also indicated that moisture detection could be easily realized using such near IR emitting LEDs. One unaddressed question regarding HgX NC LEDs is light extraction. Because of an optical index mismatch, approximately 80% of the generated photons are waveguided into the substrate. Various strategies have been reported to obtain direct emission from HgTe NCs, including the use of Bragg mirrors³⁰⁷ or coupling HgTe NCs to a photonic crystal.³⁰⁸

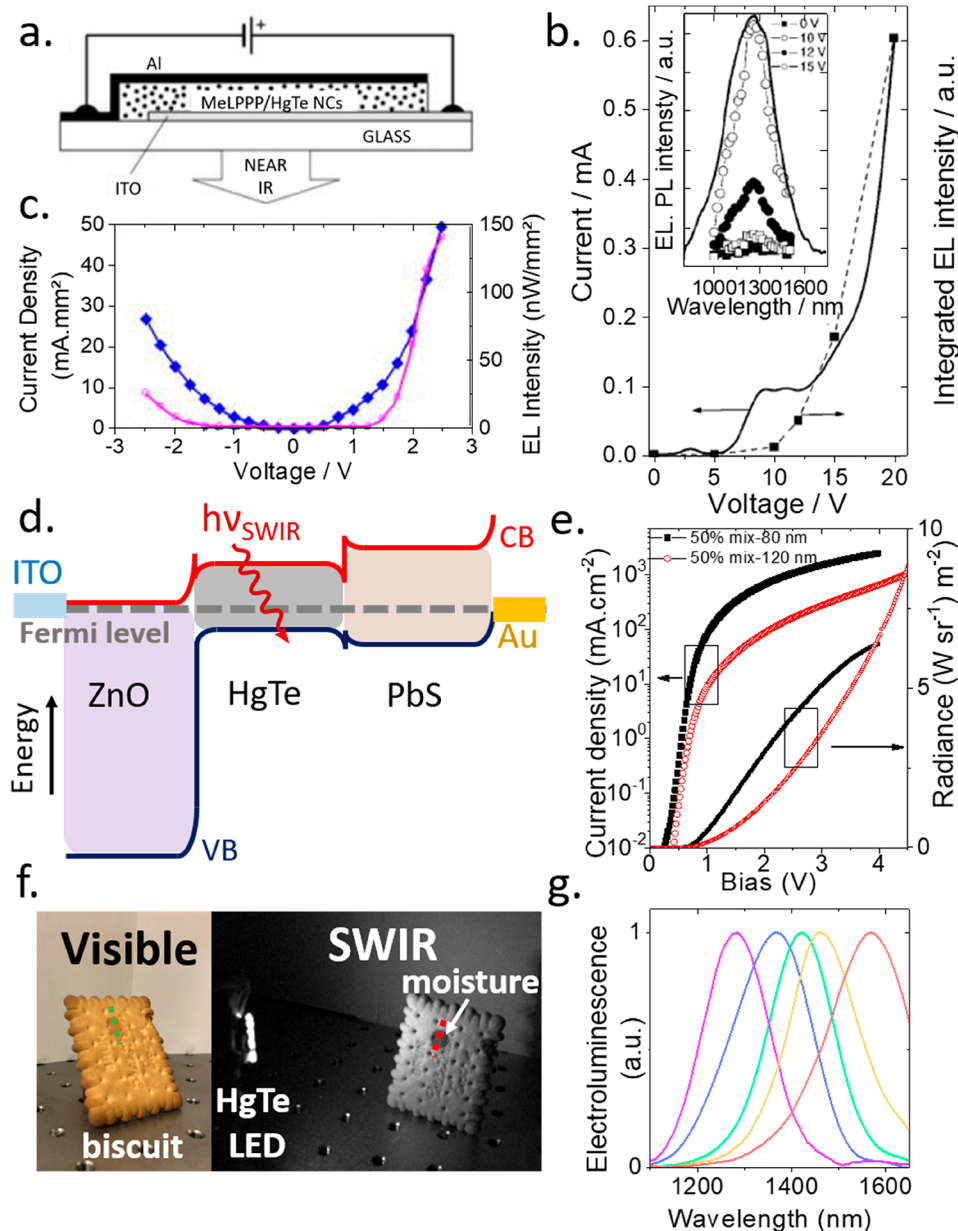


Figure 28. (a) Representation of the MeLPPP/HgTe NC device; ITO, indium tin oxide; NIR light, near-infrared light. (b) Current–voltage (—) and integrated electroluminescence (EL) intensity–voltage (squares connected by a dashed line) traces of the MeLPPP/HgTe NC device. Inset: electroluminescence spectra of the MeLPPP/HgTe NC device obtained at different voltages in comparison with the photoluminescence spectrum of HgTe NCs in a composite MeLPPP/HgTe film. (a,b) Reproduced with permission from ref 288. Copyright 2004 Wiley VCH. (c) Typical I/V/L characteristics of an NIR emissive HgTe quantum dot device (ITO/PEDOT/HgTe/Al). Blue solid diamonds, current; pink open circles, NIR luminance. (c) Reproduced with permission from ref 289. Copyright 2005 AIP Publishing. (d) Energy band profile for a diode made of ITO/ZnO/HgTe/PbS/Au. (e) Current and radiance as a function of the applied bias for two thicknesses of the emitting layer and with a 50:50 volume ratio of HgTe and the ZnO solution. (f) Visible (a) and SWIR (b) images of a biscuit presenting a moisture stain. (g) Normalized electroluminescence spectra obtained from HgTe NCs. (d–g) Adapted with permission from ref 23. Copyright 2020 American Chemical Society.

5. ELECTRONIC TRANSPORT

This section addresses the electronic transport in HgX NCs, mostly in the array form. This review is focused on dark conduction because the next section is dedicated to photoconduction and detection applications of HgX NCs. The chapter is organized as follows. We first discuss the specificity of the electronic conduction in a range of carrier densities (10^{-3} –1 carriers per NC), which is considerably higher than that typically observed for wide bandgap semiconductors but considerably lower than that observed for metals (1 electron per atom). Next, we discuss the effect of the surface chemistry on transport and

stability of the material's electrical properties under atmospheric conditions. Finally, a large part of the discussion is dedicated to the design of field effect transistors and their use in examining electronic transport.

As discussed earlier, a key specificity of HgX NCs with respect to other NCs is their large carrier density. Figure 29a summarizes the main results discussed in the section dedicated to doping. The figure provides a sense of scale for the expected carrier densities in various NCs. The large carrier density originates from a combination of the low carrier activation energy of a narrow bandgap semiconductor and self-doping that

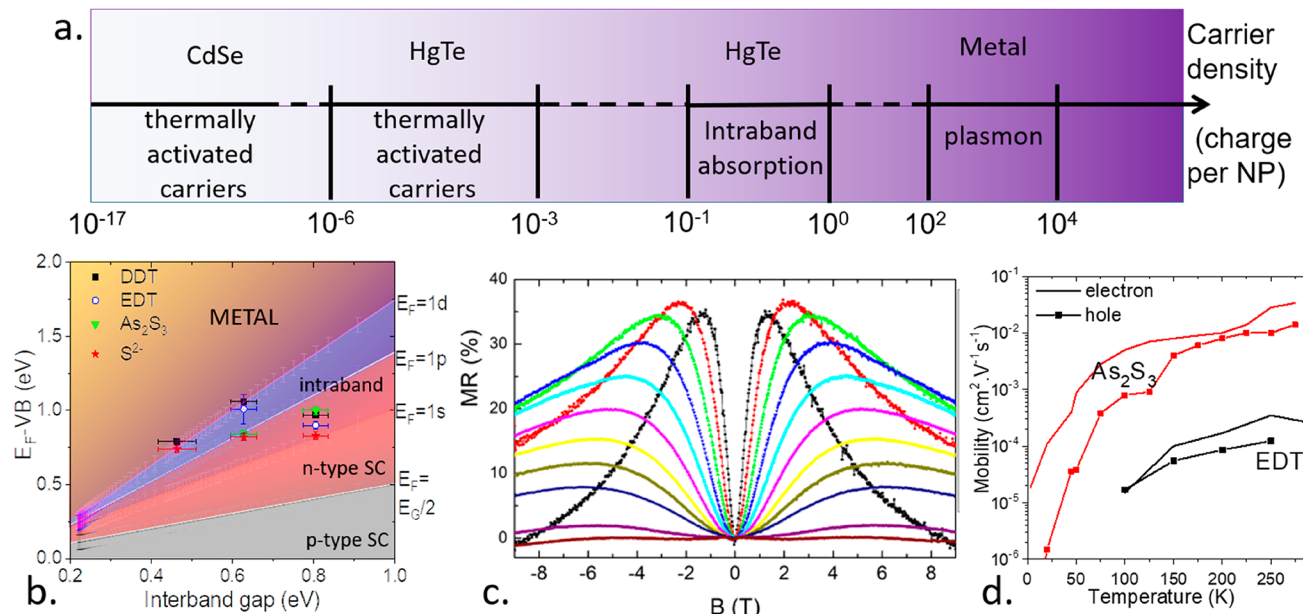


Figure 29. (a) Scale illustrating doping in various NCs. (b) Fermi level energy with respect to the valence band as a function of the interband gap for HgSe NCs. The solid lines correspond to the midinterband gap and the $1S_e$, $1P_e$, and $1D_e$ states from the three-band k-p simulation. (b) Adapted with permission from ref 122. Copyright 2017 American Chemical Society. (c) Magnetoresistance as a function of the applied magnetic field for heavily Mn-doped HgS (15%) NCs. (c) Adapted with permission from ref 206. Copyright 2015 American Chemical Society. (d) Electron and hole mobilities of a HgTe CQD film processed with As_2S_3 and EDT as a function of temperature. (d) Reproduced with permission from ref 35. Copyright 2013 Wiley VCH.

can lead to high doping levels of up to several carriers per NC. In a simple hydrodynamic model, the current density scales as the product of the carrier density and film mobility. For a given surface chemistry (*i.e.*, for a given mobility), this aspect can lead to a significantly high dark conductance for a film of HgTe with approximately 0.01 carriers per NC, especially compared to a CdSe film that has virtually no thermally activated carriers. This increased conductance provides the opportunity to examine the transport in an FET configuration, although this increase impedes IR detection. The large dark carrier density reduces the photocurrent-induced modulation and requires cryogenic conditions to achieve optimal performance. Martinez *et al.* exploited the doped nature of HgSe NCs to investigate the effect of confinement on the metal–insulator transition.¹²² Doping is an intuitive key to examine the metal–insulator transition.^{309,310} According to the Mott criterion, metallic behavior should occur if $\alpha_0 n^{1/3} > 0.25$,³¹¹ where n is the carrier density and $a_0 = \frac{\hbar^2 \epsilon_0 \epsilon_r}{\pi m^* e^2}$ is the Bohr radius, with \hbar being the Planck constant, ϵ_0 being the vacuum permittivity, ϵ_r being the material dielectric constant, m^* being the effective mass, and e being the elementary charge. This approach has been used to induce a metallic nature in doped Si NCs.³⁰⁹ In HgSe, because of self-doping, confinement influences the relative position of the bands with respect to the Fermi level, as shown in the phase diagram in Figure 29b. The authors indicated that by increasing the size (*i.e.*, reducing confinement), the $1S$, $1P$, and finally $1D$ states can be filled and, consequently, the carrier density is increased to 2, 8, and 18 electrons per NC, respectively. The authors, through temperature-dependent conductance measurements, demonstrated that the transport law driving the current in HgSe arrays is affected by the state filling at the NC level. At low carrier densities, the results obtained using Mott's law ($\sigma \propto \exp(-(T_M/T)^{1/4})$, where σ is the conductance and T is the temperature, match well with the experimental results. This

aspect reflects that the density of states drives the current magnitude at low carrier densities. At high carrier densities, typically when the HgSe 1D state is filled, the number of electrons increases and a more metallic behavior is expected. The conductance temperature dependence in this range can be defined using Efros–Shklovskii's law ($\sigma \propto \exp(-(T_{ES}/T)^{1/2})$). This law is the conventional law observed in an array of metallic nanoparticles.³¹² When the D states are filled, the density of the states increases and the availability of the empty states no longer drives the current magnitude. At this point, the current is determined by the Coulombic cost of the nanoparticle charging. This aspect shows that at the nanoparticle level, the NC switches from a semiconductor behavior to a metallic behavior.¹²² However, owing to the weak coupling between the individual nanoparticles at the film level, the collective behavior (*i.e.*, the film behavior) is similar to that of a semiconductor, with an overall increase in the electrical resistance upon cooling.

5.1. Surface Chemistries for Transport in the HgX Array

In NC arrays, transport is driven by the surface chemistry, which determines the carrier mobility.^{24,313} However, in early studies involving transport in arrays of HgX NCs ligand exchange procedures were not as developed as they are today, and sintering was the classic strategy to enhance the transport properties. The latter processes can be implemented through the aggregation of particles during synthesis³ or induced by mild temperature annealing (150 °C).^{194,314} Annealing at temperatures above the growth temperature leads to a strong particle sintering, which favors the nanoparticle coupling associated with a higher carrier mobility. However, this strategy is not viable to design IR sensors because aggregation considerably broadens the optical features of the NCs, leading to an inferiorly defined cutoff wavelength and delocalized oscillator strength. The material obtained from this approach presents p-type conduction with hole mobility of the order of $1 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. This p-

type nature may be the result of material oxidation occurring during annealing or may result from the expected p-type nature of small HgTe NCs even without annealing.^{314,315} Moreover, a low film quality was achieved from aggregated NCs.³ Although electronic coupling within a grain can be large, it remains weak among aggregates. Moreover, the surface of the film obtained from the aggregated NCs is extremely rough, which explains why all the early reports related to HgX for IR detection were based on planar geometry devices for which high quality films are not required.

To achieve milder conditions to obtain conductive films of HgX NCs, solid-state ligand exchange with various short organic ligands was examined. The library of reported ligands includes short amines²⁶⁶ and thiols²⁷⁸ due to the strong affinity of mercury atoms for sulfide ions. During this procedure, the film is exposed to a dilute solution of short ligands in a non-solvent, commonly an alcohol. The concentration of ligands used during this procedure differs from group to group but is typically approximately 1% in mass. This concentration is critical to obtain sufficient ligand exchange: when the value is extremely low, the exchange is not complete. When the ligand concentration is extremely high, the film tends to peel off from the substrate. As ligand exchange occurs, the volume of the particles is reduced. For example, on a 5 nm particle initially capped with C₁₂ chains (\approx 1.5 nm) exchanged for ethanedithiol (EDT), the volume reduction reaches a factor of 2.4. This phenomenon leads to a strong volume contraction of the film, which induces crack formation at the mesoscopic scale. Consequently, multilayer deposition must be implemented for the following two reasons: (i) Because of diffusion, ligand exchange only occurs on a thin slab (*i.e.*, ligand exchange on thick films is not complete) and (ii) the cracks formed during the previous steps must be filled. When implementing solid-state ligand exchange, the band edge feature is more preserved than that in the case of particle sintering; however, the carrier mobility is considerably reduced, ranging from 10^{-4} to 10^{-3} cm² V⁻¹s⁻¹. However, this approach is limited in terms of the achievable film thicknesses, as the overall film quality degrades when too many layers are deposited. The procedure was later developed by Guyot-Sionnest and co-workers, who reported EDT ligand exchange under acidic conditions.^{301,303} The mixture used for ligand exchange was composed of EDT and HCl diluted in ethanol. The acid protonates the long thiolate grafted at the surface, which eases its removal from the NC surface, while chloride ions likely contribute to electronic surface passivation. More recently, the combination of thiol and chloride was further updated for a combination of HgCl₂ and mercaptoethanol. Using this approach, the carrier mobility can be expanded to more than 1 cm²V⁻¹s⁻¹ for both HgTe^{36,316} and HgSe.³¹⁷ This increased carrier mobility was also observed by other groups,^{146,318} although values above 1 cm²V⁻¹ s⁻¹ were not achieved. The discrepancy could likely be attributed to the different shapes of HgTe nanoparticles used by the different teams (spheres *vs* branched particles), which may impact the particle packing and thus the interparticle coupling. This approach is compatible with liquid phase ligand exchange, allowing the formation of concentrated solutions of HgTe nanoparticles. During this procedure, the NCs initially dispersed in a nonpolar solvent are mixed with a polar phase containing short ionic ligands.²⁴ Native long ligands (typically, DDTs) are replaced by short ligands. Owing to this ionic surface, the NCs become soluble in the polar phase. These inks (NCs capped with short ligands compatible with transport and dispersed in a polar

solvent) can be used for the deposition of thick (>500 nm) layers, which considerably increases the absorption of the film compared with that in the previous generation of ligand exchange procedures. In addition, this dual ligand method leads to increased operation stability in air, with a weak increase in the dark current and monotonic *I*–*T* curves.

For infrared nanocrystals, energy transfer from the exciton to ligand absorption is often seen as a key limitation. Consequently, fully inorganic ligands have been investigated. The As₂S₃ material has attracted the most interest, despite its toxicity. Initially, the As₂S₃-based surface chemistry was introduced by Talapin and co-workers as an inorganic strategy to obtain bright near-infrared PbS NCs.³¹⁹ This strategy was later used to build PbS NC arrays with strong photoconduction.³²⁰ The method can also be applied for HgX NCs: in this case, As₂S₃ powder is first dissolved in a short amine, such as propylamine or butylamine. If a HgX NC film is directly exposed to this solution, then the film dissolves, and thus, this ligand solution must be diluted in ethanol before conducting a typical ligand exchange. This method has been applied to HgTe^{35,42,45} and HgSe.^{29,122} Compared to conventional ligand exchange with short organic thiols, As₂S₃-based ligands lead to an increased mobility, as shown in Figure 29d, typically by 2 orders of magnitude. The enhanced stability leads to monotonic *I*–*T* curves, even for air processing.

Furthermore, atomic layer deposition (ALD) has been explored to realize inorganic infilling among NCs^{321,322} and develop conducting films.^{323,324} Alumina³¹⁵ and CdS³²⁵ deposition are performed on HgTe NC films. A clear reduction in film resistance is observed compared to that before ALD. However, this method suffers from two key limitations. First, the approach is generally conducted at a high temperature (*i.e.*, at approximately or above the NC growth temperature), which leads to sintering, and thus, the drop in resistance is presumably a convolution of the mobility increase and bandgap reduction. Second, the photoresponse is reduced after ALD deposition.³²⁵ More work is required to clarify the potential of ALD as a strategy to passivate HgX NC films.

5.2. Magnetotransport

Transport measurement is relatively straightforward in HgX NC arrays, and the potential of these materials for spintronics has also been investigated. Liu *et al.*²⁶⁶ compared the magnetotransport in n- and p-type doped HgTe NCs. The field-resolved conductance presented first, a positive magnetoresistance (MR) at a low field ($B < 0.2$ T), resulting from the spin blockade. When a larger magnetic field was applied, the wave function was compressed, leading to a negative MR signal. This second effect strongly depends on the carrier mass. In HgTe, the hole band is barely dispersed (*i.e.*, the hole mass is similar to the free electron mass), and the wave function compression is limited. However, the conduction band has a low mass, owing to which the MR signal is considerably larger than the hole band, reaching an MR of -10% under 4 T. To enhance the MR signal, the same team proposed to dope the NCs (HgS) with manganese.²⁰⁶ For doping levels below 1%, the authors observed a behavior similar to that previously reported for pristine HgTe CQD films; see the bottom lines of Figure 29c. At higher doping, the authors observe a nonmonotonic magnetoresistance for HgS CQDs. At a low field, a strong positive magnetoresistance of up to 35% is observed, 10 times higher than the spin blockade induced positive MR signal observed in undoped NCs. This strong positive MR signal can be attributed to an electron magnetic

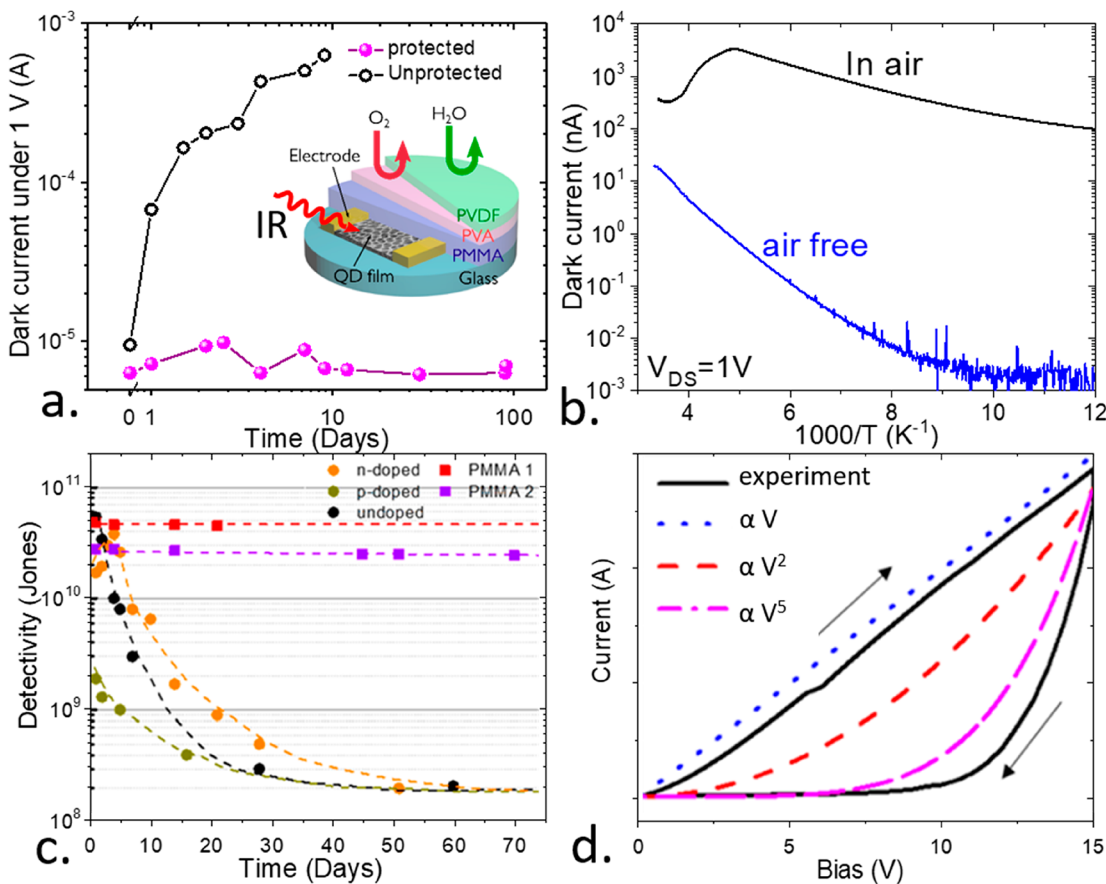


Figure 30. (a) Dark current as a function of time spent in air for a HgTe CQD film with and without protective layers. Time zero corresponds to the sample in the air free glovebox. Inset: scheme of the HgTe CQD film device with polymer protective layers. (a) Adapted with permission from ref 302. Copyright 2018 American Chemical Society. (b) Current as a function of temperature for a film of HgTe CQDs processed with ethanedithiol under air (black) and air-free (blue) conditions. (b) Reproduced with permission from ref 35. Copyright 2013 Wiley VCH. (c) Detectivity of photoconductive devices using a HgTe/hybrid with and without encapsulation. (c) Adapted with permission from ref 36. Copyright 2019 American Chemical Society. (d) I – V hysteresis under illumination (2.5 mW) at 300 K; the forward sweep exhibits linear and high carrier density, the backward sweep corresponds to diffusion. (d) Adapted with permission from ref 327. Copyright 2018 American Chemical Society.

polaron. At higher fields, the polaron binding energy is reduced, leading to a reduced hopping activation energy and a decrease in resistance. Nevertheless, the observed effect remains small. In comparison, it can be considered that the Fe/Cr layers³²⁶ lead to a modulation of more than 50% under a similar magnetic field, and tunnel magnetoresistance can lead to a similar resistance modulation with a 100 times lower field. In this sense, the magnetoresistance in an array of HgX NCs is a promising technique to examine wave function localization.

Jeong *et al.*¹¹⁸ studied the transition from diamagnetic to superparamagnetic characteristics in self-doped HgSe NCs as the carrier density increased and filled the 1Se state. The authors estimated the g-factor to be slightly above 2, and the EPR measurement in a solid-state sample yielded a value of 2.09.

5.3. Stability and Air Operation

A key concern for detector integration relates to the stability of the transport properties, especially when air operation is targeted. As stated previously, transport represents a sensitive approach to identify material modifications. For example, transport properties are usually considerably affected by the formation of an oxide shell or grafting of molecules on the NC surface.

The solid-state ligand exchange conducted on the HgTe NC array enhances not only the material conductivity but also the air

sensitivity. The latter phenomenon manifests as two distinct effects. First, a continuous rise in dark conductance is observed over time;³⁰² see Figure 30a. This increase in dark conductance can reach a factor of 100 after 10 days. The second effect is the nonmonotonic behavior of the temperature dependence of the dark current;³⁵ see Figure 30b. These two effects are particularly problematic for detection applications. The slow rise of the dark current requires regular updates of the correction table used in cameras, and the signal-to-noise ratio of the detector is degraded. Owing to the second effect, cooling, which is the most common strategy to enhance the performance of IR sensors, is rendered ineffective. The origin of this increased dark current remains unexplained. A simple strategy to avoid this dark current increase involves air-free processing of the sample (see Figure 30b). Alternatively, Cryer *et al.* exploited material oxidation.³²⁷ The authors observed that after a certain time, the I – V curves strongly deviated from ohmic behavior and exhibited hysteretic shapes (see Figure 30d). The authors considered this nonlinear I – V curve to design a memistor. To enable air operation while maintaining the initial transport properties, film encapsulation has been considered. However, this aspect remains one of the main challenges to be addressed by HgX NCs in the coming years. Most encapsulation methods, such as those developed for organic electronics, require high temperature annealing, which leads to detrimental sintering in

Table 5. Figures of Merit for Field Effect Transistors Based on HgX Nanocrystal Arrays

HgX	surface chemistry	on/off ratio	temperature (K)	μ_e/μ_h ($\text{cm}^2 \cdot (\text{V} \cdot \text{s})^{-1}$)	gating method	comments	year/ref
HgTe	sintering at 150 °C	10 ³	RT	−/≈2	SiO ₂ /Al ₂ O ₃	dual gate	2006 ³¹⁴
HgTe	likely OH [−] + Al ₂ O ₃ by ALD	10 ³	RT	−/1	Al ₂ O ₃	flexible	2008 ³¹⁵
HgTe	MPOH/HgCl ₂	10 ³	220	4 × 10 ^{−2} /1.5 × 10 ^{−2}	LaF ₃	ionic glass	2019 ³¹⁸
HgTe	MPOH/HgCl ₂	10 ³	80	4.5/0.3	SiO ₂	high mobility	2019 ³⁶
HgS	SCN	200	RT	1.5/NR	SiO ₂	annealed	2017 ¹⁹⁴
HgTe	1-thioglycerol	20	RT	NR/10 ^{−3}	SiO ₂	spray coating	2017 ³³⁰
HgTe	1,7-heptanedithiol	200	RT	0.1/0.1	electrochemical		2012 ²⁶⁶
HgTe	MPOH/HgCl ₂	3 × 10 ³	75	8/1.5	SiO ₂	high mobility	2020 ³¹⁶
HgTe	As ₂ S ₃	<10 @RT 10 ⁷ at 45 K		3 × 10 ^{−2} /1 × 10 ^{−2}	SiO ₂		2013 ³⁵
HgSe	As ₂ S ₃	100	RT	90/NR	SiO ₂ /electrolyte	intraband	2016 ²⁹
HgSe	MPOH/HgCl ₂	100	80	0.7/0.3	SiO ₂		2020 ³¹⁷

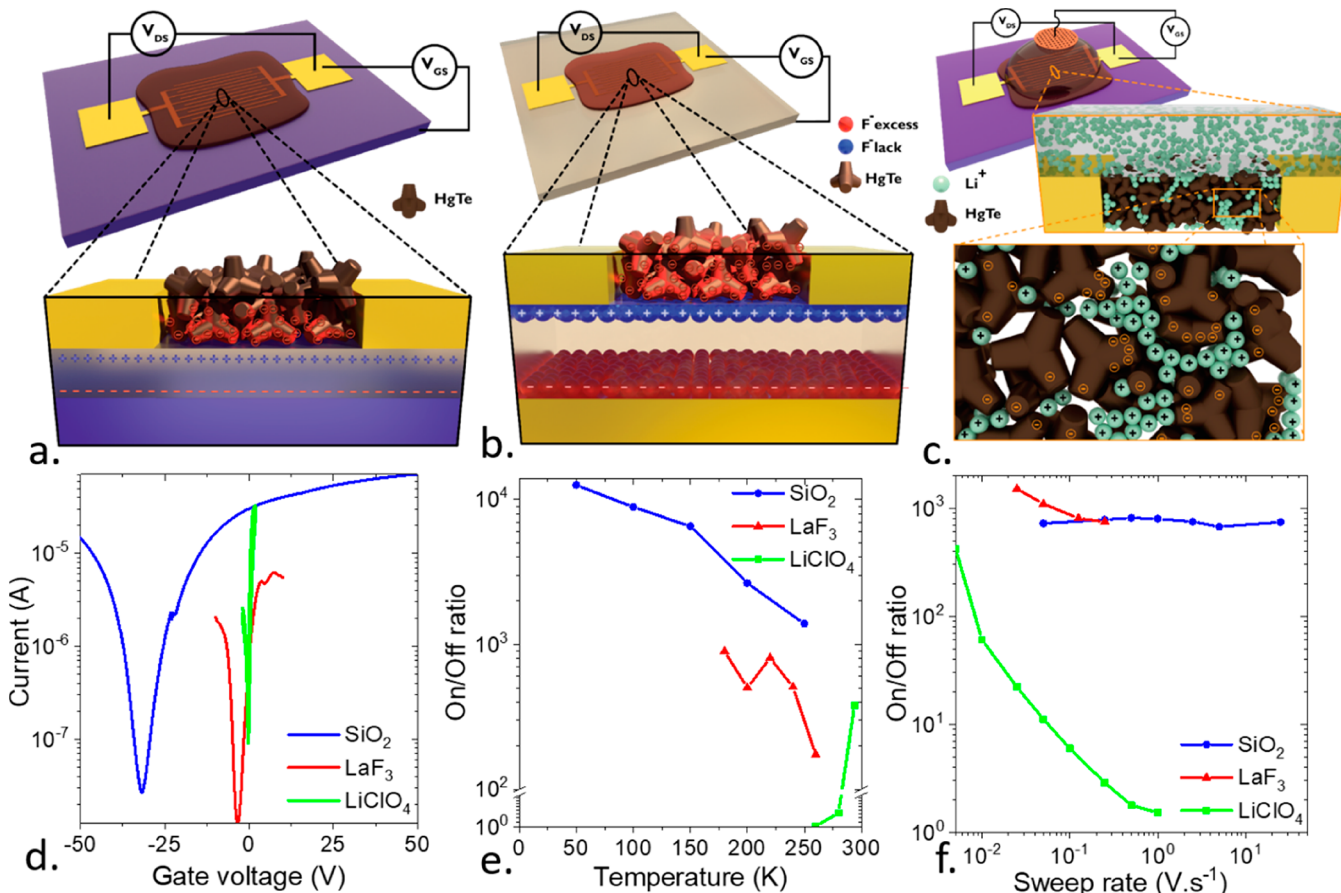


Figure 31. Schemes of HgTe back gate FETs on (a) SiO₂ and (b) LaF₃ when $V_{\text{gs}} > 0$. (c) Scheme of a HgTe top gate FET with an electrolyte (LiClO₄ in a PEG matrix). (d) Typical transfer curves obtained for HgTe nanocrystals with different gating techniques. SiO₂ and LaF₃ are measured at 200 K, and the electrolyte is measured at 300 K. (e) On/off ratios of FET curves measured at different temperatures for different gating configurations. The ratios are normalized according to the ratio at the highest temperature for each gating configuration. (f) On/off ratios normalized according to the ratio at the lowest sweep rate for each gating configuration measured at different sweep rates of V_{GS} for each gating configuration. Adapted with permission from ref 318. Copyright 2019 American Chemical Society.

HgX arrays. Jagtap *et al.* showed that any annealing step above 60 °C leads to a clear red-shift of the absorption spectrum and increase in dark conductance.³⁰² In addition, the growth of core–shell heterostructures is not a viable alternative to enhance the temperature stability. To overcome these difficulties, the authors used a stack of polymer layers deposited at room temperature (see the inset in Figure 30a). The authors

combined water repelling layers (PMMA, PVDF) with an oxygen repelling layer (PVA). The thickness of the stack was approximately 1 μm, sufficiently large to avoid water and oxygen diffusion but sufficiently small to avoid excessive shortwave infrared absorption. Using this approach, air operation with a quasiconstant dark current was obtained over a three-month period. Later, Talapin and co-workers demonstrated that all

these layers were not required and that stable detection properties could be realized using only the PMMA capping layer (see Figure 30c), which simplifies the encapsulation process.³⁶

In addition to device encapsulation, significant progress has been made in building more stable films owing to the enhanced film passivation. The use of As_2S_3 -based surface chemistry was initially motivated by the identification of a capping ligand enabling air processing of the film without observing non-monotonic $I-T$ curves.³⁵ As_2S_3 leads to improved air stability (*i.e.*, monotonic $I-T$ curve even when using air processing); however, the optimal condition is obtained under air free conditions. More recently, the introduction of the dual ligand strategy (thiol + chloride) has facilitated the limitation of postfabrication air oxidation.¹⁴⁶

5.4. HgX Field Effect Transistors

Transport is one of the most sensitive measurements for material investigation: while techniques such as EDX, photoemission, or ICP-MS have resolutions of 0.1% at best, transport is highly sensitive to even small variations (ppm) in composition. In solid-state physics, several experimental reports have been presented on transport to estimate the carrier density and mobility. The Hall effect is the most common strategy for evaluating these quantities in bulk materials. In an NC array, Hall measurements are historically not as popular because of the low mobility of the films. Only recently, with the emergence of high mobility films ($\mu > 1 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$), have Hall measurements with reasonable signal-to-noise ratios been made possible.^{25,328} In 2020, Hall measurements were reported for HgTe ³¹⁶ and HgSe .³¹⁷ The authors used this characterization to extract the film mobilities and confirm the temperature dependence of the field effect mobility.

To date, field effect transistors (FETs) remain the most commonly used strategy to examine transport in HgX NC arrays while extracting information regarding the nature of the carriers. The large inherent doping of HgX NCs affects the FET properties more than that for other NC materials (CdSe , $\text{PbS}(\text{e})$). For wide bandgap materials, the off state is usually limited by FET leakage. This aspect is generally not the case for HgX NCs, for which the off current is limited by thermally activated carriers.^{35,316} This aspect reduces the current modulation in HgX NC-based FETs, typically below 10^3 at room temperature, and the value is even lower for self-doped NCs.

Different gating strategies have been used to design HgX NC-based FETs (see Table 5). Si/SiO₂ wafers, in which the silica layer is used as a dielectric, represent the easiest strategy to design a back gate FET from a film of HgS ,¹⁹⁴ HgSe ,^{29,317} and HgTe ^{35,36,314} NCs due to their commercial availability. As a bias is applied across this layer, surface charges are generated. These charges are screened by the charges injected by the electrodes into the NCs. Dry oxide is beneficial to limit the breakdown and allows for a high gate bias operation.^{36,316} Silica gating can lead to large current modulations (10^7 at 50 K has been reported³⁵) but requires a high gate bias operation (50 V and more) due to its low dielectric constant (3.9). Consequently, insulators with a higher dielectric constant, such as alumina, have been used for the design of HgX NC-based FETs,^{314,315} reducing the gate bias range to below 10 V for the same current modulation. Using this approach, Yun *et al.* combined p-type HgTe NCs with n-type HgSe NCs to design NOR and NAND logic gates.³²⁹

Alternatives to dielectrics have also been explored to realize the gating of HgX NC arrays. The quest for such alternatives was first motivated by the design of higher capacitance gates to control the carrier density in self-doped NCs. Electrolytes can be a strategy to induce gate effect in wide bandgap NCs, such as CdSe ^{331,332} or ZnO .³³³ Electrolyte gating of HgTe ^{30,96,262,266,278} and HgSe ^{29,120,122} NCs has also been well developed. An electrolyte is a dispersion of ions within a liquid or a polymer matrix,¹⁵⁸ see Figure 31c. As a gate bias is applied, the ions can percolate within the NC array, and the electrodes inject carriers to screen these charges. Moreover, intercalation of the ions within the NC atomic lattice has been reported.³³⁴ In the electrolyte, gate capacitance is driven by the ratio of the dielectric constants on each side of the ionic double layer formed at the interface of the NCs. This double layer is extremely thin (1 nm), and the gate capacitance is consequently much larger ($\approx 1 \mu\text{F}\cdot\text{cm}^{-2}$) than that for dielectric materials ($\approx 10 \text{ nF}\cdot\text{cm}^{-2}$ for 400 nm SiO₂). Because of the percolation of ions within the NC array, it is possible to obtain a gate effect in thick NC films, which is of key interest to design phototransistors in which both large absorption and gate tunability are required. The large gate capacitance enables the realization of a low bias operation (<3 V, as shown in Figure 31d). Nevertheless, the gate effect is based on the displacement of ions; this process is intrinsically slow and thus limits the sweep rate (Figure 31f). It also requires an operation of the device above the freezing point of the matrix (Figure 31e). The last point allows the preparation of a device in a given charging configuration at room temperature before cooling it below the electrolyte freezing point; however, this approach cannot be used to realize the low temperature tuning of the carrier density.

To overcome this difficulty, Gréboval *et al.* proposed the use of ionic glass³³⁵ to gate HgTe NC films.^{108,318} The authors used a LaF₃ substrate, which is also an IR transparent material. Such glass types include fluorine vacancies (F⁻ vacancies) that can move to the surface to induce polarization under an applied bias,³³⁶ see Figure 31b. Ionic glass also has a large capacitance, allowing low bias operation (see Figure 31d). Vacancy displacement is more efficient than ion diffusion in the electrolyte matrix, which enables a higher sweep rate and operation down to 150 K. The latter temperature of the glass is determined by the vacancy freezing point (Figure 31f). In this sense, ionic glass gating combines the advantages of electrolytes with solid-state handling,^{337,338} which releases the constraint from manipulating liquids/polymers.

More recently, the same group explored a complementary approach for the gating of HgX NC films. High capacitance methods (electrolytes or ionic glasses) rely on ion displacement, which corresponds to two intrinsic limitations: (i) slow operation limiting the sweep rate and (ii) a relatively high operating temperature (above the freezing point, which is always above 200 K). The latter point is detrimental for the design of phototransistors dedicated to IR detection because cryogenic conditions are required, especially for the detection of long IR wavelengths. For example, the operating temperature of a 5 μm cutoff wavelength camera based on epitaxially grown semiconductors remains below 150 K, dropping by approximately 90 K for 10 μm detection and as low as 4 K in the THz range. Consequently, a gating method to accommodate low temperature operations while still offering a large capacitance and possibly without the limitation of SiO₂ (breakdown) is of considerable interest. Gréboval *et al.* explored a ferroelectric gating strategy.³³⁹ At the Curie temperature of a ferroelectric

crystal (T_c), the material transitions from a paraelectric to a ferroelectric phase, which induces a divergence in the dielectric constant, as depicted in Figure 32a. To obtain low temperature

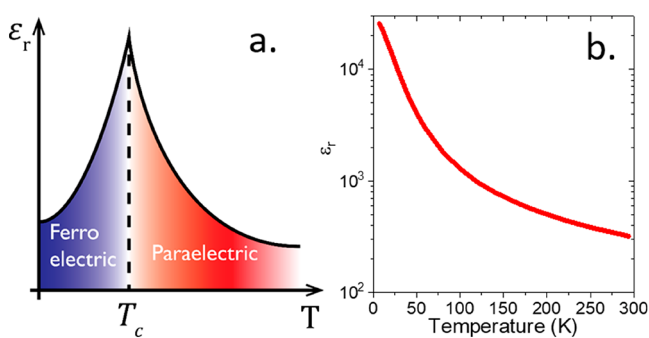


Figure 32. (a) Scheme of the divergence of the dielectric constant around the Curie temperature as a paraelectric switch toward its ferroelectric phase. (b) Dielectric constant dependence on the temperature of the STO substrate used as the gate. (b) Adapted with permission from ref 339. Copyright 2021 American Chemical Society.

gating, the authors used SrTiO₃, a frustrated ferroelectric (*i.e.*, with a negative T_c), also known as a quantum paraelectric. The dielectric constant strongly rises below 100 K and can reach 30 000 below 40 K. At this temperature, the dielectric constant

saturates owing to quantum fluctuation.³⁴⁰ Such gating has been explored previously for a 2D superconducting electron gas obtained at the LaAlO₃/SrTiO₃ interface.^{341,342} Gréboval *et al.* applied this gating to HgTe NCs with various bandgaps from 720 to 250 meV and obtained a large current modulation ($>10^3$)³³⁹ owing to the use of the SrTiO₃ gate. The reported capacitance is equivalent to that of an 80 nm thick SiO₂ film. Compared with thin silica films, quantum paraelectric gating is a breakdown free/low leakage gating method. A bias of as large as 200 V can be applied without any risk of device destruction (*i.e.*, without electrical breakdown of the gate). This method is especially promising owing to the recent progress in thin film deposition of oxide and ferroelectric materials.

While it is unlikely that HgX NC-based FETs can compete with Si-based technologies for electronics, FET appears to be a promising spectroscopic tool to investigate the confined forms of HgX. In addition to the determination of the majority carriers and internanoparticle coupling through mobility, observation of a Pauli blockade can be used to determine and finely tune the state filling of NC quantum states. The concept was initially introduced for CdSe NCs, while Guyot-Sionnest and co-workers observed nonmonotonic mobility behavior as the 1S_e state is filled using electrolytic gating.³⁴³ The 1S_e state presents a 2-fold degeneracy; thus, when the Fermi level is brought in resonance with the 1S_e state, the mobility is maximized for a state filling of 1 carrier per NC. For higher fillings, the film conductance drops

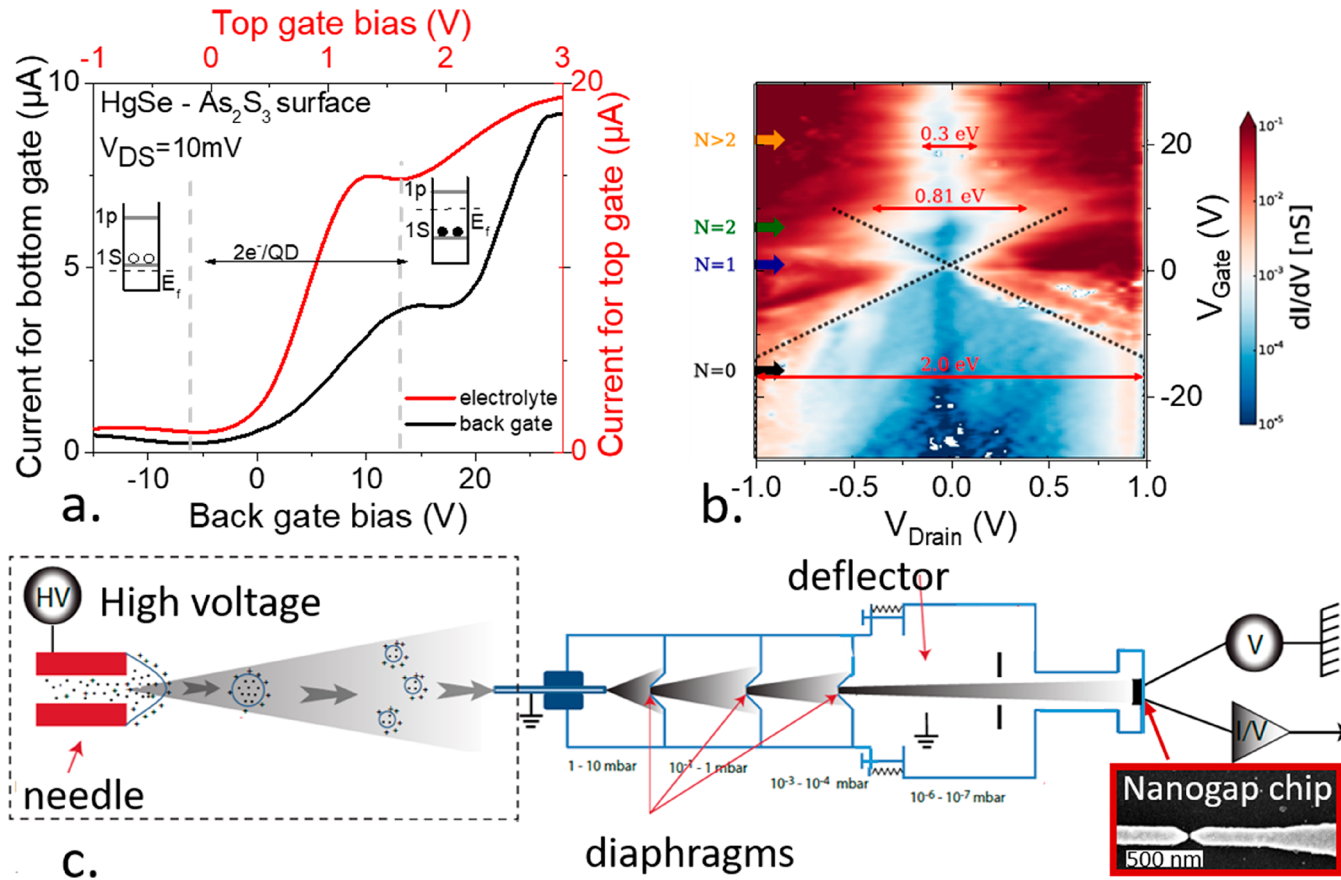


Figure 33. (a) Transfer curves (drain current vs gate voltage) for a thin HgSe CQD film using back and electrolytic gating. In both cases, the drain source bias is 10 mV. (a) Adapted with permission from ref 29. Copyright 2016 American Chemical Society. (b) Log scale color map of the differential conductance dI/dV as a function of the drain and gate voltages for a single HgSe NC. The black dashed lines are a visual guide for the closing of the gap as the 1S_e level crosses the Fermi level energy. (c) Sketch of the electrospray setup utilized to fabricate the single QD junctions. (b,c) Adapted with permission from ref 259. Copyright 2017 American Chemical Society.

until the $1P_e$ state starts to be filled. This drop in mobility is a result of the fact that in the regime between 1 and 2 carriers per NC on average, certain NCs exhibit a fully filled (2 carriers) $1S_e$ state. Consequently, these NCs cannot contribute to transport, as the only way for carriers to move is to pass through the $1P_e$ state, which requires overcoming the $1P_e$ – $1S_e$ energy barrier. Using a dual gate configuration (bottom SiO_2 gate + top electrolytic gate), Lhuillier *et al.* demonstrated that a similar behavior can be observed in an array of monodisperse HgSe NCs,²⁹ as shown in Figure 33a. This observation was later confirmed by Chen *et al.* by using only a bottom dielectric gate, discussed as state resolved transport.³¹⁷ The observation of a Pauli blockade in HgTe occurred only later. This delay is related to the fact that the observation of a Pauli blockade requires more monodisperse syntheses, which, until recently, was not the case for HgTe NCs. The introduction of new synthetic procedures by Shen *et al.*⁶³ enabled the growth of spherical and monodisperse HgTe, leading to the observation of these state-resolved transfer curves.³¹⁶

Aubin and co-workers further investigated state-resolved transport in HgSe NCs²⁵⁹ by conducting single particle measurements. The strategy for a single NC-based experiment usually relies on the measurement of a single NC PL signal. However, single particle measurements become especially difficult when using infrared materials due to their low PL efficiency and the lack of efficient single photodetectors in the infrared region. The strategy proposed by Aubin and co-workers is based on on-chip tunnel spectroscopy. Compared to conventional scanning tunneling spectroscopy, this approach offers higher temporal stability and enables gating. The above-mentioned method is based on two electrodes spaced by 5–10 nm (*i.e.*, nanogap), prepared by e-beam lithography. A solution of nanoparticles is sprayed under vacuum on a chip including 32 nanogaps; see the scheme of the setup in Figure 33c. After each spray, the conductance through the nanogap electrodes is measured, and a sudden increase in conductance indicates particle trapping. The method has been applied to a variety of nanoparticles, including gold,³⁴⁴ Fe_3O_4 ,³⁴⁵ PbS,³⁴⁶ and HgSe²⁵⁹ NCs. Once the particle is trapped, on-chip electron tunnel spectroscopy can be conducted at a low temperature. A typical tunnel spectroscopy map is shown in Figure 33b. As a function of the filling with the gate, two bandgaps are observed. The large gap, under hole injection (*i.e.*, negative gate bias), corresponds to the interband gap, while the smaller gap, under electron injection, corresponds to the intraband gap. The obtained values can be fully correlated with optical measurements from IR spectroscopy. This result indicates that self-doping is present at the single particle level and that intraband absorption is not the consequence of collective behavior. Finally, the authors also reported on photoconduction from a single HgSe NC.

6. PHOTOCONDUCTION AND IR DETECTION

The last section of this review is dedicated to infrared sensing based on HgX NCs. Bulk HgCdTe alloy is the most mature material for infrared detection because of its high performance and wide tunability from the near-infrared region to the long-wave infrared region. In this material, spectral tunability is obtained by alloying and controlling the Cd fraction. In HgTe NCs, the same magnitude of tunability is enabled through confinement control, which renders HgTe NCs a natural replacement for HgCdTe in the future design of IR sensors. However, IR detection applications only appeared 10 years after the introduction of mercury chalcogenide NCs, whose first

application was focused on light emission. This low interest in HgX NC-based IR detection in the early 2000s was likely due to the inferior transport properties of the NCs available at that time. During the 2000–2010 period, substantial progresses in ligand exchange procedures were made, allowing the design of high mobility films. This aspect demonstrated that NCs could be used for IR detector design even though the approaches were not immediately applied to films of HgX NCs.

This section is organized as follows. We first redefine the main figures of merit relative to infrared detection, including noise, because this quantity is often wrongly evaluated. In this discussion, we also mention the experimental setups used to measure these figures of merit. Next, we discuss different strategies for the devices that have been developed. First, we consider photoconductors and later discuss the refinements of the planar geometry, including phototransistors, flexible devices, intraband detection, and coupling to 2D materials. The following section is focused on photodiodes, mostly relying on vertical geometry devices. This section is valuable because all of the highest performing devices are based on this geometry. The final sections of the chapter are dedicated to three key recent developments that enhance the performance and maturity of IR detection based on NCS. These developments include (i) multicolor sensing, (ii) detectors with enhanced light–matter coupling, and (iii) multipixel devices and focal plane arrays.

6.1. Figures of Merit and Their Experimental Determination

Figures of merit are used to compare the performance of different types of photodetectors (*e.g.*, different working principles, channel materials, and geometries). In the following discussion, we briefly introduce the general figures of merit and their experimental measurements.

6.1.1. Spectral Response. The term “infrared” describes a broad part of the optical spectrum at longer wavelengths than those of the visible range. The infrared region is commonly subdivided into three main domains: the near-infrared region (1–3 μm), mid-infrared region (3–20 μm), which corresponds to the range of blackbody emission close to room temperature and in which most chemical bonds present resonances, and far-infrared region, which strongly overlaps with the THz range. In regard to IR sensing and imaging, the subdivisions are even more defined, especially to account for the transparency ranges of the atmosphere and the region in which it is possible to conduct long distance imaging. The following definitions can be considered:

Near Infrared (NIR): NIR is close to the visible spectrum and spans from 700 to 900 nm. This range overlaps with the absorption of silicon. Target applications in the NIR range are divided into biological imaging and fluorescence detection.

Shortwave Infrared (SWIR): SWIR is defined as the wavelength range from 1 to 1.7 μm . The latter value is arbitrary because it corresponds to the bandgap of InGaAs detectors, which are the most mature detectors in this range of wavelengths. In this range, several applications are possible, including active imaging, night glow-assisted imaging, and tissue imaging. It is also common to define SWIR in the range between 1.7 and 2.5 μm . In SWIR, the contrast in an image is, as in the visible region, the result of the scattered light from an external source.

Midwave Infrared (MWIR): MWIR extends from 3 to 5 μm and corresponds to an atmospheric transparency window. Above 3 μm , blackbody emission starts to prevail over reflection from external sources, enabling thermal imaging. In MWIR, hot objects are typically targeted. Moreover, thermal imaging can be

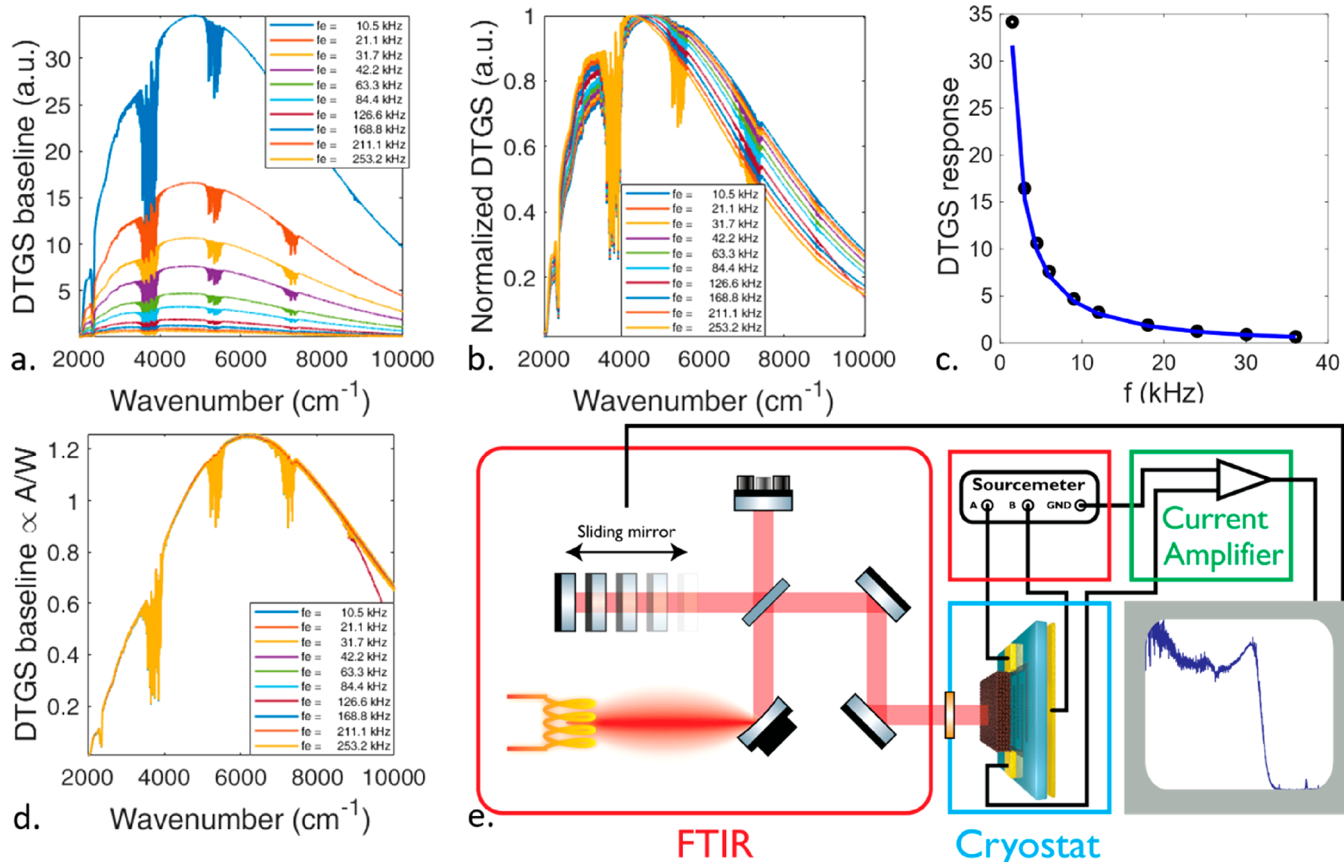


Figure 34. (a) DTGS signal of an FTIR source measured at different sampling frequencies. (b) Normalized DTGS signal at different sampling frequencies. (c) Estimated transfer function at 4500 cm^{-1} and its fit by $H(f) = (a + bf + cf^2 + df^3)^{-1}$. (d) DTGS source spectra at different sampling frequencies corrected by the transfer function. (e) Scheme of a typical setup used to measure the spectral response. FTIR illuminates the sample that is biased by a source meter and can eventually be placed in a cryostat. The current is later amplified by a transimpedance amplifier and sent back to the FTIR electronics to compute the Fourier transform.

conducted at room temperature with a greater contrast than in LWIR, albeit at the cost of a smaller number of photons. In this range, narrow bandgap semiconductors, such as InSb and HgCdTe and other type II superlattices based on InAs/GaSb, are conventional materials used for photodetection and in applications such as thermal imaging and gas sensing.

Longwave Infrared (LWIR): LWIR ranges from 8 to $12\text{ }\mu\text{m}$. This range is well matched to the maximum emission of a blackbody at room temperature ($10\text{ }\mu\text{m}$). This range is thus a useful range to perform passive thermal imaging. Ternary alloy materials, such as HgCdTe and type II superlattices, are the traditional materials used for LWIR detection. This range is also the operating range of bolometers, energy detectors used in most mass market thermal detectors and cameras. Detectors in LWIR are applied in industrial inspection, defense, and astronomy.

Terahertz (THz): Terahertz ranges from 30 to $200\text{ }\mu\text{m}$. Bolometers that operate at low temperature are typically used for THz detection. This range of wavelengths is used in astronomy and to conduct nonionizing surveillance (mostly for safety applications). In the THz range, various materials that are opaque in the visible region become transparent.

The classical techniques for detector characterization typically employ a monochromator, which is an integrating sphere, and rely on measuring the signal on a calibrated detector. In the infrared range, such techniques are difficult to set up. One convenient way to measure the spectral response is to perform

Fourier transform infrared spectroscopy (FTIR) in the continuous scan mode. The spectral response is measured by placing the detector in a FTIR instrument and amplifying the current with a low noise amplifier before sending the signal back to the FTIR electronics to perform a Fourier transform. In this case, the collected signal typically comes from the spectrum of the source multiplied by the transmission of the optics and by the spectral response of the detector. The latter is the signal of interest. A reference is thus required to extract the spectral response of the detector from this product. This background is generally measured by using the signal collected through the internal DTGS detector of the FTIR, which has a flat spectral response. Consequently, the signal measured on the DTGS is simply the product of the source spectrum by the transmission of the optics. Dividing the signal acquired on the unknown detector by that from the DTGS leads to the spectral response of the detector.

However, as explained by Gravrand *et al.*,³⁴⁷ this spectrum should be corrected by the transfer function of the instrument, taking into account the low response time of the DTGS detector; see Figure 34a,b. The measured signal (after normalization by the DTGS background) is the result of the product of this transfer function by the detector's real spectral response. The transfer function can be estimated by measuring the IR source spectra at different sampling frequencies with the DTGS detector. The transfer function can be estimated as $H(f) = \frac{1}{a + bf + cf^2 + df^3}$, where f is the temporal frequency

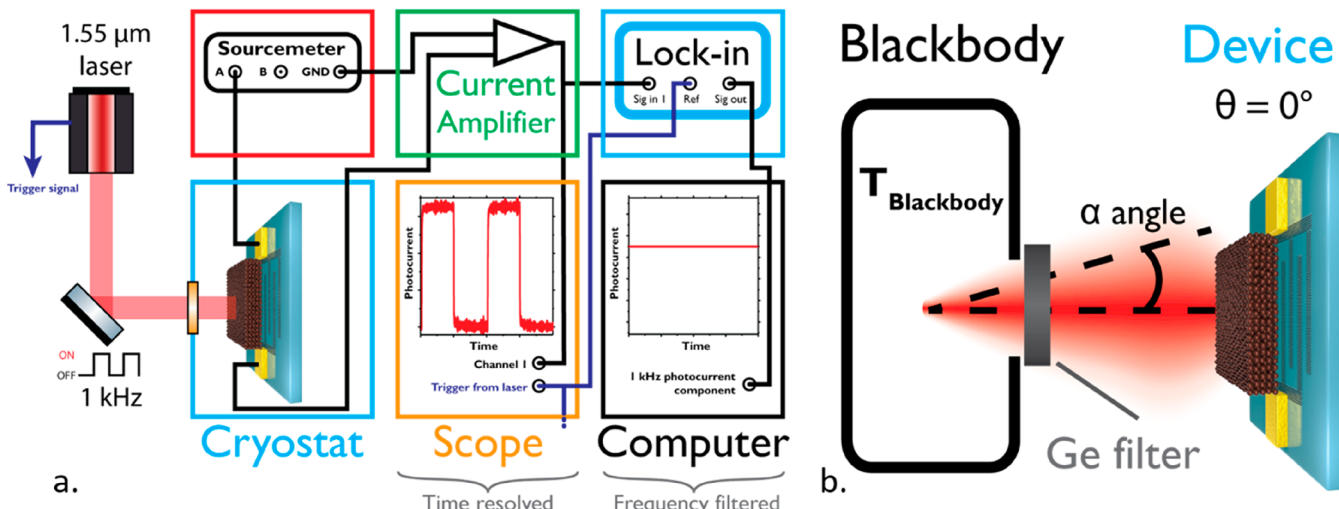


Figure 35. (a) Scheme of an experimental setup to measure the photocurrent with an electrically chopped laser at $1.55\text{ }\mu\text{m}$. The signal is biased with a sourcemeter, and the current is amplified through a transimpedance amplifier (TIA). The photocurrent can either be measured by a scope in a time-resolved setup or, for low SNR samples, a lock-in amplifier can be used to filter all the other frequencies of the signal. (b) Scheme of a response setup with blackbody illumination. The device is at a normal incidence ($\theta = 0^\circ$), α is the half viewing angle of the device by the blackbody, $T_{\text{Blackbody}}$ is the temperature of the blackbody, and the germanium filter is a high pass filter with a $1.8\text{ }\mu\text{m}$ cutoff.

associated with the wavenumber after the Fourier transform; = $\text{RR} \times f_e \times \sigma$, where RR is the retardation rate of the laser, f_e is the sampling frequency, and σ is the wavenumber; see Figure 34c. Once the fitting parameters a , b , c , and d are identified, the source spectrum is divided by the transfer function to ensure that the response of the detector becomes independent of the acquisition speed (see Figure 34d). The obtained spectrum is at this stage specified in relative units and must be calibrated in $\text{A}\cdot\text{W}^{-1}$, which requires precise knowledge of the incident photon flux. Another way of rescaling the spectral response in $\text{A}\cdot\text{W}^{-1}$ is to measure the exact device responsivity at a given wavelength with a known incident flux (e.g., with a laser diode).

6.1.2. Operation Temperature Range. To reduce the dark current in the device and its associated noise, low temperature operation is required to achieve high signal-to-noise ratios in the infrared domain. A larger targeted wavelength corresponds to a lower operating temperature. Cryogenic temperature operation is associated with a high cost, weight, and constraints regarding the lifetime. Consequently, most research on IR detection is focused on increasing the operating temperature. In the SWIR, room temperature operation is possible, but it is common to use a Peltier stage, which keeps the operating temperature as constant as possible. Depending on the targeted performance, cooling can also be implemented in SWIR, generally over a range of temperatures that is compatible with the Peltier stage. In MWIR, cryogenic operation is usually performed. In the blue part of MWIR (i.e., below the CO₂ absorption at $4.2\text{ }\mu\text{m}$), state-of-the-art operating conditions can attain temperatures of up to 150 K, while for the red part of MWIR, the usual operating temperature is approximately 120 K. In LWIR, operating conditions immediately above the temperature of liquid nitrogen can be achieved. In THz, liquid He operating conditions are usually employed. These constraints must be considered to highlight the level of performance required for the NC-based IR sensors. In particular, in SWIR, low cost sensing requires room temperature operation.

6.1.3. Responsivity (R , $\text{A}\cdot\text{W}^{-1}$). Responsivity (R) is the ability of a material to convert incident light into an electrical signal and is a common indicator to evaluate the performance of

photoconductive devices. Responsivity is the ratio of the detector's photocurrent (I_{ph}) to the total incident optical power (P_{in}) that it receives. This value can be simply described as $R = I_{\text{ph}}/P_{\text{in}}$.

In nanocrystal devices, R is strongly dependent on the incident flux. In general, a higher photon flux corresponds to a weaker response. Because of this behavior, the R values reported using a blackbody (low photon flux source) are generally higher than those measured using laser sources. Moreover, it is important to investigate the frequency dependence of R to determine the range of operating frequencies of the device.

Responsivity is usually measured at a given frequency, indicating that the sample is illuminated by electrically or mechanically chopped light. This light can either be a laser or a filtered blackbody. In the case of laser illumination, the power of the laser is directly measured using a calibrated detector. For blackbody radiation, the power can be calculated using the following formula:

$$P(W) = A_d \pi \cdot \cos(\theta) \cdot \sin^2(\alpha) \int_{\lambda_{\min}}^{\lambda_{\max}} \frac{2hc^2}{\lambda^5} \frac{1}{e^{hc/\lambda kT} - 1} d\lambda$$

where A_d is the area of the device, θ is the incidence angle, α is the half angle of the view under which the blackbody is detected by the device, λ_{\min} and λ_{\max} are the border wavelengths of the range of interest (taken as the cutoff wavelength of the filter and measured detector, respectively), h is Planck's constant, c is the speed of light, k is the Boltzmann constant, and T is the blackbody temperature.

Once the power is determined, the photocurrent can be measured either with an oscilloscope or, if the chopping signal is a sine curve, with a lock in amplifier, as shown in Figure 35a,b. The ratio between the photocurrent and incident power yields the responsivity in $\text{A}\cdot\text{W}^{-1}$.

6.1.4. Current Noise Spectral Density (δI_n in $\text{A}\cdot\text{Hz}^{-1/2}$).

The current noise spectral density is highly specific to the infrared region. Because of the narrow bandgaps of the materials used for IR detection, optical excitation is always in competition with thermal excitation. Consequently, the signal-to-noise ratio is weak compared to that for visible and UV detectors. The noise

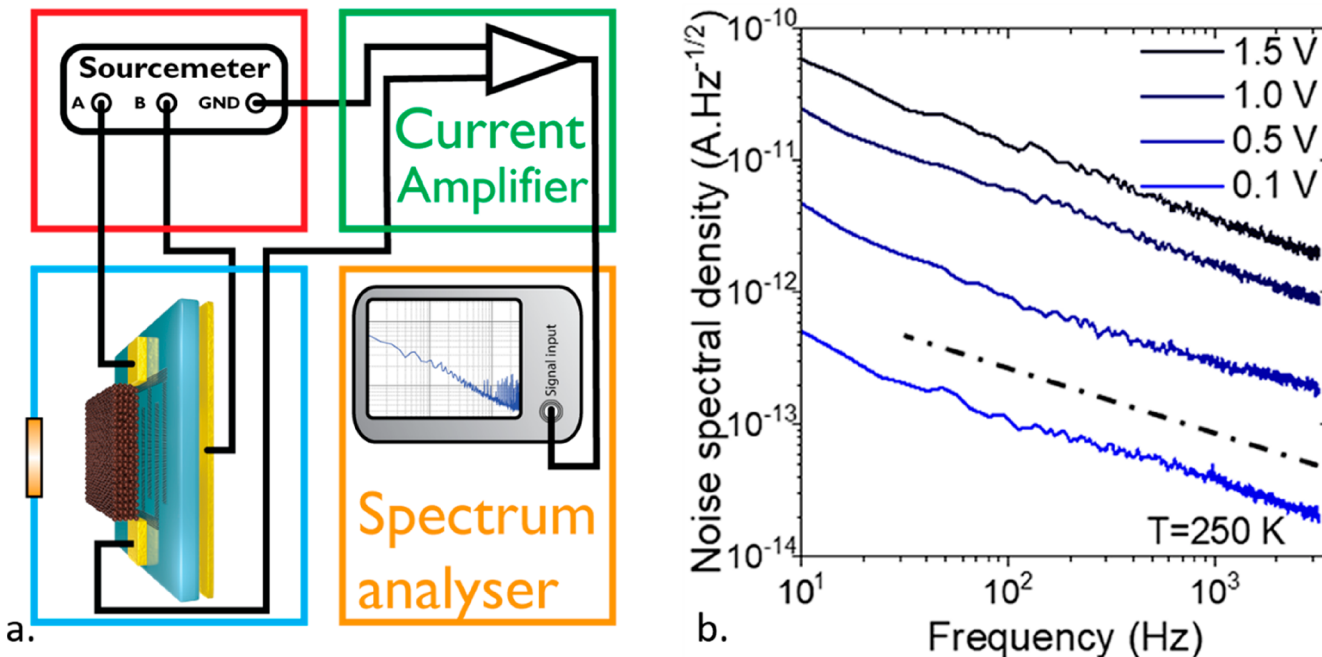


Figure 36. (a) Scheme of a typical experimental setup for making a noise spectral density measurement. The sample is biased by a source meter (or a battery), and the current is amplified through a low noise transimpedance amplifier. The amplified signal is sent to a spectrum analyzer. (b) Typical noise spectral density curves for a HgTe film under various biases at 250 K. The dashed line represents the 1/f slope.

in the photodetector mainly consists of (i) Johnson noise, (ii) shot noise, and (iii) flicker noise ($1/f$). (i) The thermal noise δI_{th} is related to the fluctuation of the carrier density in any object, which is a common behavior that is not specific to conductive materials. The corresponding value can be easily estimated as $\delta I_{\text{th}}^2 = 4 kT \Delta f / \text{Res}$, where kT is the thermal energy, Δf is the frequency bandwidth of the measurement, and Res is the device resistance. This noise is white noise (*i.e.*, independent relationship with frequency). (ii) The shot noise relates to the charge fluctuation resulting from the current flow in the device and can be defined as $\delta I_{\text{GR}}^2 = 2e\bar{g}I$, where e is the elementary charge, I is the current flowing in the structure, and \bar{g} is the photoconduction gain. As a form of thermal noise, this noise is white noise.

In nanocrystals, these two white noise sources rarely dominate. The main contribution originates from (iii) $1/f$ noise; see Figure 36b.^{348,349} Contrary to the previous sources of noise, there is no analytical expression to evaluate $1/f$ noise from easily measurable parameters (R , I). Hooge's law provides an empirical expression³⁵⁰ for $1/f$ noise $\delta I_{1/f}^2 = \frac{\alpha I^2}{Nf}$, where N is the total number of carriers involved in the transport, I is the current, f is the frequency, and α is Hooge's constant over a large range of carrier concentrations. However, for arrays of NCs, Hooge's constant does not appear to be consistent^{279,348} and is larger²⁷⁹ (approximately 1) than the value reported by Hooge in metallic thin films ($\alpha < 10^{-3}$). In NC films, α appears to be independent of the material nature (such as metal, wide bandgap semiconductor, or narrow bandgap semiconductor) and particle size.³⁴⁸ However, the quality of the film and, particularly, the internanoparticle coupling dramatically drive the actual magnitude of α . A larger coupling corresponds to a smaller α . Overall, in NC-based detectors, frequency-resolved measurements must be performed to determine the actual value of noise as it cannot be measured analytically. The usual assumption of using white noise as the limiting noise systematically under-

estimates noise, often by several orders of magnitude, which dramatically overestimates the signal-to-noise ratio (and the detectivity) of the device.

In practice, noise is usually measured by biasing the device with a battery. The current is amplified through a low noise amplifier and sent to a spectrum analyzer; see the scheme of the experimental setup in Figure 36a. The typically targeted range of frequency is $1/t_i$, where t_i is the targeted integration time. Generally, noise is acquired between 1 Hz and a few kHz. A current amplifier whose bandwidth matches the experiment must be used, especially for highly resistive samples that require large gains. The reported noise needs to be sample limited and not amplifier limited. Additionally, for uncooled amplifiers, the lowest measurable noise is of the order of a few $\text{fA}\cdot\text{Hz}^{-1/2}$. This aspect can be a general limitation to measure noise in resistive samples (especially for wide bandgap samples). A key parameter for this measurement is the noise level of the room in which the measurement is performed. The sample and electronics must be isolated from other noise sources in the room. The first source of noise to be suppressed is the 50 Hz (or 60 Hz, depending on the location) noise from the AC line. For highly conductive samples (low resistance), noise measurement may be more difficult to perform, and a Wheatstone bridge can be used to suppress the DC component of the signal,³⁴⁸ allowing the use of a higher amplification gain and resolution of the sample noise spectral density.

6.1.5. Noise Equivalent Power (NEP, $\text{W}/\text{Hz}^{1/2}$). NEP is the minimum optical power for a detector to output a signal-to-noise ratio of 1 with a 1 Hz bandwidth. This value can be expressed as $\text{NEP} = \delta I_n / R$, where δI_n is the current noise spectral density and R is the responsivity. This value is an indicator of the availability to detect the minimum illumination power; a low NEP is desirable for a photodetector.

6.1.6. Specific Detectivity (D^* , $\text{cm Hz}^{1/2} \text{ W}^{-1}$). The specific detectivity is a signal-to-noise ratio derived from the NEP and normalized by the optical area of the device to allow

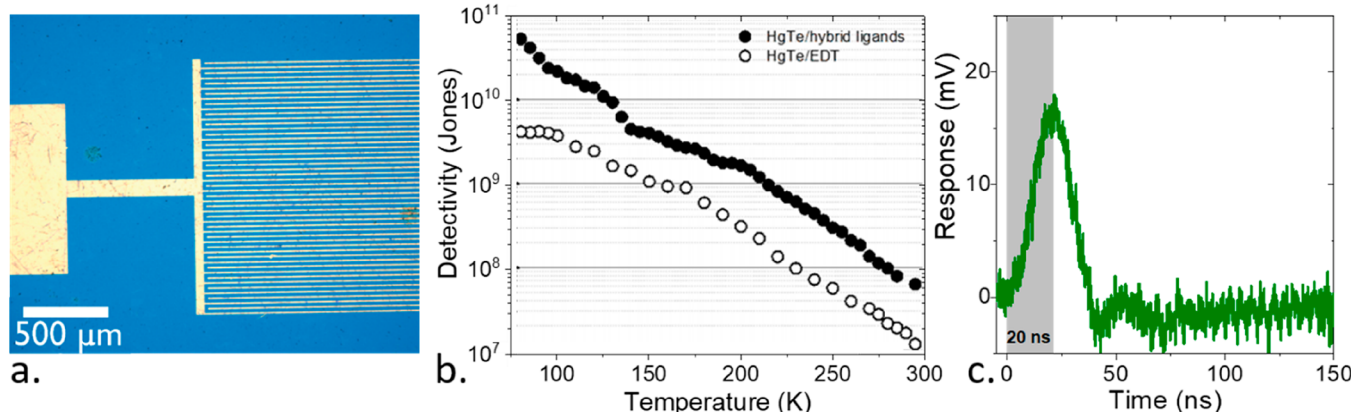


Figure 37. (a) Microscopy image of interdigitated electrodes. (b) Detectivity as a function of temperature for HgTe (HgCl_2 /mercaptoethanol surface chemistry) and HgTe/EDT films. (b) Adapted with permission from ref 36. Copyright 2019 American Chemical Society. (c) Typical response of a 5 μm cutoff wavelength film made of HgTe NCs to a 4.4 μm pulsed quantum cascade laser. (c) Adapted with permission from ref 219. Copyright 2018 American Chemical Society.

fair comparisons from detector to detector. Generally, this ratio is normalized by the optical area of the device to account for the variation from device to device. In this case, we consider the specific detectivity (D^*), which can be expressed as $D^* = RA^{-1/2}/\delta I_n$, where A is the detector optical area, R is the responsivity, and δI_n is the current noise spectral density. The corresponding unit is $\text{cm} \cdot \text{Hz}^{1/2} \cdot \text{W}^{-1}$, also known as Jones. The D^* value depends on an applied bias and signal frequency. Because the main noise source is $1/f$ noise, it may appear attractive to report detectivity at a high frequency. However, the responsivity measurement needs to be conducted at the same frequency. Typically, the detector needs to be characterized at frequencies lower than the 3 dB cutoff of its responsivity. A detectivity value should always be reported with the frequency used for its calculation, as well as the cutoff wavelength of the detector, operating temperature and bias.

6.1.7. Response Time. As opposed to solar cells, for which the response time is not a relevant figure of merit, the response time of an IR sensor is critical. Sensors dedicated to imaging must be compatible with the video frame rate (*i.e.*, exhibit a smaller response time than 1/25 s). Other applications (missile detection, LIDAR detection) require a considerably higher response time (<1 μs).

6.1.8. Noise Equivalent Temperature Difference (NETD, mK). The NETD is a useful parameter to evaluate the performance of focal plane arrays (FPAs) influenced by the signal-to-noise ratio of IR detectors. This value quantizes the minimum temperature difference that can be imaged using an infrared sensor. This quantity is only relevant for thermal sensing (*i.e.*, mostly for MWIR and LWIR applications) and can be expressed as³⁵¹ $\text{NETD} = \delta I_n h\nu P_{\text{in}} / k_B T_B^2$, where δI_n is the current noise spectral density, P_{in} is the incident optical power, h is Planck's constant, ν is the frequency of light, k_B is the Boltzmann constant, and T_B is the blackbody radiation temperature. High performance quantum sensors reach NETDs of as low as 10 mK, while thermal sensors, such as bolometers, have typical NETD values in the 50–100 mK range.

6.2. Photoconductive Devices

6.2.1. Conventional Geometries. HgTe is the most investigated material among mercury chalcogenide NCs for the following two reasons. The first reason is primarily cultural. In the field of IR detection, HgCdTe is the most investigated material, exhibiting high performance and strong tunability from

NIR to LWIR. As we switch from a bulk material to NCs, confinement plays the role of Cd alloying in the spectral response tuning. The second reason is related to the fact that among HgX NCs, this material has the lowest carrier density, which considerably reduces the dark current, leading to higher performance. Progress on the device side is generally strongly correlated with the material development. Heiss and co-workers first reported an absorption of up to 3 μm ,²⁷ and they promptly considered the potential of this material for photoconductivity.¹ The authors reported an elegant device based on an inkjet printed HgTe NC film. Using this approach, the authors drastically limited the amount of material required per device (2 μg per device). The authors reported a high response (up to 4.4 $\text{A} \cdot \text{W}^{-1}$) under extremely low power, suggesting the presence of electrical gain in the film, while the detectivity reached 3.2×10^{10} Jones for 1.4 μm peak detection at room temperature. Around the same period, Seong *et al.*² also reported a NIR device based on small HgTe NCs. The authors designed a photoconductive device with a film of HgTe NCs connected by ITO electrodes. The material was designed to present absorption at 1.3 μm corresponding to the current target for solar cell applications. At such wavelengths, HgTe NCs compete with lead chalcogenide NCs, which are more mature from both the synthesis and device perspectives. This fact is likely the reason why later, devices with longer wavelengths in the infrared region were considered to address a spectral range exempt from NC competitors. In particular, wavelengths above 2 μm are challenging to address with the use of lead chalcogenides because significantly large NCs with low colloidal stability are required. However, HgTe NCs remain small and exhibit their highest performance at such wavelengths. A critical breakthrough was obtained by Keuleyan *et al.*, who realized the synthesis of HgTe NCs with sizes up to 12 nm and an absorption cutoff wavelength reaching 5 μm at room temperature.³ The authors designed a device in which a thin film of such NCs was deposited onto interdigitated electrodes and demonstrated the first mid-infrared photoconductivity in NC arrays. With an absorption edge above 3 μm , the proposed device potentially enabled thermal detection. Many advancements have been made to improve this basic photoconductive device, including the combination of several cutoff wavelengths in a device to design multicolor devices^{41,42,45} or the introduction of light resonators to enhance the light absorption.^{37,39,40} These devices are discussed later in this

Table 6. Figures of Merit for Photoconductive Devices Based on HgTe NCs (PC = Photoconductor, PT = Phototransistor)

active material	device structure/type	spectral range (μm)	responsivity (mA·W $^{-1}$)	detectivity (Jones)	response time (s)	operating temperature (K)	specific feature	year/ref
HgTe	PC	1.4	4400	3.2×10^{10}	<0.1	RT	inkjet printed	2007 ¹
HgTe	PC	1.1	1.1×10^{-4}	N/R	2×10^{-2}	RT	$\text{H}_2\text{Te}/\text{HgClO}_4$ synthesis	2008 ²
HgTe	PC	3–6	100	2×10^9	< 10^{-7}	130	TOP:Te synthesis	2011 ²⁷⁹
HgTe	PC	3–5	250	2×10^9	N/R	100	TOP:Te synthesis	2011 ³
HgTe	PC	3.5	>100	3.5×10^{10}	< 10^{-5}	230	As_2S_3 ligand exchange	2013 ³⁵
HgTe	PC	1.1	500	2×10^{10}	10^{-6}	RT	spray coating	2014 ²⁸²
HgTe	PC	2.5	900	8×10^9	2×10^{-2}	RT	multicolor	2018 ⁴²
HgTe	PC	2.5	10	10^6	10^{-4}	RT	encapsulated	2018 ³⁰²
HgTe	PT	0.9	20	N/R	10^{-4}	RT	HgTe NPLs	2017 ²⁶²
HgTe	PT	2.2	200	2×10^{10}	10^{-4}	RT	sspray coating	2017 ³³⁰
HgTe	PT	2.5	N/R	N/R	10^{-5}	200	LaF_3 gate	2019 ³¹⁸
HgTe	PT	2.5	6.5	10^9	10^{-5}	220	2D/0D heterojunction	2020 ¹⁰⁸

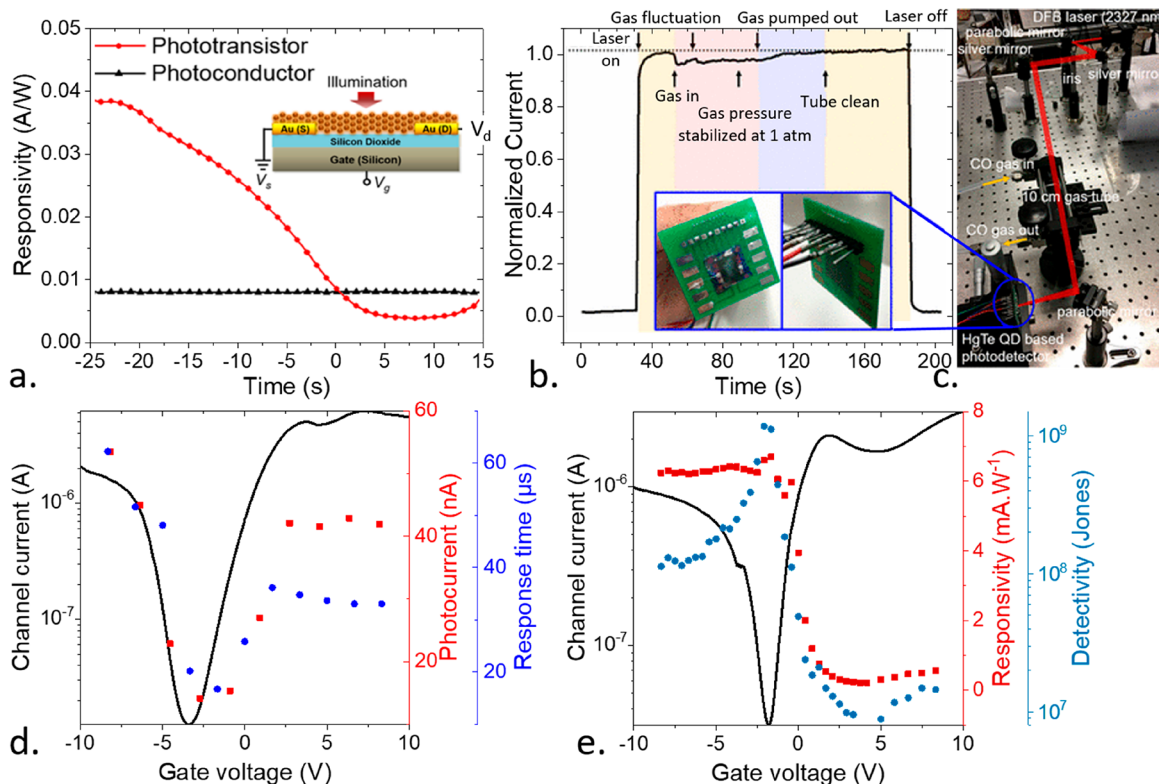


Figure 38. (a) Gate voltage dependent responsivity of a phototransistor in comparison to its responsivity at floating gate operation. Inset: shows a scheme of the device. (b) Normalized room temperature photocurrent response of a HgTe QD-based phototransistor during the CO gas sensing experiment. (c) Picture of the experimental setup of the gas sensing system. Inset in b: Photos of the packaged phototransistor. (a–c) Adapted with permission from ref 330. Copyright 2017 American Chemical Society. (d) Dark current, photocurrent, and response time as a function of the applied gate bias for a phototransistor made of a HgTe NC film, whose gate is made of LaF_3 ionic glass. (d) Adapted with permission from ref 318. Copyright 2019 American Chemical Society. (e) Dark current, photocurrent, and detectivity as a function of the applied gate bias for a phototransistor made of a HgTe NC film with graphene electrodes with a gate made of LaF_3 ionic glass. (e) Adapted with permission from ref 108. Copyright 2020 American Chemical Society.

chapter. Because of its simplicity, the progress related to photoconductive geometry strongly correlates with the material development. An interesting approach was proposed by Chen *et al.*³⁶ The authors used surface chemistry based on HgCl_2 and mercaptoethanol to achieve relatively high mobility films ($\approx 1 \text{ cm}^2 \cdot \text{V}^{-1} \text{s}^{-1}$) and correlated this enhancement in mobility to the increased detectivity. This result is not notably important for the reported performance; however, the result highlights the requirement of a high film quality, as shown in Figure 37b.

Interdigitated electrodes represent the most popular geometry for photoconductive devices;^{42,279,302} see Figure 37a. This planar geometry is robust and can accommodate low quality films resulting from thickness fluctuations and pinholes. Films with this geometry are achieved using common deposition techniques, such as drop casting,^{35,279} spin coating,³⁰² and spray coating.^{42,282,330} The interdigitated digits of the device act as a current amplifier. An advantage of photoconductive devices over their photovoltaic counterparts is the possibility of electrical

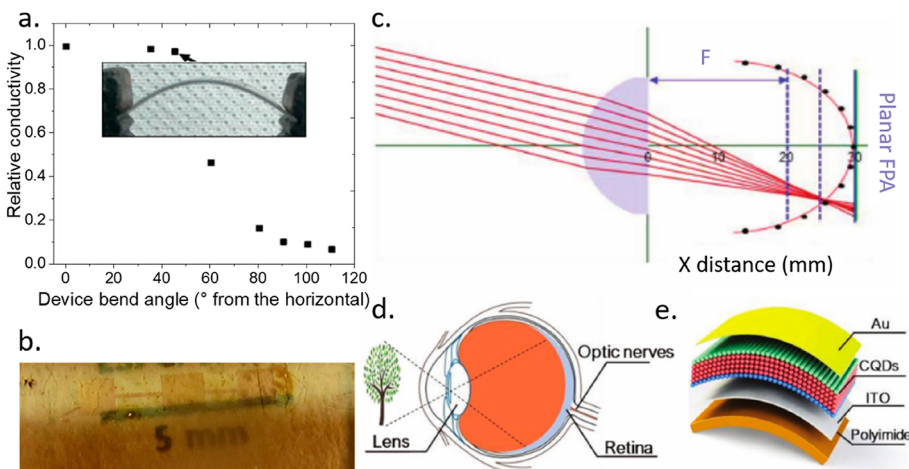


Figure 39. (a) Plot showing the relative (dark) conductivity of the device after it was bent to the angle indicated. Inset: an image of the substrate bent at 45°. (a) Adapted with permission from ref 42. Copyright 2018 American Chemical Society. (b) Image of a flexible intraband detector based on HgSe NCs. (b) Adapted with permission from ref 186. Copyright 2016 American Chemical Society. (c) Ray tracing analysis of a single lens. The black dots and red curve denote the focal surface of the lens. (d) Schematic of the ocular structure of humans. (e) Illustration of the device architecture of flexible HgTe CQD photovoltaic detectors. (c–e) Adapted with permission from ref 304. Copyright 2019 Wiley VCH.

gain.^{159,259,352,353} This process is often observed when one carrier has higher mobility than the other carrier. In this case, one carrier reaches the electrode faster than the other and is reinjected in the device to preserve the electrical neutrality of the film. Thus, a carrier (generated by a single photon) may contribute several times to the photocurrent. This aspect appears with a trade-off of a reduced detector bandwidth (defined by the longest carrier lifetime). This phenomenon is commonly referred to as the gain/bandwidth product trade-off in the electrical engineering community. This process is more efficient in small devices (<1 μm and less), which is challenging to achieve with interdigitated geometry.

Nevertheless, photoconductive devices based on HgTe have been used to design photoconductive detectors covering the 1–5 μm range with a stronger focus on the SWIR/extended SWIR (1.5–2.5 μm) and MWIR (3–5 μm) ranges. Table 6 reviews the figures of merit associated with these devices. A responsivity as high as a few A/W has been reported. However, responsivity on its own is an inferior figure of merit because it can be made arbitrarily high by increasing the bias and tuning the device geometry to generate a higher gain. The detectivity range is 10⁹–10¹⁰ Jones at RT in SWIR, and an identical detectivity is obtained in MWIR under cryogenic conditions (80–100 K). Interestingly, most studies indicate that HgTe photoconductive devices present a fast time (ms and less) response. This aspect is in contrast with PbS NC-based devices, for which longer decay times are commonly observed in the transient photocurrent. Livache *et al.* demonstrated in MWIR (*i.e.*, 5 μm cutoff wavelength) HgTe NC device response times of as small as 20 ns under resonant excitation,²¹⁹ as shown in Figure 37c.

6.2.2. Phototransistor. A phototransistor is the successful combination of a photoconductive film and an FET. Applying a gate bias leads to additional tunability compared to bare photoconductors. A basic motivation for the phototransistor is the tunability of the photoresponse with the gate, as depicted in Figure 38a. Such devices have been applied mostly to HgTe NCs^{108,318,330} and NPLs.²⁶² Chen *et al.* used such phototransistors for gas detection. The authors used the infrared range as a chemical fingerprint to detect CO (see Figure 38b,c). Such applications are particularly interesting for NC-based IR sensors because the use of powerful laser sources alleviates the

requirement for a low dark current. Nevertheless, the drift resulting from 1/f noise is a major limitation.

Phototransistors made of NCs exhibit two main limitations. While the phototransistor geometry can enhance the device capability to tune the photoresponse, the dark current is usually tuned in a similar manner, and the two currents generally follow one another,^{262,318,330,354} as shown in Figure 38d. Consequently, applying a gate to increase the photoresponse increases the dark current. Alternatively, minimizing the dark current reduces the photoresponse of the device. In this case, the gate is not used to examine the nature of the majority carrier but rather as a key to prepare the material in a given carrier density state. In particular, the gate can be used to balance an initial doping of the film and reduce the dark current by exploiting the small offset of the dark current and photoresponse as a function of the applied gate bias. Consequently, this device is used as a patch to compensate for the limited material control at the synthesis level. The second limitation is from the response time; see Figure 38d. As the photoresponse is increased, the response is slowed. In other words, phototransistors operate at a quasi-constant gain bandwidth product. It may be interesting to slightly concede on the response time to generate an increased response, especially with HgTe, which presents a fast response. The only exception to this behavior was reported by Numbé *et al.*, who used graphene electrodes instead of ordinary metallic electrodes.¹⁰⁸ Using this concept, the authors operated the transistor in a nonconventional mode, in which a large drain source bias was used to generate a p–n junction instead of a uniform doping of the film. This concept has been extensively used to generate light from a FET,^{355–361} although its application to detection using NCs has been rare. By forming such gate controlled p–n junctions, the authors could identify the operating conditions under which the minimal dark conductance corresponded to the maximal photoresponse (see Figure 38e), thereby enhancing the overall detectivity of the device.

A current limitation of phototransistors is the film thickness. FETs based on surface gating methods can only tune the conductance of the film in the vicinity (typically over ≈50 nm) of the gate interface. Such thin films cannot effectively absorb incident light, which limits the photoresponse of the device. Coupling the phototransistor to the resonators can likely help

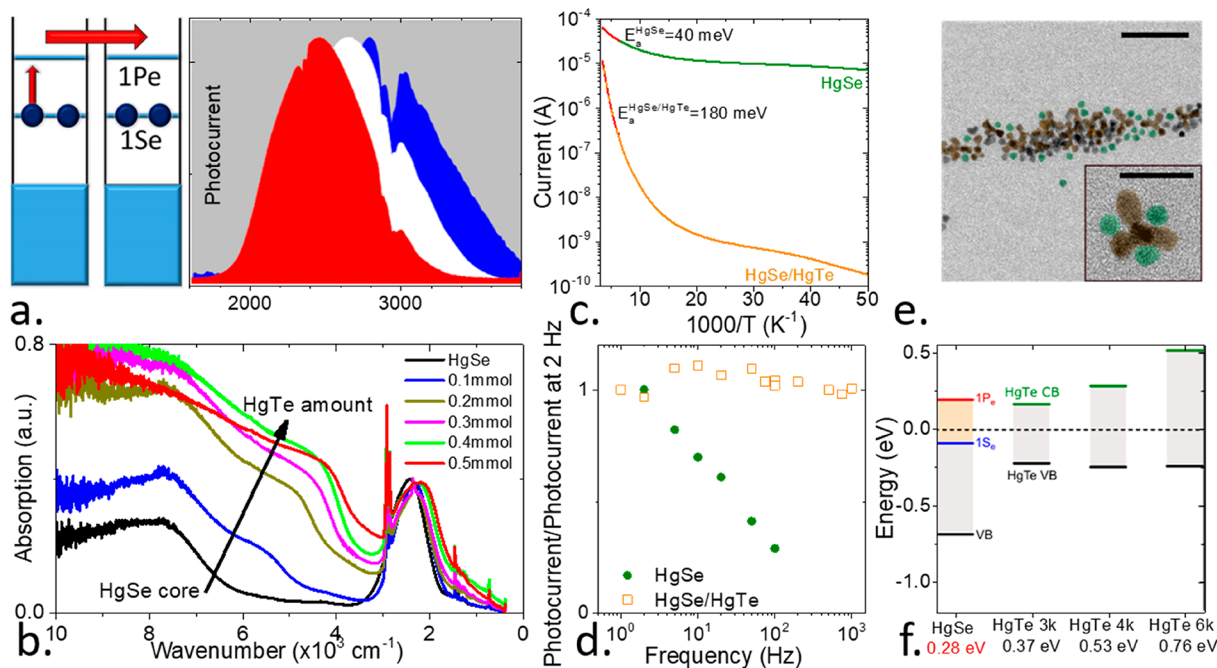


Figure 40. (a) Principle of phototransport based on intraband photoconduction (left) and intraband photoconduction in an array of HgSe NCs with various sizes (right). (a) Adapted with permission from ref 34. Copyright 2014 American Chemical Society. (b) Absorption spectra of HgSe cores and HgSe/HgTe core–shell structures for different amounts of HgTe. The intraband signal is barely affected by the addition of HgTe, while the interband signal is amplified and red-shifted. (c) Current as a function of temperature for thin films made of HgSe cores and HgSe/HgTe heterostructures. The Arrhenius fit close to the room temperature range is highlighted in red. (d) Bode diagram (normalized signal magnitude at 2 Hz) as a function of the signal frequency for thin films made of HgSe cores and a HgSe/HgTe heterostructure. (b–d) Adapted with permission from ref 120. Copyright 2018 American Chemical Society. (e) TEM image of mixed HgSe and HgTe quantum dots for random colloidal QWIP fabrication. Certain HgSe (respectively HgTe) QDs are indicated in green (respectively brown). The scale bar is 50 nm. Inset: higher resolution TEM image showing a HgTe tetrapod along with three HgSe spheres. The scale bar is 20 nm. (f) Electronic spectra of the four materials used for this study determined using a combination of XPS measurements and optical spectroscopy. The bandgap of each material is given on the x axis (the intraband gap is given for HgSe). The black dashed line represents the Fermi level. (e,f) Reproduced with permission from ref 47. Copyright 2019 Nature.

overcome the absorption trade-off, this strategy has just been reported by coupling an STO gate with a metallic grating.³³⁹ The alternative approach is the use of electrolyte gating, which enables the gating of thick films. The method has been applied to the design of HgTe NPL-based phototransistors.²⁶²

6.2.3. Flexible Devices. Among potential additional functions for infrared sensors based on HgX NCs, the concept of flexible³⁶² substrates was introduced early for both FETs^{315,363} and detectors,⁴² see Figure 39a,b for an example of such a device. This effort mostly concerns HgTe^{304,364} but also works for HgSe NCs.¹⁸⁶ In most reports, performance are tested as a function of the bending radius and number of cycles. However, the motivation for such flexible devices remains mostly unclear. Current developments for detection involve imaging applications, which require a readout circuit that is unlikely to be flexible. This aspect is even more unlikely if the device requires cooling with a Peltier or cryogenic machine. Nevertheless, one reports for a device based on a flexible substrate which was properly motivated. Tang *et al.* proposed the design of an eyelike detector;³⁰⁴ see Figure 39c–e for a scheme of the concept. This work was initially motivated by the authors' strategy to conduct imaging using a single pixel detector with a scanning lens in front of the detector to form an image. Nevertheless, despite the clear success of this approach, distortion occurs, and a curved detector is usually introduced to counterbalance the distortion introduced by the scanning method.

6.3. Intraband Detection

While intraband transitions have been observed since the 1990s, the procedures used to induce such transitions (saturation of the interband transition^{365,366} or electrochemical gating³³²) were incompatible with practical device integration. Observation of the intraband in the steady state,^{65,66} because of self-doping in HgS, has been a breakthrough.³³ Promptly after this observation, intraband photoconduction was investigated in HgSe^{29,34} and later in HgTe^{30,63,96} or PbS.²²⁷ In these cases, photoexcitation occurs between the 1S_e and 1P_e states in the conduction band. This transition is the 0D equivalent of the intersubband transition in doped quantum wells. In HgSe with particle sizes of approximately 5 nm, this transition overlaps with the 3–5 μm range (MWIR); see Figure 40a.

This transition yields opportunities to develop an effective strategy for IR detection while reducing toxicity. Current narrow bandgap interband transitions are mostly observed using Pb- and Hg-based materials and heavy atom containing materials. Narrow bandgaps can be more easily obtained from materials with a large number of electrons than from those with light atoms for which the density of the states is smaller. Thus, achieving doping in moderately doped nanocrystals may clarify intraband transitions in materials with reduced toxicity. Another key advantage of intraband transitions, compared with interband transitions in a semimetal, is the reduced sensitivity of the spectral response to sintering. In HgTe, owing to the lack of a bandgap and the extremely large Bohr radius (40 nm), annealing leads to a dramatic red-shift of the optical bandgap. This aspect

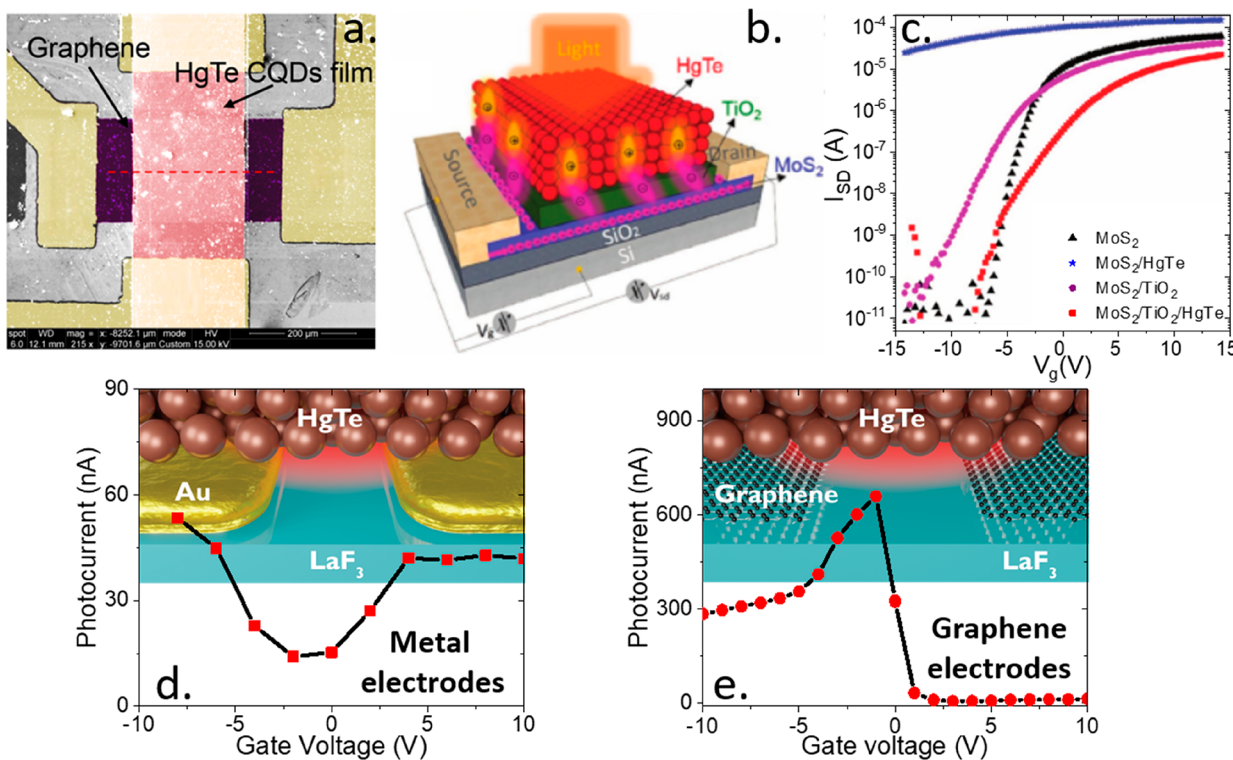


Figure 41. (a) Faux color of a graphene/HgTe CQD junction. (a) Adapted with permission from ref 375. Copyright 2019 American Chemical Society. (b) Scheme of hybrid device under light illumination and photogenerated carrier separation and transport within those photodetectors. (c) Transfer curves (drain source current as a function of the gate current) of pristine MoS₂ and MoS₂/HgTe with and without the TiO₂ from the photodetector devices presented in (b). (b,c) Reproduced with permission from ref 377. Copyright 2017 Wiley VCH. (d) (respectively e) Photocurrent as a function of the gate bias for gold/HgTe/gold (respectively graphene/HgTe/graphene) junction. The top part shows the scheme of the device. (d,e) Adapted with permission from ref 108. Copyright 2020 American Chemical Society.

not only affects the detection cutoff wavelength but also increases the dark current. In this sense, the intraband transition appears to be more robust to mild annealing. However, it has been demonstrated that the carrier mobility is more sensitive to the size distribution in the case of intraband HgSe than that for interband HgTe NCs.¹²¹

However, many researchers were skeptical regarding the possibility of achieving high performance with intraband detection. Similar concerns were raised for intersubband transitions in doped quantum wells. The intraband transition is facilitated by doping, which also modifies the dark current. In doped particles, the photoinduced change in the carrier density is reduced compared to that in interband materials. This aspect, as well as the large dark current associated with these materials, explains why, to date, the detectivity of intraband detectors remains below 10⁹ Jones.⁵⁷ The large dark current is not the only issue observed for an array of intraband nanocrystals. The photoresponse in HgSe arrays appears to be slow, with a response time of the order of seconds. This aspect raises concerns that the photoresponse may actually result from a bolometric effect rather than actual photoconduction. It has been claimed that a fast response time can be achieved when the doping magnitude corresponds to full filling of the 1S state.³⁴ This aspect appears to be practically difficult to achieve because postsynthesis ligand exchange also dramatically affects film doping.¹⁸⁶ Finally, the dark current activation energy was found to be extremely small (20–40 meV), suggesting that sample cooling may not be able to reduce the dark current in the HgSe-based detectors.

To overcome these difficulties, the most recent strategies focus on heterostructures based on narrow bandgap materials, such as HgSe and HgTe. The driving concept was to couple the intraband absorption derived from HgSe with the interband (*i.e.*, low dark current) transport properties from HgTe. The concept was inspired by III–V heterostructures, in which absorption and transport are spatially uncoupled.^{367–369} The first approach was proposed by Goubet *et al.*¹²⁰ On a HgSe core, a coating of HgTe is colloidally grown (see Figure 11*i*–*l*). The addition of the HgTe layer does not considerably influence the energy of the intraband transition, while the interband absorption exhibits a strong red-shift (see Figure 40*b*). Because of degenerate doping, the interband transition between the 1S_h and 1S_e states of HgSe is bleached due to the lack of an arrival state. However, the interband absorption coupling the valence band and the 1P_e state is still possible. HgTe valence band is closer to the vacuum level than HgSe, and its addition leads to a second path for interband absorption coupling of the valence band of HgTe with the 1P_e state of HgSe. Owing to the relative position of the HgTe and HgSe valence bands, this absorption occurs at a lower energy, explaining the observed red-shift. The introduction of HgTe increases the resistance of the sample, and the dark current activation energy reaches values higher than 150 meV (Figure 40*c*), leading to a higher benefit of sample cooling. The intraband photoresponse was evaluated using a quantum cascade laser as a selective excitation source of the intraband transition. The authors observed a significant acceleration of the intraband response (Figure 40*d*). Using an alternative approach, the same group demonstrated that similar results could be obtained while eliminating the constraint of epitaxial growth

between HgSe and HgTe.⁴⁷ The mixing of two solutions of HgSe NCs with intraband absorption and HgTe without intraband absorption (Figure 40e) yields similar results (higher resistance, higher activation energy, and faster response) while allowing more freedom for the metamaterial design. The bandgap of HgTe is thus a free parameter that can be optimized to favor charge transfer without generating dark current (see Figure 40f). Finally, the authors reported on the first colloidal intraband diode. The latter device is composed of a sapphire substrate, on which an Al grid is deposited and used as a partial mid-infrared transparent electrode. A layer of a HgSe/HgTe mixture with intraband absorption is coupled to a unipolar barrier made of HgTe NCs with a higher bandgap; see the scheme of the device in Figure 45h. The second electrode is made of gold. Interestingly, the diode exhibits a complex spectral photoresponse, in which the relative contribution of interband and intraband signals can be strongly modulated with the applied bias; see Figure 45i. Using this approach, the performance of intraband detectors has been enhanced to be similar to that of interband materials. This approach is promising, especially for longer wavelengths.

6.4. Coupling to 2D Materials

The concept of coupling nanocrystals with 2D materials was first proposed by Konstantatos *et al.*³⁷⁰ The motivation was to enhance the low mobility of nanocrystal arrays by coupling nanocrystals with graphene: the large carrier mobility of graphene ($\approx 1000 \text{ cm}^2 \cdot \text{V}^{-1} \cdot \text{s}^{-1}$ at room temperature, while the NC arrays achieved values $\ll 1 \text{ cm}^2 \cdot \text{V}^{-1} \cdot \text{s}^{-1}$) was combined to the large spectral tunability of NC absorption. The design was first used to couple graphene with NIR PbS nanocrystals, typically for the design of NC-based solar cells. In such devices, absorption in the visible- and near-IR regions occurs mostly in the nanocrystals. Depending on the exact band alignment with graphene, one type of carrier is preferentially transferred to graphene, while the second carrier is trapped in the NC array. To ensure the neutrality of the whole structure, the first type of carrier recirculates in graphene as long as the trapped carriers are not recombined, generating gain. The latter process is mostly driven by the recombination of these trapped charges with external molecules, which makes such devices sensitive to air and moisture. The recirculation process, also known as photo-reduction gain,³⁵² has been applied in the domain of nanocrystal arrays^{159,353} and was optimized due to the high mobility of graphene, which accelerated the recirculation process. Consequently, extremely large responsivities ($> 10^6 \text{ A} \cdot \text{W}^{-1}$) could be achieved.^{370,371} Following this result, the field of 0D/2D mixed dimensionality van der Waals heterostructures³⁷² rapidly expanded³⁷³ and led to the use of other types of nanocrystals.³⁷⁴ The coupling of graphene with HgTe³⁷⁵ and HgSe³⁷⁶ nanocrystals was demonstrated by Tang *et al.*, who considered the shift of the spectral response toward the mid-infrared region; the scheme of this device is shown in Figure 41a.

Despite this large response, graphene/QD heterostructures also suffer from certain key limitations. First, the high mobility of graphene is dramatically balanced with the lack of a bandgap. Indeed, the carrier density in graphene, despite being easily gate tunable, is larger than the thermally activated carrier density in a wide bandgap semiconductor. Although this claim is probably less true in regard to infrared materials, it results in a dark current and its associated $1/f$ noise,³⁷⁸ which is large in graphene QD heterostructures. In addition, the response time of these heterostructures generally presents a slow component (of the

order of seconds and sometimes, even hours), which is incompatible with the practical use of such devices. Last, the device presents an extremely low saturation with respect to the photon flux, and a high responsivity is only achieved for an extremely low irradiance. To reduce the carrier density, graphene was replaced by other 2D materials corresponding to a reasonable bandgap. For instance, graphene can be replaced by MoS₂.³⁷⁷ This transition metal dichalcogenide material has a slightly lower mobility, but the 2 eV bandgap allows it to overcome the dark current issues. As depicted in Figure 41c, the current tunability obtained with this gate is several orders of magnitude, compared to the factor of 2–5 obtained using graphene. Charge transfer between the conducting and absorbing layers can be further realized by the introduction of a TiO₂ spacer (see Figure 41b).

The most recent ideas for 2D/HgX devices do not focus on using 2D materials as high mobility transport layers but rather on other parts of the device, such as transparent conductive contacts to replace ITO or their gate tunable work function. Thus, Tang *et al.* proposed a dual color visible/SWIR detector combining a Si-based diode with a HgTe nanocrystal layer, in which the intermediate contact is graphene.⁴³ Numbé *et al.* proposed the use of graphene as an infrared contact to design planar p–n junctions, whose channel is made of a HgTe layer; see Figure 41d,e. The authors attempted to exploit the large quantum capacitance of graphene. The authors demonstrated that a phototransistor made of a high capacitance ionic glass gate coupled to a HgTe absorbing layer exhibits an extremely different behavior when using metallic contacts or graphene as electrodes. Because graphene is extremely thin, it enables the gate-induced electric field to propagate in the nanocrystal film, while the gating effect is completely screened in the case of gold electrodes. Consequently, the authors succeeded in generating a planar p–n junction that enhances the charge dissociation and boosts the detectivity by 2 orders of magnitude.¹⁰⁸

6.5. Photovoltaic Devices

Photovoltaic (PV) devices, typically based on vertical stacks, present the key advantage of a built in electric field, allowing for zero volt operation, leading to a large reduction in the dark current (and associated noise).³⁰¹ Despite this key motivation, only a limited amount of work has been dedicated to this aspect. Such devices exhibit performances that are competitive compared to commercial devices, especially in the SWIR and extended SWIR ranges.⁴⁸

Vertical geometry based infrared devices involve concerns regarding the design of transparent conductive electrodes and substrates. For visible and NIR ranges, the use of transparent conductive oxides (TCOs) is the most commonly implemented strategy. Tin-doped indium oxide (ITO) and fluorine-doped tin oxide (FTO) are the most commonly used TCOs. In the case of HgTe (work function = $4.6 \pm 0.1 \text{ eV}$),^{49,122,192} FTO (work function = 4.4 eV) is better suited than ITO (work function = 4.7 eV) for use as an electron contact. However, FTO is typically deposited by chemical vapor deposition (CVD), which cannot be realized in most laboratories. Consequently, FTO is used as a bottom contact in a limited manner. Another limitation is the corresponding absorption in the infrared range, especially above $1 \mu\text{m}$. A possible strategy to overcome this limitation is to reduce the TCO thickness³⁰² at the price of a higher contact resistance. Above $2\text{--}2.5 \mu\text{m}$, glass absorption also becomes problematic. In this case, glass must be replaced with IR transparent materials, such as CaF₂³⁰¹ or sapphire³⁰³ (Al₂O₃). Among these materials,

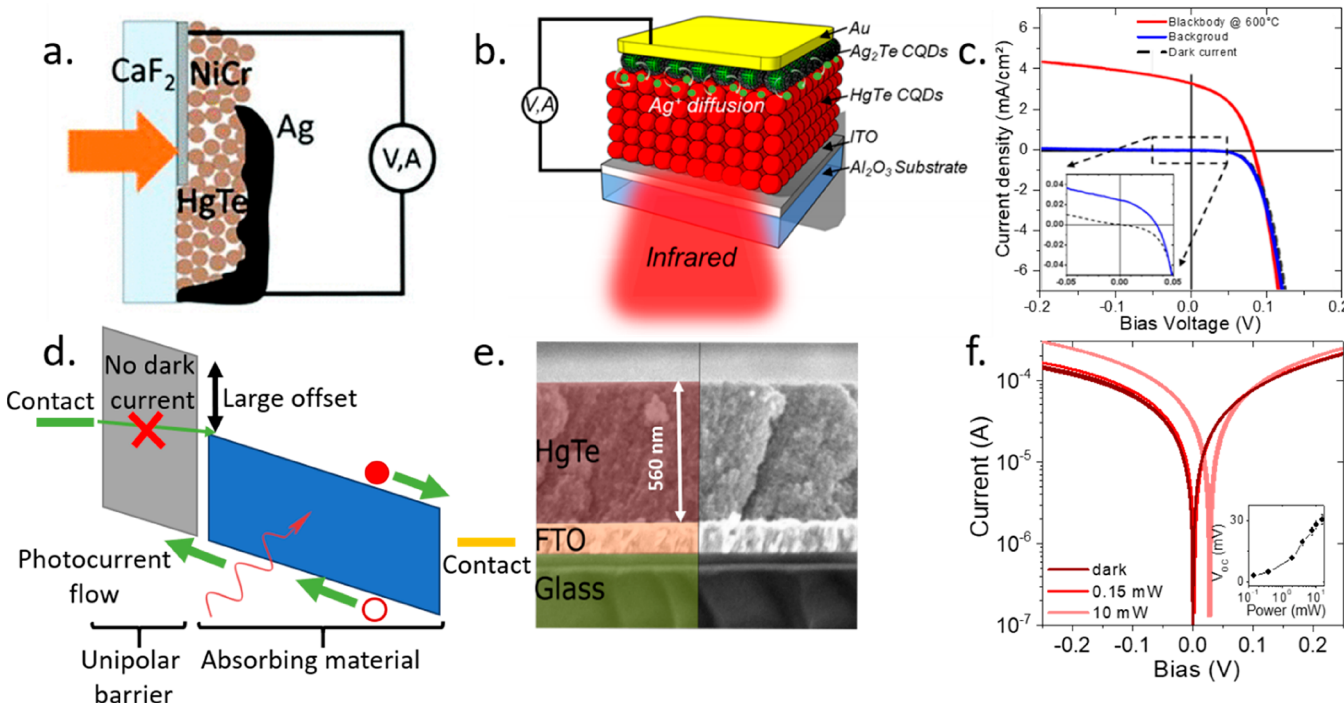


Figure 42. (a) Schematic of a MWIR photovoltaic structure based on the $\text{CaF}_2/\text{NiCr}/\text{HgTe}/\text{Ag}$ structure. Reproduced with permission from ref 301. Copyright 2015 AIP Publishing. (b) Schematic of a MWIR photovoltaic structure based on the $\text{Al}_2\text{O}_3/\text{ITO}/\text{HgTe}/\text{Ag}_2\text{Te}/\text{Au}$ structure. (c) Current density as a function of the bias voltage under dark conditions, background radiation, and $600\text{ }^\circ\text{C}$ blackbody radiation at an operation temperature of 85 K of the diode presented in (b). Inset: figure provides a zoomed in view of the dark current (closed shield) and background current (open shield). (d,e,f) Adapted with permission from ref 303. Copyright 2018 American Chemical Society. (d) Adapted with permission from ref 301. Copyright 2015 AIP Publishing. (e) SEM image of an extended SWIR photovoltaic structure based on the glass/FTO/HgTe 4k/HgTe 6k/Au structure. (f) I - V curves in the dark and under illumination by a $1.55\text{ }\mu\text{m}$ laser at different powers for the diode shown in (e). Inset: evolution of open circuit voltage (V_{OC}) as a function of increasing power. (e,f) Reproduced with permission from ref 146. Copyright 2019 Wiley VCH.

sapphire is an optimal candidate for low cost sensors, at least as long as the targeted wavelength remains below $5\text{ }\mu\text{m}$. In the MWIR range, the use of a thin (5 nm) metallic layer was proposed as a transparent contact. However, thin metallic layers are not continuous, leading to high contact resistances. Thus, Tang *et al.* proposed the use of a thin gold layer coupled to a thin ITO layer to achieve both high transparency and low contact resistance.³⁸ Livache *et al.* proposed another approach for the design of MWIR semitransparent (70% transparency) conductive contacts with a metallic grid on top of a sapphire substrate.⁴⁷ The contact design is expected to become an increasingly active field in the coming years, especially with the intense integration of resonators into devices to enhance light absorption. In this regard, resonators may reduce the constraint of transparency but raise the concern of plasmon losses.

The development of HgTe NC-based PV devices was initially motivated by the design of near-IR absorbers for solar cells.^{2,91,379,380} Extremely small HgTe NCs with the band edge at approximately the optimal value for a single junction solar cell have been grown and studied, and the design of vertical devices is clearly inspired by organic PV devices. Nam *et al.*³⁷⁹ proposed the use of HgTe NCs blended with a narrow bandgap electron donor polymer:³⁸¹ poly[(4,4'-bis(2-ethylhexyl)dithieno[3,2-*b*:2',3'-*d*]silole)-2,6-diyl-*alt*-(2,1,3-benzothiadiazole)-4,7-diyl] (PSBTBT) to obtain a bulk heterojunction. *A posteriori*, the p-type nature of small HgTe NCs does not favor the creation of a bulk heterojunction by mixing the materials with an electron-donating polymer. Im *et al.*³⁸⁰ proposed a HgTe NC-based PV device inspired by solar cells, using HgTe NCs as absorbers for a

dye-sensitized solar cell.^{382,383} The HgTe layer is sandwiched between a TiO_2 layer and P3HT layer used as an electron extractor and hole extractor, respectively.⁹¹ The presence of a polymer layer hinders the extension of the photoresponse of the device toward longer wavelengths due to the organic absorption. Consequently, Jagtap *et al.* later proposed a similar structure in which larger HgTe NCs were used to absorb light in the extended SWIR,³⁰² replacing the polymer layer with a thin inorganic transport layer of MoO_3 .¹⁹² However, the responsivity of the diode based on FTO/ TiO_2 /HgTe/ MoO_3 /Au remains weak. This aspect was attributed to the TiO_2 layer, which can help induce a rectifying behavior to the I - V curve^{192,302} but also strongly filters the photogenerated electrons, leading to low responsivity ($\ll \text{mA}\cdot\text{W}^{-1}$).

Later, the use of wide bandgap layers for hole and electron extraction was avoided. HgTe and wide bandgap materials presented a large band alignment mismatch. Consequently, the wide bandgap led to barriers in both the valence and conduction bands. The layers filtered not only the dark current but also the photocurrent. Guyot-Sionnest *et al.*³⁰¹ proposed a Schottky diode in which the HgTe NCs were connected to contacts made of NiCr and Ag, with the latter contact made of silver paste; see Figure 42a. Despite this simplistic fabrication, no electrical short was reported. This diode is the first reported to achieve background limited operation (in a background limited infrared photodetector (BLIP), the noise is limited by the fluctuation of the arriving photons).

Later, Ackerman *et al.*³⁰³ reported several key advancements in the design of HgTe NC-based PV devices; see Figure 42b.

Table 7. Figures of Merit Associated with Photovoltaic Infrared Devices Based on HgX NCs (RT = Room Temperature)

active material	device structure/type	spectral range (μm)	responsivity or EQE (mA·W $^{-1}$ or %)	detectivity (Jones)	response time (s)	operating temperature (K)	specific feature	year/ref
HgTe	FTO/TiO ₂ /HgTe/P3HT/Au	NiR 0.8–1.4	1.3%		$\approx\text{ms}$	RT	depleted heterojunction	2012 ³⁸⁰
HgTe	ITO/PEDOT:PSS/HgTe:PSBTBT/Al	Vis-NiR 0.4–1.1	1.4%			RT	bulk heterojunction	2014 ³⁷⁹
HgTe	ITO/TiO ₂ /HgTe/Au	E-SWIR	5×10^{-3}	$\approx 10^7$	$\approx\mu\text{s}$	RT	air stable via device encapsulation	2018 ³⁰²
HgTe	FTO/TiO ₂ /HgTe/HgTe-p-type/MoO ₃ /Au	E-SWIR	0.2	3×10^8	$\approx\mu\text{s}$	RT	unipolar barrier	2018 ¹⁹²
HgTe	FTO/HgTe/HgTe p-type/Au	E-SWIR	20	3×10^9	370 ns	RT	thick film made from ink	2019 ¹⁴⁶
HgTe	polyimide/ITO/Bi ₂ Se ₃ /HgTe/Ag ₂ Te/HgTe/Au	SWIR	220	7.5×10^{10}	260 ns	RT	flexible substrate	2019 ³⁰⁴
HgTe	Al ₂ O ₃ /ITO/HgTe/Ag ₂ Te/Au	SWIR	≈ 1000	$\approx 10^{11}$	1.4 μs	RT	high detectivity at RT, cross-linking with EDT/HCl	2020 ⁴⁸
HgTe	LaF ₃ /graphene gate(FET)	E-SWIR	1	10^9	<10 μs	220	gate induced p-n junction	2020 ¹⁰⁸
HgTe	CaF ₂ /NiCr/HgTe/Ag	MWIR	38	10^{10}	0.7 μs	140	BLIP detection	2015 ³⁰¹
HgTe	Al ₂ O ₃ /ITO/HgTe/Ag ₂ Te/Au	MWIR	≈ 100	3×10^9	20 μs	@RT	soping by contact layer, high detectivity	2018 ³⁰³
				380	$\approx 10^{11}$	$\approx 1 \mu\text{s}$	@85	
HgTe	Al ₂ O ₃ /ITO/Bi ₂ Se ₃ /HgTe/Ag ₂ Te/Au	SWIR MWIR	400	3×10^{10}	2.5 μs	85	bicolor SWIR/MWIR	2019 ⁴⁶
HgTe:HgSe	Al ₂ O ₃ /Al/HgTe/HgSe:HgTe/Au	MWIR-SWIR	≈ 5	$\approx 10^9$	200 ns	80	quantum well like structure with hybrid CQD. intraband detection	2019 ⁴⁷

The most critical advancement is the introduction of a layer made of Ag₂Te NCs used as a hole extractor. This layer is exposed to the Hg precursor, which leads to a cation exchange process to form a strongly p-doped layer of HgTe/Ag₂Te. The advantage of this structure is that it presents a clear diodelike transport behavior (see Figure 42c) without using wide bandgap materials, which may filter the photocarriers. Later, the success of this structure was expanded to a bicolor diode⁴⁶ and SWIR operation.⁴⁸ To date, this structure leads to the highest performances (see Table 7). For electron extraction, the same group proposed the use of Bi₂Se₃ as a strongly n-type layer with a small bandgap,^{48,304} although this layer was eliminated in their recent work.⁴⁸ This elimination was likely a result of the low colloidal stability of the large Bi₂Se₃ plates, which makes the fabrication of smooth films challenging. In parallel with the structure design, significant progress has been made in the surface chemistry of nanocrystals, switching from conventional solid-state ligand exchange mostly conducted using ethanethiol to liquid ligand exchange using a mixture of HgCl₂ and short thiols (EDT or mercaptoethanol).^{36,146} In particular, Martinez *et al.* proposed the use of an ink in which NCs are dispersed in a polar solvent to realize the deposition of nanocrystals. In this case, the HgTe NCs capped with this mixture of ligands are phase transferred²⁴ toward a polar solvent (DMF). Because of the high boiling point of DMF, it becomes

critical to use highly concentrated solutions of NCs (>250 mg·mL $^{-1}$) to facilitate the film drying. The key interest for this ink is to reduce the highly repetitive process of deposition/ligand exchange required by solid-state ligand exchange, which often leads to the generation of holes or cracks in the film. In addition, inks with high concentrations enable the deposition of thick films (>300 nm) in a single step; the cross-section of a device with a film thickness of approximately 600 nm is shown in Figure 42e.¹⁴⁶ The increase in the film thickness over recent years has been a key aspect to increase the device absorption. Pushing the film thickness from 200 to 560 nm increases the absorption from 12% to 40%, assuming a single pass of light in the absorbing layer.

To complete the discussion, a complementary approach for the design of HgTe NC-based PV devices relying on unipolar barriers must be mentioned. This concept was first developed for III–V heterostructures^{384,385} and is depicted in Figure 42d. The absorbing layer is coupled to a barrier that is used to prevent dark current injection while allowing the photocurrent to flow. To achieve this goal, a specific band alignment is required: the photocarrier conducting band of the active material must be resonant with that of the barrier, while the other band must feature a large offset, preventing the dark current. Such a design requires an accurate understanding of the band offset between the two materials, which is possible for III–V semiconductors

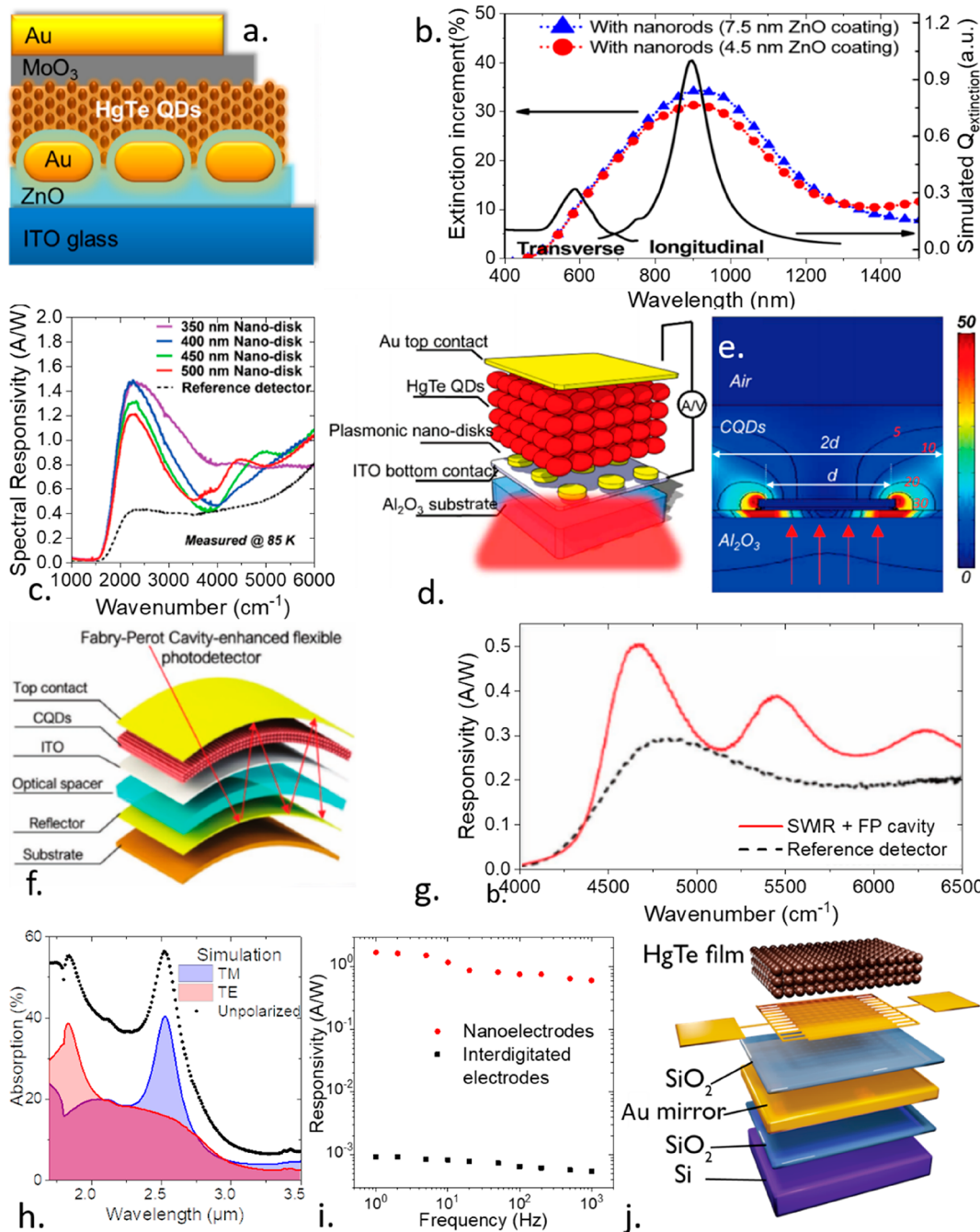


Figure 43. (a) HgTe QD/ZnO heterojunction photodiode with Au nanorods embedded within the ZnO layer. (b) Extinction increments of HgTe QD/ZnO films with ZnO-coated Au nanorods normalized by the extinction spectrum of the control sample. The normalized FDTD simulated extinction cross-section spectra with the longitudinal and transverse LSPRs of Au nanorods coated with 7.5 nm ZnO in the HgTe QD layer are also shown. (a,b) Adapted with permission from ref 389. Copyright 2014 American Chemical Society. (c) Measured spectral responsivity of HgTe CQD MWIR detectors with and without plasmonic disks. (d) Illustration of the HgTe CQD midwave infrared detector with plasmonic disks and a 30 nm Au contact. (e) Mapping of the light intensity enhancement ratio. The wavelength of incident light is $\approx 4 \mu\text{m}$. (c-e) Adapted with permission from ref 38. Copyright 2018 American Chemical Society. (f) Illustration of Fabry-Perot cavity-enhanced HgTe CQD detectors. (g) Spectral responsivity of the SWIR HgTe detector with and without the Fabry-Perot cavity. (f,g) Reproduced with permission from ref 304. Copyright 2019 Wiley VCH. (h) Simulated absorption spectra relative to the TE mode, TM mode, and total absorption for a HgTe NC-based device with a $1.8 \mu\text{m}$ period. (i) Responsivity as a function of the light signal frequency for devices based on HgTe nanocrystals using interdigitated electrodes made of gold contact deposited on a Si/SiO₂ substrate and interdigitated electrodes on a Au/SiO₂ substrate designed to induce GMR with a $1.8 \mu\text{m}$ period. (j) Schematics of the cross-section of the conventional interdigitated electrodes deposited on a Si/SiO₂ substrate. (h-j) Adapted with permission from ref 40. Copyright 2019 American Chemical Society.

but challenging for II–VI semiconductors. The band alignment results from the inverted band structure, quantum confinement, and surface chemistry. To this end, Jagtap *et al.* conducted a

systematic study of the band alignment in HgTe NCs as a function of the size and surface chemistry (see Figure 23b,c). The authors observed that the valence band offset for various

sizes of HgTe is small, and most of the confinement energy is carried by the shift of the conduction band. This observation reflects the higher effective mass of the valence band compared with that of the conduction band; see the dispersion diagram in Figure 14c or Figure 15b. Consequently, a thin layer made of smaller HgTe NCs (*i.e.*, higher bandgap) behaves as a unipolar barrier (no offset for valence band, large offset for conduction band) for larger NCs. Jagtap *et al.*¹⁹² introduced a thin layer of HgTe with a 6000 cm^{-1} ($\approx 720\text{ meV}$ or $\approx 1.8\text{ }\mu\text{m}$) band edge energy in a device in which the absorbing layer is made of HgTe with a 4000 cm^{-1} bandgap ($\approx 480\text{ meV}$ or $\approx 2.5\text{ }\mu\text{m}$). The introduction of this thin layer leads to a diode behavior for the I – V curve, with the presence of an open circuit voltage under illumination^{47,146} at room temperature (see Figure 42f). This barrier also increases the detectivity by a factor of 10.¹⁹²

To date, PV devices have exhibited the highest detection performance. Advancements have been made by combining the developments in surface chemistry and PV device design. In the next section, we discuss how the introduction of resonators has enhanced the performance due to better light–matter coupling and enhancement of device absorption. Currently, the design of energy band profiles suffers from a lack of rationality. Thus, optimal electrical properties might be realized in the future. The other key challenge relative to PV devices is the transfer of the diode structure toward FPAs. For example, gold is used as a contact in most devices, even though this metal is not CMOS compatible. A systematic investigation of the effect of the contact work function must be conducted in the future.

6.6. Enhanced Light Matter Coupling

In NC arrays, achieving decent photoconduction comes at a cost. Because of hopping transport, the mobility in NC arrays is limited. Although significant progress^{29,35,194,317} has been made in the direction of increased mobility values^{4,24,386} from below 10^{-4} to approximately $1\text{ cm}^2\cdot\text{V}^{-1}\cdot\text{S}^{-1}$ using ink-based films with hybrid passivation (ions and organics) to passivate different facets, the value of mobility remains 2 to 3 orders of magnitude below the bulk values. Consequently, the charge diffusion length extends over several NCs (10–100 nm) but remains shorter than the absorption depth (few μm) in such films. As discussed earlier in this review, films with thicknesses of a few micrometers are required to absorb most of the incident light. For instance, a 200 nm thick film, which is the typical thickness of films obtained from solid-state ligand exchange, with a band edge at $2.5\text{ }\mu\text{m}$ (0.5 eV or 4000 cm^{-1}) absorbs only 12% of the incident light for a single pass of the light. The trade-off is a result of the fact that building thicker films is not useful because the charges cannot be collected.

Two approaches can be used to overcome these severe limitations. The first approach is to work on the material side to fabricate higher quality (*i.e.*, mirror-like, with a film roughness of the order of the NC size) thick (>500 nm) films with improved internanocrystal coupling (*i.e.*, higher μ). Such efforts have been conducted by considering the design of photodiodes in vertical geometry,³⁰³ and HgTe-based inks have been used to facilitate thick film deposition with a reduced number of steps.^{36,146} The second method to overcome the diffusion length/absorption depth trade-off is to control the light–matter coupling within the NC film. The driving concept is to modify the electromagnetic field to concentrate it into the absorbing layer. Resonators can be introduced, enabling an increase in the magnitude of the incident field in a thin slab of the absorbing semiconductor, resulting in an increase in absorption. The field

enhancement can be considered to be a reduction of the effective absorption depth to eliminate the previously mentioned trade-off. To date, most efforts have been focused in this direction. Another aspect that has not been extensively examined for NCs is the planar focusing of the light to uncouple the optically absorbing area of the device from its electrical area. The concept is to collect photons over an area that is as large as possible while reducing the electrical area as much as possible, thereby proportionately decreasing the dark current.³⁸⁷ Zhu *et al.* recently demonstrated³⁸⁸ a device based on a silicon photonic waveguide with a small volume of HgTe used as an IR absorbing material. This structure shows an enhanced detectivity compared to the bare HgTe film.

The coupling of resonators with HgX NC films is, for most devices, based on the introduction of plasmonic resonators. The first demonstration was made by Zhao and co-workers. The authors coupled gold nanorods to absorbing films of HgTe NCs,³⁸⁹ as shown in Figure 43a. Gold nanorods present two resonances, and the resonance associated with the long direction of the nanorods falls in the near-infrared region (see Figure 43b). Small HgTe NCs were synthesized to match this wavelength. To avoid exciton quenching in the gold, a ZnO spacer was deposited on top of the gold nanorods. By using this approach, the photocurrent is enhanced by a factor of 2. Following this work, various approaches to couple HgX NC films with plasmonic resonators have been demonstrated using a top-down approach for the fabrication.

Yifat *et al.*³⁷ used a photoconductive device consisting of interdigitated electrodes coupled to HgTe absorbing films in the MWIR range. Using e-beam lithography, the authors functionalized the substrate with an array of gold nanorods. The authors matched the resonance with the HgTe band edge and observed an increase in the photocurrent up to a factor of 3. Because the authors used gold rods as a pattern, the photocurrent was polarization dependent and observed only for light polarization along the rods. The same geometry was later applied by Tang *et al.* to HgSe,³⁹ for which the resonance matched the intraband absorption in HgSe. To avoid the polarization effect, the authors used disks instead of rods as a pattern. Chu *et al.* proposed another version of this device.^{40,337} In previous devices, transport was supported by interdigitated electrodes and was completely uncoupled from the array of resonators. Chu *et al.* combined these electrical contacts and optical resonance in the same set of electrodes; see Figure 43j. In other words, the period and size of the interdigitated electrodes were used to induce resonances. A second improvement is the introduction of a backside mirror to induce a second pass of the light through the absorbing layer. This resonator is known as the guided mode resonator (GMR) because it reshapes the incident plane wave into a steady wave propagating along the substrate within the NC film. Because of the interdigitated electrode pattern, the field enhancement was polarized, and the two resonances were used to broaden the spectral response (see Figure 43h). The design obtained $\approx 100\%$ absorption along one polarization (70% of the total signal). The other key advantage of merging plasmonic resonators with transport electrodes was the generation of gain. Because the electrode spacing was reduced, the transit time in the structure was also reduced, which generated electrical gain. Consequently, by using these GMR electrodes, the photocurrent enhancement was considerably higher than the absorption enhancement, reaching a factor of approximately 1000 (see Figure 43i). Finally, the authors demonstrated that a clear tunability of the spectral response can be attained by tuning the

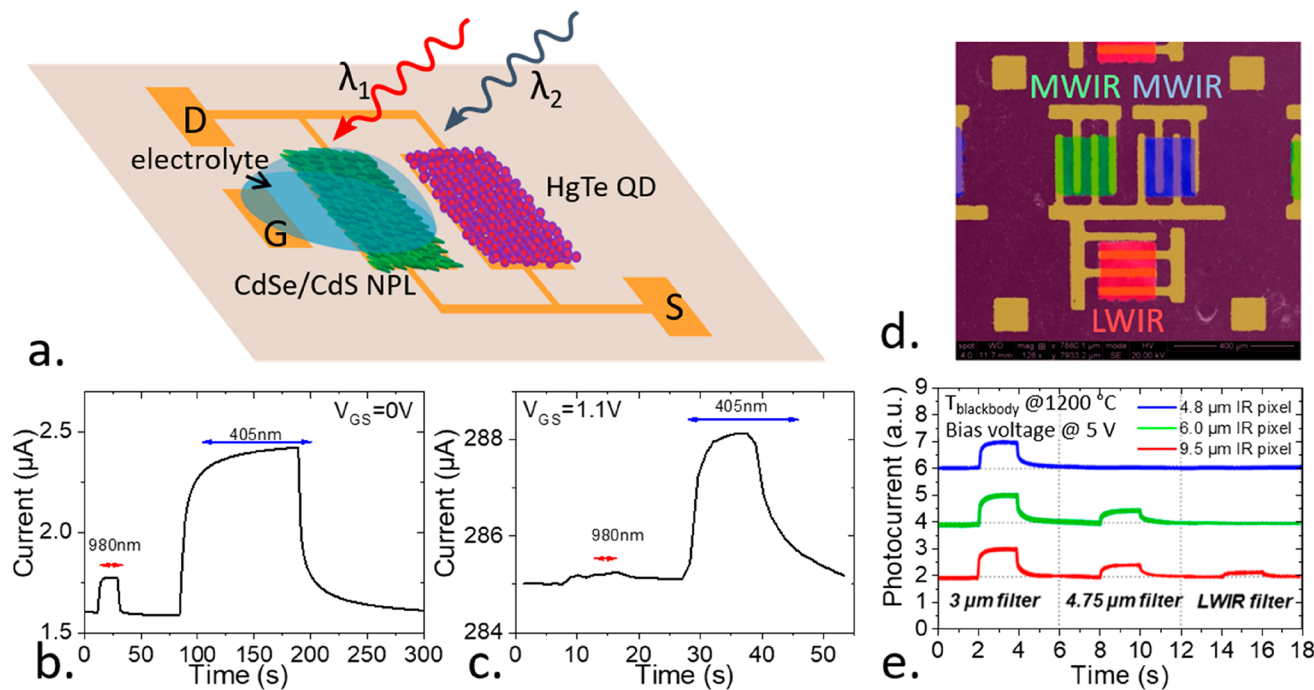


Figure 44. (a) Scheme of a dual color device. (b) (respectively c) Current as a function of time under illumination at 405 and 980 nm at an applied gate bias = 0 V (respectively 1.1 V). (a–c) Adapted with permission from ref 41. Copyright 2014 American Chemical Society. (d) Faux color SEM image of a multispectral photodetector. (e) Photocurrent as a function of time of all three pixels under pulses of illumination from a blackbody with different filters. (d,e) Adapted with permission from ref 45. Copyright 2016 American Chemical Society.

period of the grating. In this sense, this geometrical tunability, through the period, offers a new postsynthesis degree of freedom for the design of photodetectors in addition to that due to the quantum confinement in the material. Tang *et al.* proposed this method for vertical geometry devices.³⁸ In this case, the authors added an array of gold nanodisks on top of the ITO electrode (see Figure 43d) in a diode structure reported previously.³⁰³ The photocurrent enhancement reached a factor of 3–4 (see Figure 43c).

Plasmonic resonators have been successful; however, these devices introduce several losses in the metal and ITO electrodes. For example, in the GMR electrode proposed by Chu *et al.*, ≈25% of the absorption occurs in the electrodes and does not generate photocurrent. Other types of resonators have been proposed, including Fabry–Perot resonators;^{303,304} see Figure 43f. In this case, a dielectric spacing layer is required, which leads to several resonances due to the harmonic behavior; see Figure 43g. A distributed feedback mirror cavity has also been proposed by the same group. In this case, rather than the enhancement of the photocurrent, the spectral shaping of the photoresponse is targeted;⁴⁴ see Figure 45g.

In all these reports, fabrication of the resonators was performed before the deposition of the HgX nanocrystal film. Such an approach accounts for the fragility of HgX nanocrystals, which easily sinter when annealed or exposed to non solvents. Thus, most of the groups choose a strategy in which all the lithography steps are performed at the early stage of fabrication and HgX NCs are deposited as a top layer or require only minimal postdeposition processing (top metal contact). An exception was reported by Tang *et al.* last year.³⁹⁰ The grating generating the optical features was directly implemented by stamping the HgTe film.

The coupling of HgX films to resonators has been comparatively successful over recent years, and this direction

must be pursued. The losses within the metallic parts of the resonator must be addressed, likely through the introduction of dielectric resonators. Another promising direction is to supplement the introduction of a single resonance. In solar cells, up to 9 resonances can be implemented to achieve broad band absorption.³⁹¹

6.7. Multicolor Detection

Multicolor detectors can be used to combine the specificities of each spectral band to clarify the exact nature of a scene. Visible sensors have the key advantage of being broadly available and being low cost. Shortwave infrared is commonly used for active imaging or moisture detection. The mid-infrared domain corresponds to thermal imaging, which can be used for temperature determination or to conduct fully passive imaging. The combination of detectors to design bi- or multicolor detectors has already been well established.³⁹² This high level of functionality was only recently developed for NC arrays. The developments of such multicolor detectors can only be achieved owing to the progress of the single color material detectors discussed earlier.

The first category of multicolor sensors includes a combination of several single color sensors. The first bicolor sensor based on nanocrystals was proposed by Lhuillier *et al.*⁴¹ The device is a visible/SWIR detector; see the scheme of the device in Figure 44a. The SWIR detection part is made of HgTe NCs, while the visible part is made of CdSe/CdS 2D nanoplatelets.^{67,183} The films of the two materials are connected in parallel, while the switch from a SWIR to visible response is enabled by the gating of the CdSe/CdS NPLs. When no gate bias is applied, the HgTe film, due to its narrow bandgap nature, is more conductive, and the photoresponse is driven by this material: both visible and SWIR present photoconduction (see Figure 44b). When a gate bias is applied, the conductance of the NPL film increases and overcomes that of the HgTe film, owing

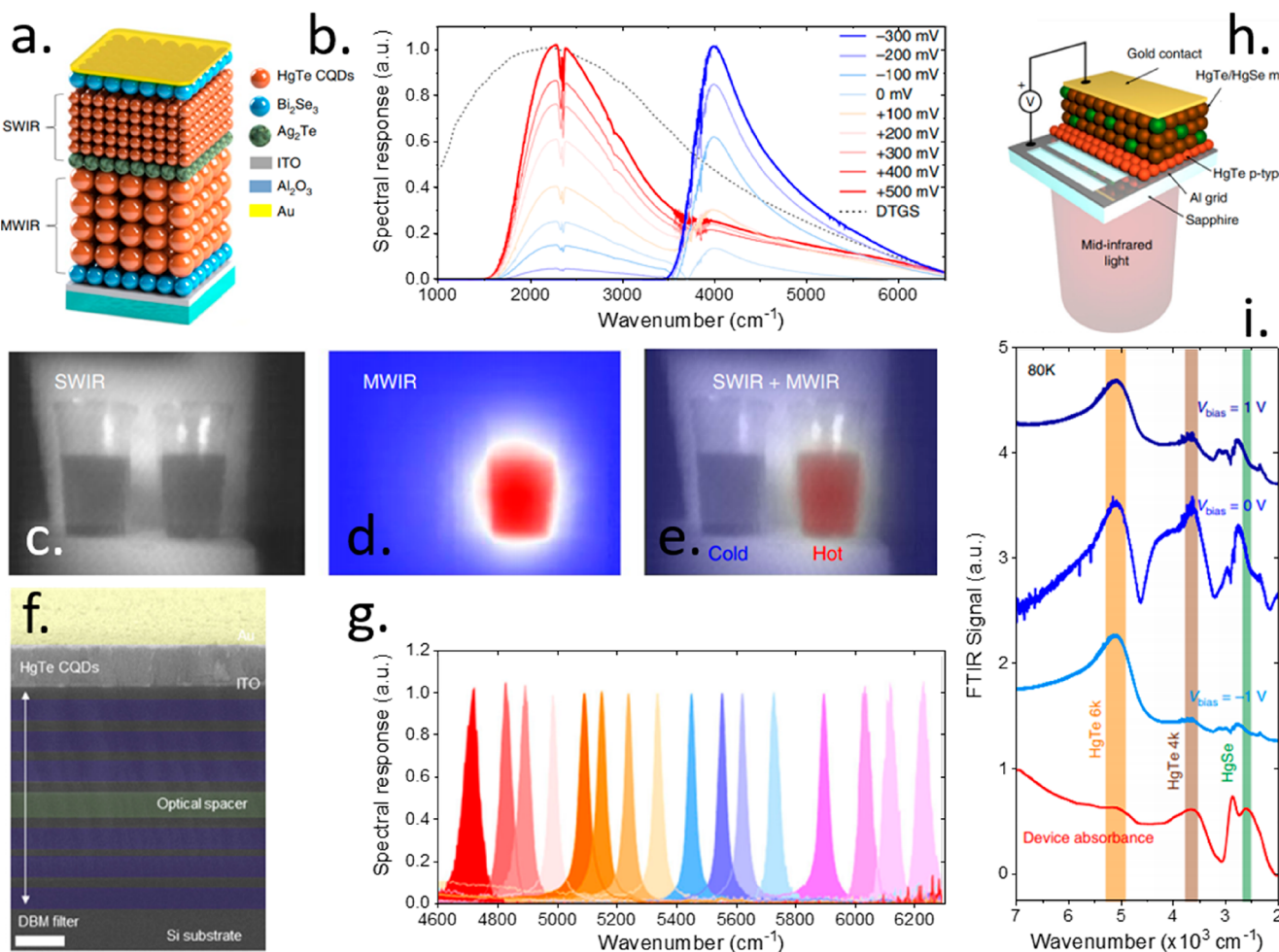


Figure 45. (a) Illustration of the structure of a dual band CQD imaging device. A bias voltage is applied between indium tin oxide (ITO) and the Au contact (grounded). (b) Spectral response of the dual band detector given in (a) under different bias voltages from positive (+500 mV) to negative (−300 mV) voltages at 85 K. The dashed black line is the response of the internal deuterated triglycine sulfate (DTGS) detector in the Fourier transform infrared (FTIR) spectrometer, which reflects the spectral irradiance of the internal FTIR blackbody. The threshold bias voltages for the MWIR photodiode dominated and SWIR photodiode dominated photocurrents were +300 mV and −200 mV, respectively. SWIR (c), MWIR (d), and merged (e) dual band images of hot and cold water. The SWIR and MWIR images are obtained at room temperature and 85 K, respectively. (a–e) Reproduced with permission from ref 46. Copyright 2019 Nature. (f) False color cross-sectional scanning electron microscopy image of the CQD hyperspectral detector. The overlapping region between ITO and Au defines the active sensing area. Scale bar: 500 nm. (g) Measured spectral responsivity of fabricated CQD hyperspectral detectors. (f,g) Reproduced with permission from ref 44. Copyright 2019 Wiley VCH. (h) Scheme of the HgTe:HgSe diode. Illumination is provided from the backside through a sapphire substrate and patterned aluminum electrode, allowing 70% of light transmission in the MWIR. (i) Photocurrent spectra and device absorbance measured at 80 K for several diode biases. The spectral contributions (maximum absorbance) of HgSe (green), HgTe 4k (bandgap at 4000 cm⁻¹, brown), and HgTe 6k (bandgap at 6000 cm⁻¹, orange) are highlighted. (h,i) Adapted with permission from ref 47. Copyright 2019 Nature.

to which only a visible response is observed (see Figure 44c). Tang *et al.*⁴⁵ proposed a three color IR sensor at 5 μm, 6 μm (MWIR), and 9.5 μm (LWIR). Fabrication of the device was based on stamping. Each active material was selectively deposited on interdigitated electrodes; see Figures 44d,e. Cryer *et al.*⁴² proposed a similar device based on four colors in the SWIR and MWIR. The selective deposition of the material was realized by combining spray coating and shadow masking on a flexible PET substrate.

A second concept to design bi- and multicolor NC-based sensors is to design a single device in which the spectral response can typically be tuned by applying a bias. Tang *et al.* proposed a SWIR/MWIR photodiode⁴⁶ consisting of a stack of an MWIR³⁰³ diode and a SWIR³⁰⁴ diode; see Figure 45a. The bias was used to switch the main contribution of the photoresponse from the SWIR to the MWIR diode; see Figure

45b. By coupling this diode to a mechanical scanning method, the authors realized bicolor imaging (see Figure 45c–e). The same group recently proposed a visible/SWIR bicolor detector.⁴³ The visible response was obtained from a Si wafer, while the SWIR response was collected from a HgTe NC film. A graphene layer was used as the intermediate transparent electrode. This device is especially promising because it can be easily transposed to a complementary metal oxide semiconductor (CMOS) readout circuit. Finally, a SWIR/MWIR sensor was proposed by Livache *et al.* by coupling the interband and intraband absorption from HgTe and HgSe, respectively; see Figure 45h,i.⁴⁷

Although the existing devices allow the switching from one spectral range to another, Tang *et al.* proposed a hyperspectral sensor. In this case, the multicolor detection was no longer driven by the material but by the optical filtering of the

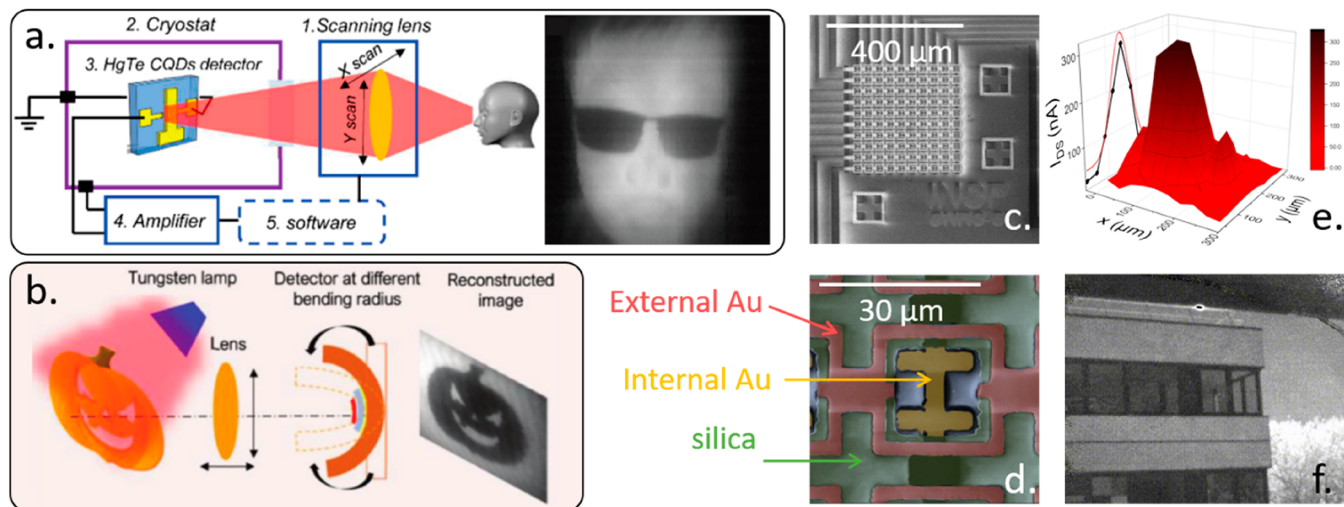


Figure 46. (a) (left) Illustration of the HgTe CQD detector thermal imaging system. (right) Thermal images captured by HgTe CQD MWIR detectors. The detector was operated at 90 K. (a) Adapted with permission from ref 38. Copyright 2018 American Chemical Society. (b) Simplified setup of infrared imaging with the flexible detector. A tungsten lamp is used to provide active infrared illumination. (b) Reproduced with permission from ref 304. Copyright 2019 Wiley VCH. (c) SEM image of the 100 pixel matrix (magnified 200 \times). (d) Zoomed SEM image of a pixel with faux colors corresponding to different layers: yellow, green, and red represent the internal gold contact (columns), insulating silica, and external gold contact (lines) (magnified 3000 \times), respectively. Blue is the glass substrate. (e) Intensity of the 97 working pixels with the laser spot focused on the matrix. The Gaussian shape of the beam has a full width at half-maximum (FWHM) of 75 μm . (c–e) Reproduced with permission from ref 146. Copyright 2019 Wilwy VCH. (f) Image from a HgTe nanocrystal film connected to CMOS ROIC, with no other lighting except for natural light (*i.e.*, passive imaging). The picture is obtained with a long-pass filter to remove the visible part of the solar spectrum. (f) Adapted from ref 50.

photoresponse. To build such a device, a HgTe layer is coupled to an array of Bragg mirrors; see Figure 45f. This resonator allowed extremely narrow band selection (approximately 30 cm^{-1}), enabling the selection of 64 bands in the SWIR and MWIR (see Figure 45g).

6.8. Integration and Imaging

Infrared nanocrystals have been discussed extensively and emerged as a possible alternative for low cost infrared imaging. The demonstration of imaging with infrared nanocrystals is relatively recent and enabled by their higher material maturity. Device integration, as opposed to single particle measurements, requires concentrated syntheses with a production of at least 100 mg (1 g is preferable even at the research level).^{29,96} Moreover, the material must present a certain degree of stability (toward air and aggregation). The demonstration of single pixel devices is a second mandatory step that allows one to understand the potential of the material compared to the existing alternatives. Alternatives to III–V and II–VI semiconductors for IR detection have been extensively studied, including colloidal materials; however, only a few recent reports presenting credible competitors with historical materials have been published. Infrared absorption is not sufficient to achieve high performance devices. Imaging devices must combine large photon-to-current conversion, fast response time (compatible with video frame rate), low noise, and material processability to obtain high quality thick films that strongly and homogeneously absorb over at least 1 cm^2 . All these reasons have delayed the demonstration of NC-based imaging setups.⁸⁶ HgX, and particularly HgTe, has reached this level of maturity and emerged as a versatile platform for infrared optoelectronics. Consequently, the coupling of HgTe films to CMOS electronics has become a recent trend in the field, stimulated by the demonstration of focal plane arrays.

The design of nanocrystal-based devices relies on non-functionalized substrates, such as glass or silicon wafers. The substrate is mostly used as a mechanical sample holder, with

possible constraints regarding the transparency. FETs are among the few early devices in which electrical functionalities from the substrate were expected. As the material maturity increased, coupling to Si electronics became possible. In the field of solar cells, it is unlikely that NCs can substitute for Si technology. The trend is generally to design tandem cells in which NCs lead to additional absorption to the silicon layer.³⁹³

A similar concept was proposed by Guyot-Sionnest and co-workers to design a bicolor detector in which a Si layer absorbing in the visible and near IR was coupled to HgTe NCs to achieve SWIR photoconduction.⁴³ Rogach and co-workers also proposed a device architecture that extended the advantage to Si coupling. The authors proposed the integration of HgTe NCs as IR absorbers in a silicon waveguide.³⁸⁸ This waveguide enhanced the incident electromagnetic field and thus obtained high absorption in a small volume. A reduced volume was expected to reduce the dark current of the device. A second benefit from this volume reduction is the reduction in the amount of HgTe NCs, which is a promising approach to obtain greener devices. By using silicon-based resonators, the advancements of Si photonics can be exploited, particularly for the design of high quality factor resonators.

Before discussing the design of multipixel sensors and focal plane arrays, it is worth noting that imaging can be obtained from a single pixel or a line of pixels coupled to a scanning system. This method has been extensively investigated by Tang *et al.*,^{38,304} as illustrated in Figure 46a,b, with application to dual color detection,⁴⁶ as shown in Figure 45c–e. Generally, the device operates under cryogenic conditions (*i.e.*, inside a cryostat), which makes displacement of the detector difficult. Therefore, the authors scanned the position of the optical lens to reconstruct an image. This method is slow due to mechanical displacement and is more suitable for picture acquisition than live imaging. Another issue is the presence of aberrations from the lens. The authors overcame this difficulty by designing a

device with enhanced light–matter coupling on a flexible substrate to obtain an eyelike sensor.³⁰⁴

Currently, the design of FPAs is a topic that attracts the most interest relative to the coupling of HgX layers to Si-based electronics. This step is critical to realize the design of low cost sensors. The cost of an IR sensor is split over three main contributions: (i) growth of the active layer (which can be significant for liquid phase epitaxy of HgCdTe epitaxially matched on a CdZnTe substrate), (ii) hybridization, in which the active layer is coupled to the readout circuit (ROIC), and (iii) the packaging (including the cooling system if needed and optics, among other aspects). To obtain a notable cost reduction for the final FPA, it is mandatory to optimize all the aforementioned aspects in addition to the simple cost difference of the colloidal and epitaxial growth. In this scenario, it is unlikely for low cost sensors to have a cryogenic operating temperature, the cost of cryocoolers remaining prohibitive. If near room temperature is required, then it would be more promising to employ SWIR materials compared to MWIR materials because their larger bandgap corresponds to a reasonably low thermally activated carrier density. NCs represent the most promising materials to reduce the hybridization cost. At present, the active layers of III–V or II–VI semiconductors are coupled to the CMOS readout circuit through indium bumps (see Figure 47a). This process presents

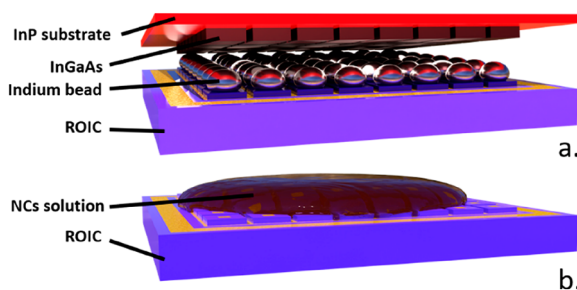


Figure 47. (a) Scheme of an InGaAs layer grown on its InP substrate and coupled to a readout circuit (ROIC) through indium bumps. (b) Scheme of an NC thin film deposited on a readout circuit (ROIC).

two severe drawbacks. First, this process is an expensive step with a limited yield, which further increases the cost. Second, this process is challenging for small pixel pitches. Existing cameras have a $15\text{ }\mu\text{m}$ pitch, and the trend is to reduce this size by approximately $10\text{ }\mu\text{m}$.³⁹⁴ This value is considerably larger than that achieved for CMOS visible sensors, for which pixel pitches of $2\text{ }\mu\text{m}$ are commonly used. Thus, the quality of the SWIR image is not diffraction limited. Using NCs, it is possible to directly spread the NC solution onto the surface of the ROIC, which eases the fabrication and provides opportunities for smaller pixel design.

Ironically, the functionalization of ROICs with HgTe NCs was first reported in the MWIR range.^{51,52} HgTe NCs with a cutoff of approximately $5\text{ }\mu\text{m}$ were spin-coated on top of the ROIC (see Figure 47b), and this step was followed by a solid-state ligand exchange procedure. Images with clear evidence of thermal contrast and NETD below 100 mK were obtained. In the SWIR, the first multipixel devices were based on small arrays of 10×10 pixels, as shown in Figure 46c,d. This matrix size was extremely small to achieve high quality images but could allow the function as an infrared beam profiler,¹⁴⁶ as shown in Figure 46e. This application is particularly well suited for nanocrystal-based FPAs, for which imaging scenes presenting a large number

of photons occur frequently because the amount of dark current is higher than that in the case of InGaAs detectors. Later, Chu *et al.* deposited an ink of HgTe NCs on a VGA ROIC and used this FPA as an infrared beam profiler, albeit with a significantly improved resolution.⁴⁹ Finally, by enhancing the quality of ink deposition, Gréboval *et al.* demonstrated passive imaging with these VGA FPAs; see the image in Figure 46f. No memory effect was observed in these devices, and framerates compatible with live imaging could be achieved.⁵⁰

7. CONCLUSIONS

Over the past few decades, significant progress has been achieved for HgX NCs and their integration in devices. For material growth, a key aspect relates to size control, which has allowed the absorption of HgX materials to increase from $3.7\text{ }\mu\text{m}$ ²⁷ in 2006 to $60\text{ }\mu\text{m}$ at present.³⁰ The other key development from NC growth is the discovery of self-doped NCs presenting intraband absorption features.^{33,65,66} The use of such transitions has been facilitated by the development of optimal experimental methods to clarify the doping mechanism and strategies to tune the corresponding magnitude. At this stage, intraband devices remain less effective than their interband competitors. However, intraband NCs are particularly interesting for the design of new devices operating at longer wavelengths and for the development of low toxicity alternatives.

The global knowledge and understanding of the material have also been considerably improved. Ten years ago, the modeling of HgX compounds was more or less limited to an effective mass approximation to determine the energy *vs* size dependence. Significant theoretical efforts based on multiband k-p,⁶² tight binding,^{28,61,216} and DFT calculations have been dedicated to the understanding of properties such as the conduction band fine structure, temperature dependence of the spectroscopic properties, or the emergence of four-level systems for stimulated emission.

In terms of light emissions, which represented the most mature aspect 10 years ago, the initial trend was to extend PL emissions toward redder wavelengths. A value of $5\text{ }\mu\text{m}$ at room temperature appears to be the current limit owing to the low PL efficiency.⁹⁵ PL magnitude reduction is intrinsic to infrared wavelengths, and the room for enhancement (*e.g.*, with heterostructures) is typically 1 order of magnitude. Stimulated emission²² and electroluminescence²³ have been demonstrated over recent years, although a laser has not been developed yet.

On the detection side, tremendous progress has been made, including mid-IR photoconduction, high mobility films, near unity absorption devices,⁴⁰ multicolor detection, demonstration of stability up to a three month time scale, and demonstration of focal plane arrays in the SWIR and MWIR ranges. It was believed that from the detection perspective, interest in HgTe would be driven by thermal imaging in the MWIR. This early belief was actually driven by the material and academic motivations to use HgTe NCs in a range of wavelengths beyond those achieved by lead chalcogenide materials. In the MWIR, the applications mostly relate to defense and astronomy, for which cost is not a major constraint and extremely high performance and reliability are desired. The SWIR domain is presently exhibiting an interest in low cost detection. This range of wavelengths suffers from the vicinity of the visible range, for which extremely high performance imaging is available with inexpensive applications. A VGA format Si-based FPA in the visible region based on CMOS technology costs less than \$1 and achieves excellent performances up to $1\text{ }\mu\text{m}$, while the InGaAs equivalent operating

at up to $1.7\text{ }\mu\text{m}$ is considerably more expensive (more than \$100). This notable difference in the cost for a marginal shift of the cutoff wavelength is typically not well understood by customers outside the detection community. Infrared NC-based FPAs exhibit considerably lower performances than those of FPAs based on InGaAs; consequently, applications for which a dark current is not a limitation should be targeted. A clear application is infrared beam profiling. Because of the large number of photons available in this application, this scenario is compatible with moderate responsivity devices. Active imaging is another application in which a dark current is not a limiting factor because the scene is illuminated. Various applications relative to industrial vision fall in this type of application, including waste sorting (SWIR is used to distinguish metal from plastic or paper), moisture detection (water strongly absorbs in the SWIR range), and food damage detection (broken eggs, fruit damage, etc.). Detection for LIDAR may also be considered owing to the short response time of HgTe devices, although this aspect is expected to be challenging because of the reduced photon flux.

8. FUTURE DEVELOPMENTS

8.1. Need for Better Materials

8.1.1. Stability. While the performance of HgX NC-based devices has been enhanced, transfer toward the mass market requires a demonstration of the viability of long-term operations. Stability is the main issue due to the corresponding performance enhancement. If this issue can be effectively addressed, then HgX NCs will enter a new phase of development. The objective is for the material to be operated without irreversible damage up to $60\text{ }^\circ\text{C}$ while being processable with limited damage up to $150\text{ }^\circ\text{C}$. Hybrid passivation combining halides and organic ligands can enhance film stability,³⁶ and multilayer encapsulation based on water and oxygen repelling layers can ensure room temperature stability.³⁰² These objectives have not been attained yet. Low temperature damage prevents the use of various encapsulation methods. Material developments must be facilitated to enable such encapsulations. The solutions will likely require the development of new core–shell growths, making large scale syntheses more complicated. However, this aspect can be realized because display applications have demonstrated that complex core–shell nanoparticles can be grown at a large scale. However, even if the scale up of HgX NC can benefit from the developments of CdSe and InP, the change in material will require a certain update of the procedure to account for the specificity of the HgX compound, such as low T growth.

8.1.2. Toxicity. Although this review is focused on HgX materials, the quest to achieve low toxicity must be considered. The current situation regarding toxicity is not as complex as that for solar cells. When CdTe-based solar cells were developed for third generation solar cells, in addition to the undeniable toxicity of Cd, the supply of a large amount of Te was under debate. There is a fundamental size difference between IR sensors and solar cell devices. A VGA format focal plane array is a 1 cm^2 square chip, while a solar cell requires m^2 devices. Livache *et al.* estimated that introducing a HgX NC-based focal plane array in every car sold in Europe (20 million unit/year) will “only” require a few tens of kg of HgX NCs.³⁹⁵

Another focus is on alternatives materials. The current industry dedicated to IR sensing is already used to managing II–VI compounds. While this industry has demonstrated interest in

the NC form, the presence of Hg still retards the industrial transfer of HgX NCs despite their reported promising performance. A heavy metal free alternative with a similar performance to that achievable using HgTe may be a breakthrough technique. Various materials have been developed to achieve IR photoconduction, including graphene, black phosphorus,^{396–400} platinum,^{401,402} and palladium^{403,404} dichalcogenides, silver chalcogenides,^{77–79,405,406} and doped oxides (ITO)s.⁷⁶ For example, Ag₂Se can easily be grown in NC form. The infrared spectrum is surprisingly similar to that of HgSe. In addition, Ag₂Se is a chalcogenide NC, and it is natural to assume that the knowledge transfer from PbX, CdS, and HgX materials can ease the integration of this material into devices. Recent demonstrations have been rather disappointing. In particular, materials exhibit low electrochemical stability, and the silver atoms undergo changes in redox states while obtaining *I*–*V* curves.⁷⁹ The performance for IR photodetection is several orders of magnitude lower than that measured for HgX NCs. Similarly, the potential of ITO NCs has been evaluated. The ITO spectrum is widely tunable in the mid-infrared region through the doping magnitude. ITO is a plasmonic material, and a higher cross-section was anticipated compared to that for intraband absorption. However, this benefit is more than balanced by the extremely short lifetime of plasmons (a few fs). Consequently, the infrared photoresponse obtained in devices made from this material is not due to photoconduction but to bolometric effects.⁷⁶

One of the challenges, while trying to develop new materials for IR sensing is to move beyond the photoconduction proof of concept. The latter aspect corresponds to a technological readiness level of 1, while a level of 3 or 4 is required for the industry to take over. In addition to photoconduction, noise and stability also must be evaluated. The next level of maturity addresses the technology transfer from a single pixel to imaging setups. This effort is generally performed in parallel to on-chip integration. To date, only PbS and HgTe NCs are sufficiently mature to be integrated into devices.

While the quest for achieving low toxicity materials has been moderately successful for the design of IR sensors, we can try to develop “greener” HgX NCs. In particular, all the environments of HgX NCs can be made greener. One step in this direction has been taken with the report of highly concentrated syntheses.⁹⁶ The authors dramatically reduced solvent and ligand consumption. A similar effort for the ligand exchange step required to make the film conductive is also required. This phenomenon may be achieved through direct synthesis of the NCs into polar solvents, as proposed by Kershaw *et al.*⁹² Film-making techniques should also be revised: the current trend is to use spin-coating deposition onto a substrate, electrodes, or readout circuit. This technique is responsible for a considerable loss of NCs, which is currently the major source of waste, generating high costs and excessive consumption of heavy metals. Better deposition techniques must be developed. Inkjet printing^{407,408} or stamping⁴⁵ may be able to be used, as only the needed amount of material must be deposited. In addition, this aspect may enable the deposition of NCs at the wafer scale rather than the chip scale. While NC deposition by these methods has been reported, one must not neglect the time required to transfer these skills to HgX NCs.

8.2. Need for Deeper Material Understanding

A clear trend for both light emission and detection is the development of light–matter coupling. In other words, the

spectral response of a detector is no longer restricted to the intrinsic optical properties of the material. The introduction of photonic structures (plasmonic resonators, distributed feedback mirrors⁴⁴) has enabled the introduction of new degrees of freedom for the design of spectral responses, PL efficiency enhancement,²¹ or light focusing.^{38,40} This direction will undoubtedly be expanded in the coming years. While the design of the light–matter coupling is strongly correlated to electromagnetic simulation tools, the amount of work dedicated to the measurement of the optical properties of HgX NCs remains comparatively low.^{114,228} Deeper and more systematic studies on the complex optical index need to be conducted to serve as inputs for simulations. One reason for this lack of data is the low availability of ellipsometers operating at wavelengths above 1.7 μm.^{409,410} Significant improvements have been achieved in terms of film quality,^{146,303} particularly in terms of film roughness, and consequently, no fundamental limitation is expected to prevent such measurements in the coming years.

The determination of the band alignment of HgX materials with other materials or transport layers is critical for the growth of colloidal heterostructures and the design of photodiodes. To date, HgX NC-based photodiode design does not have a rational basis and is dictated by empirical laws. Future designs must account for the specificity of narrow bandgap materials and, consequently, must be increasingly inspired by solar cells, for which the required bandgap (and the associated band alignment) is two to three times larger. For this aspect, electrochemistry and photoemission must be more systematically used to clarify the band alignment as a function of the size and surface chemistry. It is likely that the interaction of the absorbing layer with electron and hole transport layers can help facilitate the design process.

8.3. Need for Better Devices

As mentioned previously, IR sensing based (mostly) on HgTe NCs has seen dramatically increased performances over the past few years, and the performance in SWIR is similar to this of the existing commercial devices.⁴⁸ Although there remains room for device improvements, a breakthrough could be the realization of room temperature operation for MWIR detectors. From our perspective, the main technological challenge will be integration. None of the existing studies have reported on the integration of high performing new concepts (use of gates, plasmonic resonators) developed for single-pixel devices for FPAs. For instance, most resonators designed to achieve strongly absorbing films are not compatible with the pitch size of a FPA. In addition, in laboratory devices, illumination can be performed from both sides of the device; however, the presence of the readout circuit renders backside illumination impossible. These aspects must be considered by examining the interface between academic laboratories and industrial companies. Considering that the development cost of a dedicated ROIC is typically a million dollars, the cost required might also be significantly higher.

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Notes

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Biographies

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Audrey Chu received her Bachelor's degree from École Supérieure de Physique et de Chimie Industrielle de la ville de Paris. After an internship in Sandrine Ithurria's group, she received a Master's degree in material science and nanotechnology from Sorbonne University. She later joined the group of Emmanuel Lhuillier in 2018 to conduct her Ph.D. Her primary research interests include the design and study of light–matter coupled devices to enhance the photodetection properties of NC films.

Nicolas Goubet received a Ph.D. in 2016 from the University Pierre and Marie Curie. He has 15 years of experience in metal nanocrystal synthesis. He worked as a postdoctoral researcher with Emmanuel Lhuillier's group, developing new mercury chalcogenide synthesis techniques. He is currently an engineer assistant at the MONARIS laboratory at Sorbonne University.

Clément Livache received his Chimie ParisTech–ENSCP engineering degree in 2016. He also holds a Master's degree in materials sciences from Sorbonne University. He joined both Benoit Dubertret's group at ESPCI and Emmanuel Lhuillier's group at Sorbonne University in 2016 for a joint Ph.D. He was involved with device fabrication and characterization, with a focus on carrier dynamics and the exploitation of intraband transitions. He obtained a Ph.D. degree in 2019 and is presently working as a postdoctoral research fellow at the Los Alamos National Laboratory in the team of Victor Klimov.

Sandrine Ithurria is an ESPCI engineer and holds a Master's degree in chemistry from Sorbonne Université. She was a Ph.D. student with Benoit Dubertret at ESPCI, where she realized the first growth of 2D zinc blende nanoplatelets. She later joined Talapin's group at the University of Chicago to study nanocrystal surface chemistry and develop a new method for low temperature shell growth. Since 2012,

she has been an associate professor at ESPCI, with a research focus on 2D colloidal materials. In 2019, she was awarded an ERC starting grant.

Emmanuel Lhuillier is an ESPCI engineer and holds a Master's degree in condensed matter from Sorbonne University. To attain his Ph.D., he joined ONERA under the supervision of Emmanuel Ronsencher, where he examined the transport in a quantum well structure to be used as an infrared sensor. In the group of Philippe Guyot-Sionnest at the University of Chicago, he investigated the photoconduction properties of HgTe nanocrystals. He later joined the group of Benoit Dubertret at ESPCI, focusing on the optoelectronics of the colloidal quantum well. In 2015, he joined CNRS at Sorbonne Université at the Nanoscience Institute of Paris, where his group studies the optoelectronics of confined nanomaterials, specifically, infrared nanocrystals.

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ABBREVIATIONS

- BLIP = background limited infrared photodetection
CdX = cadmium chalcogenides
CQD = colloidal quantum dots
c-ALD = colloidal atomic layer deposition
DDT = dodecanethiol
DMF = dimethylformamide
DMSO = dimethylsulfoxide
EDT = ethanedithiol
EDX = energy dispersive X-ray spectroscopy
EL = electroluminescence
FET = field effect transistor
FPA = focal plane array
HgX = mercury chalcogenides
LWIR = longwave infrared
MEG = multi-exciton generation
MWIR = midwave infrared
NC = nanocrystal
NIR = near infrared
OLA = oleylamine
PL = photoluminescence
QCL = quantum cascade laser
QD = quantum dots
ROIC = readout integrated circuit
SEM = scanning electron microscopy
SWIR = short wave infrared
TEM = transmission electron microscopy
XPS (UPS) = X-ray (UV) photoemission spectroscopy

REFERENCES

- (1) Böberl, M.; Kovalenko, M. V.; Gamerith, S.; List, E. J. W.; Heiss, W. Inkjet-Printed Nanocrystal Photodetectors Operating up to 3 μm Wavelengths. *Adv. Mater.* **2007**, *19*, 3574–3578.

- (2) Seong, H.; Cho, K.; Kim, S. Photocurrent Characteristics of Solution-Processed HgTe Nanoparticle Thin Films under the Illumination of 1.3 μm Wavelength Light. *Semicond. Sci. Technol.* **2008**, *23*, 075011.
(3) Keuleyan, S.; Lhuillier, E.; Brajuskovic, V.; Guyot-Sionnest, P. Mid-Infrared HgTe Colloidal Quantum Dot Photodetectors. *Nat. Photonics* **2011**, *5*, 489–493.
(4) Kovalenko, M. V.; Scheele, M.; Talapin, D. V. Colloidal Nanocrystals with Molecular Metal Chalcogenide Surface Ligands. *Science* **2009**, *324*, 1417–1420.
(5) Lawson, W. D.; Nielsen, S.; Putley, E. H.; Young, A. S. Preparation and Properties of HgTe and Mixed Crystals of HgTe-CdTe. *J. Phys. Chem. Solids* **1959**, *9*, 325–329.
(6) Murray, C. B.; Norris, D. J.; Bawendi, M. G. Synthesis and Characterization of Nearly Monodisperse CdE (E = Sulfur, Selenium, Tellurium) Semiconductor Nanocrystallites. *J. Am. Chem. Soc.* **1993**, *115*, 8706–8715.
(7) Peng, X.; Manna, L.; Yang, W.; Wickham, J.; Scher, E.; Kadavanich, A.; Alivisatos, A. P. Shape Control of CdSe Nanocrystals. *Nature* **2000**, *404*, 59–61.
(8) Ithurria, S.; Tessier, M. D.; Mahler, B.; Lobo, R. P. S. M.; Dubertret, B.; Efros, A. L. Colloidal Nanoplatelets with Two-Dimensional Electronic Structure. *Nat. Mater.* **2011**, *10*, 936–941.
(9) Hines, M. A.; Guyot-Sionnest, P. Synthesis and Characterization of Strongly Luminescing ZnS-Capped CdSe Nanocrystals. *J. Phys. Chem.* **1996**, *100*, 468–471.
(10) Bourzac, K. Quantum Dots Go on Display. *Nature* **2013**, *493*, 283.
(11) Huang, Y.-M.; Singh, K. J.; Liu, A.-C.; Lin, C.-C.; Chen, Z.; Wang, K.; Lin, Y.; Liu, Z.; Wu, T.; Kuo, H.-C. Advances in Quantum-Dot-Based Displays. *Nanomaterials* **2020**, *10*, 1327.
(12) Leroy, S. *Phosphors and Quantum Dots 2015: LED Down-converters for Lighting and Displays*; Yole Développement 2015.
(13) Leroy, S. *Quantum Dots and Wide Color Gamut Display Technologies*; Yole Développement, 2017.
(14) Rogach, A. L.; Eychmüller, A.; Hickey, S. G.; Kershaw, S. V. Infrared-Emitting Colloidal Nanocrystals: Synthesis, Assembly, Spectroscopy, and Applications. *Small* **2007**, *3*, 536–557.
(15) Eychmüller, A.; Mews, A.; Weller, H. A Quantum Dot Quantum Well: CdS/HgS/CdS. *Chem. Phys. Lett.* **1993**, *208*, 59–62.
(16) Reiss, P.; Protière, M.; Li, L. Core/Shell Semiconductor Nanocrystals. *Small* **2009**, *5*, 154–168.
(17) Beberwyck, B. J.; Surendranath, Y.; Alivisatos, A. P. Cation Exchange: A Versatile Tool for Nanomaterials Synthesis. *J. Phys. Chem. C* **2013**, *117*, 19759–19770.
(18) Son, D. H.; Hughes, S. M.; Yin, Y.; Alivisatos, A. P. Cation Exchange Reactions in Ionic Nanocrystals. *Science* **2004**, *306*, 1009–1012.
(19) De Trizio, L.; Manna, L. Forging Colloidal Nanostructures via Cation Exchange Reactions. *Chem. Rev.* **2016**, *116*, 10852–10887.
(20) Rogach, A.; Kershaw, S. V.; Burt, M.; Harrison, M. T.; Kornowski, A.; Eychmüller, A.; Weller, H. Colloidally Prepared HgTe Nanocrystals with Strong Room-Temperature Infrared Luminescence. *Adv. Mater.* **1999**, *11*, 552–555.
(21) Sergeev, A. A.; Pavlov, D. V.; Kuchmizhak, A. A.; Lapine, M. V.; Yiu, W. K.; Dong, Y.; Ke, N.; Juodkazis, S.; Zhao, N.; Kershaw, S. V.; Rogach, A. L. Tailoring Spontaneous Infrared Emission of HgTe Quantum Dots with Laser-Printed Plasmonic Arrays. *Light: Sci. Appl.* **2020**, *9*, 16.
(22) Geiregat, P.; Houtepen, A. J.; Sagar, L. K.; Infante, I.; Zapata, F.; Grigel, V.; Allan, G.; Delerue, C.; Van Thourhout, D.; Hens, Z. Continuous-Wave Infrared Optical Gain and Amplified Spontaneous Emission at Ultralow Threshold by Colloidal HgTe Quantum Dots. *Nat. Mater.* **2018**, *17*, 35–42.
(23) Qu, J.; Rastogi, P.; Gréboval, C.; Lagarde, D.; Chu, A.; Dabard, C.; Khalili, A.; Cruguel, H.; Robert, C.; Xu, X. Z.; et al. Electroluminescence from HgTe Nanocrystals and Its Use for Active Imaging. *Nano Lett.* **2020**, *20*, 6185–6190.

- (24) Nag, A.; Kovalenko, M. V.; Lee, J.-S.; Liu, W.; Spokoyny, B.; Talapin, D. V. Metal-Free Inorganic Ligands for Colloidal Nanocrystals: S^{2-} , HS^- , Se^{2-} , HSe^- , Te^{2-} , HTe^- , TeS_3^{2-} , OH^- , and NH^{2-} as Surface Ligands. *J. Am. Chem. Soc.* **2011**, *133*, 10612–10620.
- (25) Fafarman, A. T.; Koh, W.; Diroll, B. T.; Kim, D. K.; Ko, D.-K.; Oh, S. J.; Ye, X.; Doan-Nguyen, V.; Crump, M. R.; Reifsnyder, D. C.; et al. Thiocyanate-Capped Nanocrystal Colloids: Vibrational Reporter of Surface Chemistry and Solution-Based Route to Enhanced Coupling in Nanocrystal Solids. *J. Am. Chem. Soc.* **2011**, *133*, 15753–15761.
- (26) Tang, J.; Kemp, K. W.; Hoogland, S.; Jeong, K. S.; Liu, H.; Levina, L.; Furukawa, M.; Wang, X.; Debnath, R.; Cha, D.; et al. Colloidal-Quantum-Dot Photovoltaics Using Atomic-Ligand Passivation. *Nat. Mater.* **2011**, *10*, 765–771.
- (27) Kovalenko, M. V.; Kaufmann, E.; Pachinger, D.; Roither, J.; Huber, M.; Stangl, J.; Hesser, G.; Schäffler, F.; Heiss, W. Colloidal HgTe Nanocrystals with Widely Tunable Narrow Band Gap Energies: From Telecommunications to Molecular Vibrations. *J. Am. Chem. Soc.* **2006**, *128*, 3516–3517.
- (28) Keuleyan, S. E.; Guyot-Sionnest, P.; Delerue, C.; Allan, G. Mercury Telluride Colloidal Quantum Dots: Electronic Structure, Size-Dependent Spectra, and Photocurrent Detection up to 12 μ m. *ACS Nano* **2014**, *8*, 8676–8682.
- (29) Lhuillier, E.; Scarafaglio, M.; Hease, P.; Nadal, B.; Aubin, H.; Xu, X. Z.; Lequeux, N.; Patriarche, G.; Ithurria, S.; Dubertret, B. Infrared Photodetection Based on Colloidal Quantum-Dot Films with High Mobility and Optical Absorption up to THz. *Nano Lett.* **2016**, *16*, 1282–1286.
- (30) Goubet, N.; Jagtap, A.; Livache, C.; Martinez, B.; Portalès, H.; Xu, X. Z.; Lobo, R. P. S. M.; Dubertret, B.; Lhuillier, E. Terahertz HgTe Nanocrystals: Beyond Confinement. *J. Am. Chem. Soc.* **2018**, *140*, 5033–5036.
- (31) Izquierdo, E.; Robin, A.; Keuleyan, S.; Lequeux, N.; Lhuillier, E.; Ithurria, S. Strongly Confined HgTe 2D Nanoplatelets as Narrow Near-Infrared Emitters. *J. Am. Chem. Soc.* **2016**, *138*, 10496–10501.
- (32) Rogach, A. L.; Harrison, M. T.; Kershaw, S. V.; Kornowski, A.; Burt, M. G.; Eychmüller, A.; Weller, H. Colloidally Prepared CdHgTe and HgTe Quantum Dots with Strong Near-Infrared Luminescence. *Phys. Status Solidi B* **2001**, *224*, 153–158.
- (33) Jeong, K. S.; Deng, Z.; Keuleyan, S.; Liu, H.; Guyot-Sionnest, P. Air-Stable n-Doped Colloidal HgS Quantum Dots. *J. Phys. Chem. Lett.* **2014**, *5*, 1139–1143.
- (34) Deng, Z.; Jeong, K. S.; Guyot-Sionnest, P. Colloidal Quantum Dots Intraband Photodetectors. *ACS Nano* **2014**, *8*, 11707–11714.
- (35) Lhuillier, E.; Keuleyan, S.; Zolotavin, P.; Guyot-Sionnest, P. Mid-Infrared HgTe/As₂S₃ Field Effect Transistors and Photodetectors. *Adv. Mater.* **2013**, *25*, 137–141.
- (36) Chen, M.; Lan, X.; Tang, X.; Wang, Y.; Hudson, M. H.; Talapin, D. V.; Guyot-Sionnest, P. High Carrier Mobility in HgTe Quantum Dot Solids Improves Mid-IR Photodetectors. *ACS Photonics* **2019**, *6*, 2358–2365.
- (37) Yifat, Y.; Ackerman, M.; Guyot-Sionnest, P. Mid-IR Colloidal Quantum Dot Detectors Enhanced by Optical Nano-Antennas. *Appl. Phys. Lett.* **2017**, *110*, 041106.
- (38) Tang, X.; Ackerman, M. M.; Guyot-Sionnest, P. Thermal Imaging with Plasmon Resonance Enhanced HgTe Colloidal Quantum Dot Photovoltaic Devices. *ACS Nano* **2018**, *12*, 7362–7370.
- (39) Tang, X.; Wu, G. fu; Lai, K. W. C. Plasmon Resonance Enhanced Colloidal HgSe Quantum Dot Filterless Narrowband Photodetectors for Mid-Wave Infrared. *J. Mater. Chem. C* **2017**, *5*, 362–369.
- (40) Chu, A.; Gréboval, C.; Goubet, N.; Martinez, B.; Livache, C.; Qu, J.; Rastogi, P.; Bresciani, F. A.; Prado, Y.; Suffit, S.; et al. Near Unity Absorption in Nanocrystal Based Short Wave Infrared Photodetectors Using Guided Mode Resonators. *ACS Photonics* **2019**, *6*, 2553–2561.
- (41) Lhuillier, E.; Robin, A.; Ithurria, S.; Aubin, H.; Dubertret, B. Electrolyte-Gated Colloidal Nanoplatelets-Based Phototransistor and Its Use for Bicolor Detection. *Nano Lett.* **2014**, *14*, 2715–2719.
- (42) Cryer, M. E.; Halpert, J. E. 300 nm Spectral Resolution in the Mid-Infrared with Robust, High Responsivity Flexible Colloidal Quantum Dot Devices at Room Temperature. *ACS Photonics* **2018**, *5*, 3009–3015.
- (43) Tang, X.; Chen, M.; Kamath, A.; Ackerman, M. M.; Guyot-Sionnest, P. Colloidal Quantum-Dots/Graphene/Silicon Dual-Channel Detection of Visible Light and Short-Wave Infrared. *ACS Photonics* **2020**, *7*, 1117–1121.
- (44) Tang, X.; Ackerman, M. M.; Guyot-Sionnest, P. Acquisition of Hyperspectral Data with Colloidal Quantum Dots. *Laser Photonics Rev.* **2019**, *13*, 1900165.
- (45) Tang, X.; Tang, X.; Lai, K. W. C. Scalable Fabrication of Infrared Detectors with Multispectral Photoresponse Based on Patterned Colloidal Quantum Dot Films. *ACS Photonics* **2016**, *3*, 2396–2404.
- (46) Tang, X.; Ackerman, M. M.; Chen, M.; Guyot-Sionnest, P. Dual-Band Infrared Imaging Using Stacked Colloidal Quantum Dot Photodiodes. *Nat. Photonics* **2019**, *13*, 277–282.
- (47) Livache, C.; Martinez, B.; Goubet, N.; Gréboval, C.; Qu, J.; Chu, A.; Royer, S.; Ithurria, S.; Silly, M. G.; Dubertret, B.; Lhuillier, E. A Colloidal Quantum Dot Infrared Photodetector and Its Use for Intraband Detection. *Nat. Commun.* **2019**, *10*, 2125.
- (48) Ackerman, M. M.; Chen, M.; Guyot-Sionnest, P. HgTe Colloidal Quantum Dot Photodiodes for Extended Short-Wave Infrared Detection. *Appl. Phys. Lett.* **2020**, *116*, 083502.
- (49) Chu, A.; Martinez, B.; Ferré, S.; Noguier, V.; Gréboval, C.; Livache, C.; Qu, J.; Prado, Y.; Casaretto, N.; Goubet, N.; et al. HgTe Nanocrystals for SWIR Detection and Their Integration up to the Focal Plane Array. *ACS Appl. Mater. Interfaces* **2019**, *11*, 33116–33123.
- (50) Greboval, C.; Ferre, S.; Noguier, V.; Chu, A.; Qu, J.; Chee, S.-S.; Vincent, G.; Lhuillier, E. Infrared Narrow Band Gap Nanocrystals: Recent Progresses Relative to Imaging and Active Detection. *arXiv* **2020**, 2001.11554.
- (51) Buurma, C.; Pimpinella, R. E.; Ciani, A. J.; Feldman, J. S.; Grein, C. H.; Guyot-Sionnest, P. MWIR Imaging with Low Cost Colloidal Quantum Dot Films. In *Optical Sensing, Imaging, and Photon Counting: Nanostructured Devices and Applications 2016*; International Society for Optics and Photonics, 2016; Vol. 9933, p 993303.
- (52) Ciani, A. J.; Pimpinella, R. E.; Grein, C. H.; Guyot-Sionnest, P. Colloidal Quantum Dots for Low-Cost MWIR Imaging. In *Infrared Technology and Applications XLII*; International Society for Optics and Photonics, 2016; Vol. 9819, p 981919.
- (53) Rogalski, A.; Kopytko, M.; Martyniuk, P. Comparison of Performance Limits of HOT HgCdTe Photodiodes and Colloidal Quantum Dot Infrared Detectors. In *Infrared Technology and Applications XLVI*; International Society for Optics and Photonics, 2020; Vol. 11407, p 114070V.
- (54) Green, M.; Mirzai, H. Synthetic Routes to Mercury Chalcogenide Quantum Dots. *J. Mater. Chem. C* **2018**, *6*, S097–S112.
- (55) Kershaw, S. V.; Rogach, A. L. Infrared Emitting HgTe Quantum Dots and Their Waveguide and Optoelectronic Devices. *Z. Phys. Chem.* **2015**, *229*, 23–64.
- (56) Lhuillier, E.; Keuleyan, S.; Liu, H.; Guyot-Sionnest, P. Mid-IR Colloidal Nanocrystals. *Chem. Mater.* **2013**, *25*, 1272–1282.
- (57) Lhuillier, E.; Guyot-Sionnest, P. Recent Progresses in Mid Infrared Nanocrystal Optoelectronics. *IEEE J. Sel. Top. Quantum Electron.* **2017**, *23*, 6000208.
- (58) Guyot-Sionnest, P.; Ackerman, M. M.; Tang, X. Colloidal Quantum Dots for Infrared Detection beyond Silicon. *J. Chem. Phys.* **2019**, *151*, 060901.
- (59) Chen, H.; Li, L.; Cui, S.; Mahounga, D.; Zhang, J.; Gu, Y. Folate Conjugated CdHgTe Quantum Dots with High Targeting Affinity and Sensitivity for In Vivo Early Tumor Diagnosis. *J. Fluoresc.* **2011**, *21*, 793–801.
- (60) Schimpf, A. M.; Knowles, K. E.; Carroll, G. M.; Gamelin, D. R. Electronic Doping and Redox-Potential Tuning in Colloidal Semiconductor Nanocrystals. *Acc. Chem. Res.* **2015**, *48*, 1929–1937.
- (61) Hudson, M. H.; Chen, M.; Kamysbayev, V.; Janke, E. M.; Lan, X.; Allan, G.; Delerue, C.; Lee, B.; Guyot-Sionnest, P.; Talapin, D. V. Conduction Band Fine Structure in Colloidal HgTe Quantum Dots. *ACS Nano* **2018**, *12*, 9397–9404.

- (62) Moghaddam, N.; Gréboval, C.; Qu, J.; Chu, A.; Rastogi, P.; Livache, C.; Khalili, A.; Xu, X. Z.; Baptiste, B.; Klotz, S.; et al. The Strong Confinement Regime in HgTe Two-Dimensional Nanoplatelets. *J. Phys. Chem. C* **2020**, *124*, 23460–23468.
- (63) Shen, G.; Chen, M.; Guyot-Sionnest, P. Synthesis of Non-aggregating HgTe Colloidal Quantum Dots and the Emergence of Air-Stable n-Doping. *J. Phys. Chem. Lett.* **2017**, *8*, 2224–2228.
- (64) Zhang, H.; Guyot-Sionnest, P. Shape-Controlled HgTe Colloidal Quantum Dots and Reduced Spin–Orbit Splitting in the Tetrahedral Shape. *J. Phys. Chem. Lett.* **2020**, *11*, 6860–6866.
- (65) Kim, J.; Choi, D.; Jeong, K. S. Self-Doped Colloidal Semiconductor Nanocrystals with Intraband Transitions in Steady State. *Chem. Commun.* **2018**, *54*, 8435–8445.
- (66) Jagtap, A.; Livache, C.; Martinez, B.; Qu, J.; Chu, A.; Gréboval, C.; Goubet, N.; Lhuillier, E. Emergence of Intraband Transitions in Colloidal Nanocrystals. *Opt. Mater. Express* **2018**, *8*, 1174–1183.
- (67) Nasilowski, M.; Mahler, B.; Lhuillier, E.; Ithurria, S.; Dubertret, B. Two-Dimensional Colloidal Nanocrystals. *Chem. Rev.* **2016**, *116*, 10934–10982.
- (68) Chen, M.; Guyot-Sionnest, P. Reversible Electrochemistry of Mercury Chalcogenide Colloidal Quantum Dot Films. *ACS Nano* **2017**, *11*, 4165–4173.
- (69) Rogalski, A. HgCdTe Infrared Detector Material: History, Status and Outlook. *Rep. Prog. Phys.* **2005**, *68*, 2267–2336.
- (70) Svane, A.; Christensen, N. E.; Cardona, M.; Chantis, A. N.; van Schilfgaarde, M.; Kotani, T. Quasiparticle Band Structures of β -HgS, HgSe, and HgTe. *Phys. Rev. B: Condens. Matter Mater. Phys.* **2011**, *84*, 205205.
- (71) Delin, A.; Klüner, T. Excitation Spectra and Ground-State Properties from Density-Functional Theory for the Inverted Band-Structure Systems β -HgS, HgSe, and HgTe. *Phys. Rev. B: Condens. Matter Mater. Phys.* **2002**, *66*, 035117.
- (72) Man, P.; Pan, D. S. Infrared Absorption in HgTe. *Phys. Rev. B: Condens. Matter Mater. Phys.* **1991**, *44*, 8745–8758.
- (73) Tennant, W. E. Rule 07” Revisited: Still a Good Heuristic Predictor of *p/n* HgCdTe Photodiode Performance? *J. Electron. Mater.* **2010**, *39*, 1030–1035.
- (74) Tennant, W. E.; Lee, D.; Zandian, M.; Piquette, E.; Carmody, M. MBE HgCdTe Technology: A Very General Solution to IR Detection, Described by “Rule 07”, a Very Convenient Heuristic. *J. Electron. Mater.* **2008**, *37*, 1406–1410.
- (75) Garcia, G.; Buonsanti, R.; Llordes, A.; Runnerstrom, E. L.; Bergerud, A.; Milliron, D. J. Near-Infrared Spectrally Selective Plasmonic Electrochromic Thin Films. *Adv. Opt. Mater.* **2013**, *1*, 215–220.
- (76) Qu, J.; Livache, C.; Martinez, B.; Gréboval, C.; Chu, A.; Meriggio, E.; Ramade, J.; Cruguel, H.; Xu, X. Z.; Proust, A.; et al. Transport in ITO Nanocrystals with Short- to Long-Wave Infrared Absorption for Heavy-Metal-Free Infrared Photodetection. *ACS Appl. Nano Mater.* **2019**, *2*, 1621–1630.
- (77) Hafiz, S. B.; Scimeca, M. R.; Zhao, P.; Paredes, I. J.; Sahu, A.; Ko, D.-K. Silver Selenide Colloidal Quantum Dots for Mid-Wavelength Infrared Photodetection. *ACS Appl. Nano Mater.* **2019**, *2*, 1631–1636.
- (78) Park, M.; Choi, D.; Choi, Y.; Shin, H.; Jeong, K. S. Mid-Infrared Intraband Transition of Metal Excess Colloidal Ag₂Se Nanocrystals. *ACS Photonics* **2018**, *5*, 1907–1911.
- (79) Qu, J.; Goubet, N.; Livache, C.; Martinez, B.; Amelot, D.; Gréboval, C.; Chu, A.; Ramade, J.; Cruguel, H.; Ithurria, S.; et al. Intraband Mid-Infrared Transitions in Ag₂Se Nanocrystals: Potential and Limitations for Hg-Free Low-Cost Photodetection. *J. Phys. Chem. C* **2018**, *122*, 18161–18167.
- (80) Liu, W.; Chang, A. Y.; Schaller, R. D.; Talapin, D. V. Colloidal InSb Nanocrystals. *J. Am. Chem. Soc.* **2012**, *134*, 20258–20261.
- (81) Crisp, R. W.; Grimaldi, G.; De Trizio, L. D.; Evers, W. H.; Kirkwood, N.; Kinge, S.; Manna, L.; Siebbeles, L. D. A.; Houtepen, A. J. Selective Antimony Reduction Initiating the Nucleation and Growth of InSb Quantum Dots. *Nanoscale* **2018**, *10*, 11110–11116.
- (82) Yarema, M.; Kovalenko, M. V. Colloidal Synthesis of InSb Nanocrystals with Controlled Polymorphism Using Indium and Antimony Amides. *Chem. Mater.* **2013**, *25*, 1788–1792.
- (83) Hines, M. A.; Scholes, G. D. Colloidal PbS Nanocrystals with Size-Tunable Near-Infrared Emission: Observation of Post-Synthesis Self-Narrowing of the Particle Size Distribution. *Adv. Mater.* **2003**, *15*, 1844–1849.
- (84) Weidman, M. C.; Beck, M. E.; Hoffman, R. S.; Prins, F.; Tisdale, W. A. Monodisperse, Air-Stable PbS Nanocrystals via Precursor Stoichiometry Control. *ACS Nano* **2014**, *8*, 6363–6371.
- (85) Shulga, A. G.; Pivotteau, L.; Bisri, S. Z.; Kovalenko, M. V.; Loi, M. A. Double Gate PbS Quantum Dot Field-Effect Transistors for Tuneable Electrical Characteristics. *Adv. Electron. Mater.* **2016**, *2*, 1500467.
- (86) Rauch, T.; Böberl, M.; Tedde, S. F.; Fürst, J.; Kovalenko, M. V.; Hesser, G.; Lemmer, U.; Heiss, W.; Hayden, O. Near-Infrared Imaging with Quantum-Dot-Sensitized Organic Photodiodes. *Nat. Photonics* **2009**, *3*, 332–336.
- (87) Konstantatos, G.; Howard, I.; Fischer, A.; Hoogland, S.; Clifford, J.; Klem, E.; Levina, L.; Sargent, E. H. Ultrasensitive Solution-Cast Quantum Dot Photodetectors. *Nature* **2006**, *442*, 180–183.
- (88) McDonald, S. A.; Konstantatos, G.; Zhang, S.; Cyr, P. W.; Klem, E. J. D.; Levina, L.; Sargent, E. H. Solution-Processed PbS Quantum Dot Infrared Photodetectors and Photovoltaics. *Nat. Mater.* **2005**, *4*, 138–142.
- (89) Özdemir, O.; Ramiro, I.; Gupta, S.; Konstantatos, G. High Sensitivity Hybrid PbS CQD-TMD Photodetectors up to 2 μm . *ACS Photonics* **2019**, *6*, 2381–2386.
- (90) Killilea, N.; Wu, M.; Sytnyk, M.; Yousefi Amin, A. A.; Mashkov, O.; Spiecker, E.; Heiss, W. Pushing PbS/Metal-Halide-Perovskite Core/Epitaxial-Ligand-Shell Nanocrystal Photodetectors beyond 3 μm Wavelength. *Adv. Funct. Mater.* **2019**, *29*, 1807964.
- (91) Günes, S.; Neugebauer, H.; Sariciftci, N. S.; Roither, J.; Kovalenko, M.; Pillwein, G.; Heiss, W. Hybrid Solar Cells Using HgTe Nanocrystals and Nanoporous TiO₂ Electrodes. *Adv. Funct. Mater.* **2006**, *16*, 1095–1099.
- (92) Kershaw, S. V.; Yiu, W. K.; Sergeev, A.; Rogach, A. L. Development of Synthetic Methods to Grow Long-Wavelength Infrared-Emitting HgTe Quantum Dots in Dimethylformamide. *Chem. Mater.* **2020**, *32*, 3930–3943.
- (93) Green, M.; Wakefield, G.; Dobson, P. J. A Simple Metalorganic Route to Organically Passivated Mercury Telluride Nanocrystals. *J. Mater. Chem.* **2003**, *13*, 1076–1078.
- (94) Piepenbrock, M.-O. M.; Stirner, T.; Kelly, S. M.; O'Neill, M. A Low-Temperature Synthesis for Organically Soluble HgTe Nanocrystals Exhibiting Near-Infrared Photoluminescence and Quantum Confinement. *J. Am. Chem. Soc.* **2006**, *128*, 7087–7090.
- (95) Keuleyan, S.; Lhuillier, E.; Guyot-Sionnest, P. Synthesis of Colloidal HgTe Quantum Dots for Narrow Mid-IR Emission and Detection. *J. Am. Chem. Soc.* **2011**, *133*, 16422–16424.
- (96) Goubet, N.; Thomas, M.; Gréboval, C.; Chu, A.; Qu, J.; Rastogi, P.; Chee, S.-S.; Goyal, M.; Zhang, Y.; Xu, X. Z.; et al. Near- to Long-Wave-Infrared Mercury Chalcogenide Nanocrystals from Liquid Mercury. *J. Phys. Chem. C* **2020**, *124*, 8423–8430.
- (97) Müllenborn, M.; Jarvis, R. F.; Yacobi, B. G.; Kaner, R. B.; Coleman, C. C.; Haegel, N. M. Characterization of Solution-Synthesized CdTe and HgTe. *Appl. Phys. A: Mater. Sci. Process.* **1993**, *56*, 317–321.
- (98) Rogach, A. L.; Katsikas, L.; Kornowski, A.; Su, D.; Eychmüller, A.; Weller, H. Synthesis and Characterization of Thiol-Stabilized CdTe Nanocrystals. *Ber. Bunsenges. Phys. Chem.* **1996**, *100*, 1772–1778.
- (99) Gao, M.; Kirstein, S.; Möhwald, H.; Rogach, A. L.; Kornowski, A.; Eychmüller, A.; Weller, H. Strongly Photoluminescent CdTe Nanocrystals by Proper Surface Modification. *J. Phys. Chem. B* **1998**, *102*, 8360–8363.
- (100) Harrison, M. T.; Kershaw, S. V.; Burt, M. G.; Rogach, A.; Eychmüller, A.; Weller, H. Investigation of Factors Affecting the Photoluminescence of Colloidally-Prepared HgTe Nanocrystals. *J. Mater. Chem.* **1999**, *9*, 2721–2722.

- (101) Abdelazim, N. M.; Zhu, Q.; Xiong, Y.; Zhu, Y.; Chen, M.; Zhao, N.; Kershaw, S. V.; Rogach, A. L. Room Temperature Synthesis of HgTe Quantum Dots in an Aprotic Solvent Realizing High Photoluminescence Quantum Yields in the Infrared. *Chem. Mater.* **2017**, *29*, 7859–7867.
- (102) Wang, Y.; Liu, Z.; Huo, N.; Li, F.; Gu, M.; Ling, X.; Zhang, Y.; Lu, K.; Han, L.; Fang, H.; et al. Room-Temperature Direct Synthesis of Semi-Conductive PbS Nanocrystal Inks for Optoelectronic Applications. *Nat. Commun.* **2019**, *10*, 5136.
- (103) Kim, H.; Park, B.; Cho, K.; Kim, J.-H.; Lee, J. W.; Kim, D.-W.; Kim, S. Transport of Charge Carriers in HgTe/CdTe Core-Shell Nanoparticle Film. *Jpn. J. Appl. Phys.* **2005**, *44*, 5703.
- (104) Endicott, K. The Trembling Edge of Science. *Dartmouth Alumni Magazine* **1998**, 22–31.
- (105) Green, M. Production of Metal Chalcogenide Nanoparticles. US Patent 20040086444A1, May 6, 2004.
- (106) Kim, S.; Kim, T.; Im, S. H.; Seok, S. I.; Kim, K. W.; Kim, S.; Kim, S.-W. Bandgap Engineered Monodisperse and Stable Mercury Telluride Quantum Dots and Their Application for Near-Infrared Photodetection. *J. Mater. Chem.* **2011**, *21*, 15232–15236.
- (107) Li, L. S.; Wang, H.; Liu, Y.; Lou, S.; Wang, Y.; Du, Z. Room Temperature Synthesis of HgTe Nanocrystals. *J. Colloid Interface Sci.* **2007**, *308*, 254–257.
- (108) Noumbé, U. N.; Gréboval, C.; Livache, C.; Chu, A.; Majjad, H.; Parra López, L. E.; Mouafo, L. D. N.; Doudin, B.; Berciaud, S.; Chaste, J.; et al. Reconfigurable 2D/0D p–n Graphene/HgTe Nanocrystal Heterostructure for Infrared Detection. *ACS Nano* **2020**, *14*, 4567–4576.
- (109) Grigel, V.; Sagar, L. K.; De Nolf, K.; Zhao, Q.; Vantomme, A.; De Roo, J.; Infante, I.; Hens, Z. The Surface Chemistry of Colloidal HgSe Nanocrystals, toward Stoichiometric Quantum Dots by Design. *Chem. Mater.* **2018**, *30*, 7637–7647.
- (110) Pan, Y.; Bai, H.; Pan, L.; Li, Y.; Tamargo, M. C.; Sohel, M.; Lombardi, J. R. Size Controlled Synthesis of Monodisperse PbTe Quantum Dots: Using Oleylamine as the Capping Ligand. *J. Mater. Chem.* **2012**, *22*, 23593–23601.
- (111) Wehrenberg, B. L.; Wang, C.; Guyot-Sionnest, P. Interband and Intraband Optical Studies of PbSe Colloidal Quantum Dots. *J. Phys. Chem. B* **2002**, *106*, 10634–10640.
- (112) Rinnerbauer, V.; Hingerl, K.; Kovalenko, M.; Heiss, W. Effect of Quantum Confinement on Higher Transitions in HgTe Nanocrystals. *Appl. Phys. Lett.* **2006**, *89*, 193114.
- (113) Jean, J.; Xiao, J.; Nick, R.; Moody, N.; Nasilowski, M.; Bawendi, M.; Bulović, V. Synthesis Cost Dictates the Commercial Viability of Lead Sulfide and Perovskite Quantum Dot Photovoltaics. *Energy Environ. Sci.* **2018**, *11*, 2295–2305.
- (114) Lhuillier, E.; Keuleyan, S.; Guyot-Sionnest, P. Optical Properties of HgTe Colloidal Quantum Dots. *Nanotechnology* **2012**, *23*, 175705.
- (115) Harrison, M. T.; Kershaw, S. V.; Burt, M. G.; Rogach, A. L.; Kornowski, A.; Eychmüller, A.; Weller, H. Colloidal Nanocrystals for Telecommunications. Complete Coverage of the Low-Loss Fiber Windows by Mercury Telluride Quantum Dot. *Pure Appl. Chem.* **2000**, *72*, 295–307.
- (116) Kuno, M.; Higginson, K. A.; Qadri, S. B.; Yousuf, M.; Lee, S. H.; Davis, B. L.; Matoussi, H. Molecular Clusters of Binary and Ternary Mercury Chalcogenides: Colloidal Synthesis, Characterization, and Optical Spectra. *J. Phys. Chem. B* **2003**, *107*, 5758–5767.
- (117) Howes, P.; Green, M.; Johnston, C.; Crossley, A. Synthesis and Shape Control of Mercury Selenide (HgSe) Quantum Dots. *J. Mater. Chem.* **2008**, *18*, 3474–3480.
- (118) Jeong, J.; Yoon, B.; Kwon, Y.-W.; Choi, D.; Jeong, K. S. Singly and Doubly Occupied Higher Quantum States in Nanocrystals. *Nano Lett.* **2017**, *17*, 1187–1193.
- (119) Choi, D.; Park, M.; Jeong, J.; Shin, H.-B.; Choi, Y. C.; Jeong, K. S. Multifunctional Self-Doped Nanocrystal Thin-Film Transistor Sensors. *ACS Appl. Mater. Interfaces* **2019**, *11*, 7242–7249.
- (120) Goubet, N.; Livache, C.; Martinez, B.; Xu, X. Z.; Ithurria, S.; Royer, S.; Cruguel, H.; Patriarche, G.; Ouerghi, A.; Silly, M.; et al. Wave-Function Engineering in HgSe/HgTe Colloidal Heterostructures To Enhance Mid-Infrared Photoconductive Properties. *Nano Lett.* **2018**, *18*, 4590–4597.
- (121) Chen, M.; Shen, G.; Guyot-Sionnest, P. Size Distribution Effects on Mobility and Intraband Gap of HgSe Quantum Dots. *J. Phys. Chem. C* **2020**, *124*, 16216–16221.
- (122) Martinez, B.; Livache, C.; Notemgnou Mouafo, L. D.; Goubet, N.; Keuleyan, S.; Cruguel, H.; Ithurria, S.; Aubin, H.; Ouerghi, A.; Doudin, B.; et al. HgSe Self-Doped Nanocrystals as a Platform to Investigate the Effects of Vanishing Confinement. *ACS Appl. Mater. Interfaces* **2017**, *9*, 36173–36180.
- (123) Gonzales, M. A.; Han, H.; Moyes, A.; Radinos, A.; Hobbs, A. J.; Coombs, N.; Oliver, S. R. J.; Mascharak, P. K. Light-Triggered Carbon Monoxide Delivery with Al-MCM-41-Based Nanoparticles Bearing a Designed Manganese Carbonyl Complex. *J. Mater. Chem. B* **2014**, *2*, 2107–2113.
- (124) Choi, D.; Yoon, B.; Kim, D.-K.; Baik, H.; Choi, J.-H.; Jeong, K. S. Major Electronic Transition Shift from Bandgap to Localized Surface Plasmon Resonance in Cd_xHg_{1-x}Se Alloy Nanocrystals. *Chem. Mater.* **2017**, *29*, 8548–8554.
- (125) Schooss, D.; Mews, A.; Eychmüller, A.; Weller, H. Quantum-Dot Quantum Well CdS/HgS/CdS: Theory and Experiment. *Phys. Rev. B: Condens. Matter Mater. Phys.* **1994**, *49*, 17072–17078.
- (126) Kamalov, V. F.; Little, R.; Logunov, S. L.; El-Sayed, M. A. Picosecond Electronic Relaxation in CdS/HgS/CdS Quantum Dot Quantum Well Semiconductor Nanoparticles. *J. Phys. Chem.* **1996**, *100*, 6381–6384.
- (127) Mews, A.; Kadavanich, A. V.; Banin, U.; Alivisatos, A. P. Structural and Spectroscopic Investigations of CdS/HgS/CdS Quantum-Dot Quantum Wells. *Phys. Rev. B: Condens. Matter Mater. Phys.* **1996**, *53*, R13242–R13245.
- (128) Dorfs, D.; Henschel, H.; Kolny, J.; Eychmüller, A. Multilayered Nanoheterostructures: Theory and Experiment. *J. Phys. Chem. B* **2004**, *108*, 1578–1583.
- (129) Higginson, K. A.; Kuno, M.; Bonevich, J.; Qadri, S. B.; Yousuf, M.; Matoussi, H. Synthesis and Characterization of Colloidal β-HgS Quantum Dots. *J. Phys. Chem. B* **2002**, *106*, 9982–9985.
- (130) Yang, Y.-G.; Liu, H.-G.; Chen, L.-J.; Chen, K.-C.; Ding, H.-P.; Hao, J. One-Step Synthesis and Assembly of Two-Dimensional Arrays of Mercury Sulfide Nanocrystal Aggregates at the Air/Water Interface. *Langmuir* **2010**, *26*, 14879–14884.
- (131) Wichaiansee, W.; Nordin, M. N.; Green, M.; Curry, R. J. Synthesis and Optical Characterization of Infra-Red Emitting Mercury Sulfide (HgS) Quantum Dots. *J. Mater. Chem.* **2011**, *21*, 7331–7336.
- (132) Yang, J.; Zhang, W.-H.; Hu, Y.-P.; Yu, J.-S. Aqueous Synthesis and Characterization of Glutathione-Stabilized β-HgS Nanocrystals with near-Infrared Photoluminescence. *J. Colloid Interface Sci.* **2012**, *379*, 8–13.
- (133) Zhang, L.; Yang, G.; He, G.; Wang, L.; Liu, Q.; Zhang, Q.; Qin, D. Synthesis of HgS Nanocrystals in the Lysozyme Aqueous Solution through Biomimetic Method. *Appl. Surf. Sci.* **2012**, *258*, 8185–8191.
- (134) Goswami, N.; Giri, A.; Kar, S.; Bootharaju, M. S.; John, R.; Xavier, P. L.; Pradeep, T.; Pal, S. K. Protein-Directed Synthesis of NIR-Emitting, Tunable HgS Quantum Dots and Their Applications in Metal-Ion Sensing. *Small* **2012**, *8*, 3175–3184.
- (135) Han, L.; Hou, P.; Feng, Y.; Liu, H.; Li, J.; Peng, Z.; Yang, J. Phase Transfer-Based Synthesis of HgS Nanocrystals. *Dalton Trans.* **2014**, *43*, 11981–11987.
- (136) Shen, G.; Guyot-Sionnest, P. HgS and HgS/CdS Colloidal Quantum Dots with Infrared Intraband Transitions and Emergence of a Surface Plasmon. *J. Phys. Chem. C* **2016**, *120*, 11744–11753.
- (137) Mews, A.; Eychmüller, A.; Giersig, M.; Schooss, D.; Weller, H. Preparation, characterization, and photophysics of the quantum dot quantum well system cadmium sulfide/mercury sulfide/cadmium sulfide. *J. Phys. Chem.* **1994**, *98*, 934–941.
- (138) Ben-Moshe, A.; Govorov, A. O.; Markovich, G. Enantioselective Synthesis of Intrinsically Chiral Mercury Sulfide Nanocrystals. *Angew. Chem.* **2013**, *125*, 1313–1317.

- (139) Wang, P.; Yu, S.-J.; Govorov, A. O.; Ouyang, M. Cooperative Expression of Atomic Chirality in Inorganic Nanostructures. *Nat. Commun.* **2017**, *8*, 14312.
- (140) Kuno, J.; Immura, Y.; Katouda, M.; Tashiro, M.; Kawai, T.; Nakashima, T. Inversion of Optical Activity in the Synthesis of Mercury Sulfide Nanoparticles: Role of Ligand Coordination. *Angew. Chem., Int. Ed.* **2018**, *57*, 12022–12026.
- (141) Kuno, J.; Kawai, T.; Nakashima, T. The Effect of Surface Ligands on the Optical Activity of Mercury Sulfide Nanoparticles. *Nanoscale* **2017**, *9*, 11590–11595.
- (142) Wang, P.; Yu, S.-J.; Ouyang, M. Assembled Suprastructures of Inorganic Chiral Nanocrystals and Hierarchical Chirality. *J. Am. Chem. Soc.* **2017**, *139*, 6070–6073.
- (143) Manna, L.; Milliron, D. J.; Meisel, A.; Scher, E. C.; Alivisatos, A. P. Controlled Growth of Tetrapod-Branched Inorganic Nanocrystals. *Nat. Mater.* **2003**, *2*, 382–385.
- (144) Ithurria, S.; Dubertret, B. Quasi 2D Colloidal CdSe Platelets with Thicknesses Controlled at the Atomic Level. *J. Am. Chem. Soc.* **2008**, *130*, 16504–16505.
- (145) Zhou, G.; Lü, M.; Xiu, Z.; Wang, S.; Zhang, H.; Zhou, Y.; Wang, S. Controlled Synthesis of High-Quality PbS Star-Shaped Dendrites, Multipods, Truncated Nanocubes, and Nanocubes and Their Shape Evolution Process. *J. Phys. Chem. B* **2006**, *110*, 6543–6548.
- (146) Martinez, B.; Ramade, J.; Livache, C.; Goubet, N.; Chu, A.; Gréboval, C.; Qu, J.; Watkins, W. L.; Becerra, L.; Dandeu, E.; et al. HgTe Nanocrystal Inks for Extended Short-Wave Infrared Detection. *Adv. Opt. Mater.* **2019**, *7*, 1900348.
- (147) Riedinger, A.; Ott, F. D.; Mule, A.; Mazzotti, S.; Knüsel, P. N.; Kress, S. J. P.; Prins, F.; Erwin, S. C.; Norris, D. J. An Intrinsic Growth Instability in Isotropic Materials Leads to Quasi-Two-Dimensional Nanoplatelets. *Nat. Mater.* **2017**, *16*, 743–748.
- (148) Ithurria, S.; Bousquet, G.; Dubertret, B. Continuous Transition from 3D to 1D Confinement Observed during the Formation of CdSe Nanoplatelets. *J. Am. Chem. Soc.* **2011**, *133*, 3070–3077.
- (149) Tessier, M. D.; Javaux, C.; Maksimovic, I.; Loriette, V.; Dubertret, B. Spectroscopy of Single CdSe Nanoplatelets. *ACS Nano* **2012**, *6*, 6751–6758.
- (150) Grim, J. Q.; Christodoulou, S.; Di Stasio, F.; Krahne, R.; Cingolani, R.; Manna, L.; Moreels, I. Continuous-Wave Biexciton Lasing at Room Temperature Using Solution-Processed Quantum Wells. *Nat. Nanotechnol.* **2014**, *9*, 891–895.
- (151) She, C.; Fedin, I.; Dolzhnikov, D. S.; Demortière, A.; Schaller, R. D.; Pelton, M.; Talapin, D. V. Low-Threshold Stimulated Emission Using Colloidal Quantum Wells. *Nano Lett.* **2014**, *14*, 2772–2777.
- (152) Guzelturk, B.; Kelestemur, Y.; Olutas, M.; Delikanli, S.; Demir, H. V. Amplified Spontaneous Emission and Lasing in Colloidal Nanoplatelets. *ACS Nano* **2014**, *8*, 6599–6605.
- (153) Liu, B.; Altintas, Y.; Wang, L.; Shendre, S.; Sharma, M.; Sun, H.; Mutlugun, E.; Demir, H. V. Record High External Quantum Efficiency of 19.2% Achieved in Light-Emitting Diodes of Colloidal Quantum Wells Enabled by Hot-Injection Shell Growth. *Adv. Mater.* **2020**, *32*, 1905824.
- (154) Chen, Z.; Nadal, B.; Mahler, B.; Aubin, H.; Dubertret, B. Quasi-2D Colloidal Semiconductor Nanoplatelets for Narrow Electroluminescence. *Adv. Funct. Mater.* **2014**, *24*, 295–302.
- (155) Kelestemur, Y.; Shynkarenko, Y.; Anni, M.; Yakunin, S.; De Giorgi, M. L.; Kovalenko, M. V. Colloidal CdSe Quantum Wells with Graded Shell Composition for Low-Threshold Amplified Spontaneous Emission and Highly Efficient Electroluminescence. *ACS Nano* **2019**, *13*, 13899–13909.
- (156) Qu, J.; Rastogi, P.; Greboval, C.; Livache, C.; Dufour, M.; Chu, A.; Chee, S.-S.; Ramade, J.; Xu, X. Z.; Ithurria, S.; et al. Nanoplatelet-Based Light Emitting Diode and Its Use for All-Nanocrystal LiFi-like Communication. *ACS Appl. Mater. Interfaces* **2020**, *12*, 22058–22065.
- (157) Diroll, B. T. Colloidal Quantum Wells for Optoelectronic Devices. *J. Mater. Chem. C* **2020**, *8*, 10628–10640.
- (158) Lhuillier, E.; Ithurria, S.; Descamps-Mandine, A.; Douillard, T.; Castaing, R.; Xu, X. Z.; Taberna, P.-L.; Simon, P.; Aubin, H.; Dubertret, B. Investigating the n- and p-Type Electrolytic Charging of Colloidal Nanoplatelets. *J. Phys. Chem. C* **2015**, *119*, 21795–21799.
- (159) Lhuillier, E.; Dayen, J.-F.; Thomas, D. O.; Robin, A.; Doudin, B.; Dubertret, B. Nanoplatelets Bridging a Nanotrench: A New Architecture for Photodetectors with Increased Sensitivity. *Nano Lett.* **2015**, *15*, 1736–1742.
- (160) Schliehe, C.; Juarez, B. H.; Pelletier, M.; Jander, S.; Greshnykh, D.; Nagel, M.; Meyer, A.; Foerster, S.; Kornowski, A.; Klinke, C.; et al. Ultrathin PbS Sheets by Two-Dimensional Oriented Attachment. *Science* **2010**, *329*, 550–553.
- (161) Pedetti, S.; Nadal, B.; Lhuillier, E.; Mahler, B.; Bouet, C.; Abécassis, B.; Xu, X.; Dubertret, B. Optimized Synthesis of CdTe Nanoplatelets and Photoresponse of CdTe Nanoplatelets Films. *Chem. Mater.* **2013**, *25*, 2455–2462.
- (162) Galle, T.; Samadi Khoshkhoo, M.; Martin-Garcia, B.; Meerbach, C.; Sayevich, V.; Koitzsch, A.; Lesnyak, V.; Eychmüller, A. Colloidal PbSe Nanoplatelets of Varied Thickness with Tunable Optical Properties. *Chem. Mater.* **2019**, *31*, 3803–3811.
- (163) Galle, T.; Kazes, M.; Hübner, R.; Lox, J.; Samadi Khoshkhoo, M.; Sonntag, L.; Tietze, R.; Sayevich, V.; Oron, D.; Koitzsch, A.; et al. Colloidal Mercury-Doped CdSe Nanoplatelets with Dual Fluorescence. *Chem. Mater.* **2019**, *31*, 5065–5074.
- (164) Peterson, J. J.; Krauss, T. D. Fluorescence Spectroscopy of Single Lead Sulfide Quantum Dots. *Nano Lett.* **2006**, *6*, 510–514.
- (165) Gréboval, C.; Izquierdo, E.; Livache, C.; Martinez, B.; Dufour, M.; Goubet, N.; Moghaddam, N.; Qu, J.; Chu, A.; Ramade, J.; et al. Impact of Dimensionality and Confinement on the Electronic Properties of Mercury Chalcogenide Nanocrystals. *Nanoscale* **2019**, *11*, 3905–3915.
- (166) Izquierdo, E.; Dufour, M.; Chu, A.; Livache, C.; Martinez, B.; Amelot, D.; Patriarche, G.; Lequeux, N.; Lhuillier, E.; Ithurria, S. Coupled HgSe Colloidal Quantum Wells through a Tunable Barrier: A Strategy To Uncouple Optical and Transport Band Gap. *Chem. Mater.* **2018**, *30*, 4065–4072.
- (167) Smith, A. M.; Lane, L. A.; Nie, S. Mapping the Spatial Distribution of Charge Carriers in Quantum-Confinement Heterostructures. *Nat. Commun.* **2014**, *5*, 4506.
- (168) Keuleyan, S.; Kohler, J.; Guyot-Sionnest, P. Photoluminescence of Mid-Infrared HgTe Colloidal Quantum Dots. *J. Phys. Chem. C* **2014**, *118*, 2749–2753.
- (169) Deng, Z.; Guyot-Sionnest, P. Intraband Luminescence from HgSe/CdS Core/Shell Quantum Dots. *ACS Nano* **2016**, *10*, 2121–2127.
- (170) Kershaw, S. V.; Harrison, M.; Rogach, A. L.; Kornowski, A. Development of IR-Emitting Colloidal II-VI Quantum-Dot Materials. *IEEE J. Sel. Top. Quantum Electron.* **2000**, *6*, 534–543.
- (171) Lifshitz, E.; Porteanu, H.; Glozman, A.; Weller, H.; Pflughoefft, M.; Eychmüller, A. Optically Detected Magnetic Resonance Study of CdS/HgS/CdS Quantum Dot Quantum Wells. *J. Phys. Chem. B* **1999**, *103*, 6870–6875.
- (172) Eychmüller, A.; Voßmeyer, T.; Mews, A.; Weller, H. Transient Photobleaching in the Quantum Dot Quantum Well CdS/HgS/CdS. *J. Lumin.* **1994**, *58*, 223–226.
- (173) Mews, A.; Eychmüller, A. Quantum Wells within Quantum Dots, a CdS/HgS Nanoheterostructure with Global and Local Confinement. *Ber. Bunsenges. Phys. Chem.* **1998**, *102*, 1343–1357.
- (174) Porteanu, H. E.; Lifshitz, E.; Pflughoefft, M.; Eychmüller, A.; Weller, H. Optical Properties of CdS/HgS/CdS Quantum Dot–Quantum Well Structures. *Phys. Status Solidi B* **2001**, *226*, 219–232.
- (175) Smith, A. M.; Nie, S. Bright and Compact Alloyed Quantum Dots with Broadly Tunable Near-Infrared Absorption and Fluorescence Spectra through Mercury Cation Exchange. *J. Am. Chem. Soc.* **2011**, *133*, 24–26.
- (176) Kershaw, S. V.; Burt, M.; Harrison, M.; Rogach, A.; Weller, H.; Eychmüller, A. Colloidal CdTe/HgTe Quantum Dots with High Photoluminescence Quantum Efficiency at Room Temperature. *Appl. Phys. Lett.* **1999**, *75*, 1694–1696.
- (177) Pietryga, J. M.; Werder, D. J.; Williams, D. J.; Casson, J. L.; Schaller, R. D.; Klimov, V. I.; Hollingsworth, J. A. Utilizing the Lability

- of Lead Selenide to Produce Heterostructured Nanocrystals with Bright, Stable Infrared Emission. *J. Am. Chem. Soc.* **2008**, *130*, 4879–4885.
- (178) Lambert, K.; Geyter, B. D.; Moreels, I.; Hens, Z. PbTelCdTe Core/Shell Particles by Cation Exchange, a HR-TEM Study. *Chem. Mater.* **2009**, *21*, 778–780.
- (179) Zhou, H. S.; Honma, I.; Komiyama, H.; Haus, J. W. Coated Semiconductor Nanoparticles; the Cadmium Sulfide/Lead Sulfide System's Synthesis and Properties. *J. Phys. Chem.* **1993**, *97*, 895–901.
- (180) Justo, Y.; Geiregat, P.; van Hoecke, K.; Vanhaecke, F.; De Mello Donega, C.; Hens, Z. Optical Properties of PbS/CdS Core/Shell Quantum Dots. *J. Phys. Chem. C* **2013**, *117*, 20171–20177.
- (181) Ithurria, S.; Talapin, D. V. Colloidal Atomic Layer Deposition (c-ALD) Using Self-Limiting Reactions at Nanocrystal Surface Coupled to Phase Transfer between Polar and Nonpolar Media. *J. Am. Chem. Soc.* **2012**, *134*, 18585–18590.
- (182) Hazarika, A.; Fedin, I.; Hong, L.; Guo, J.; Srivastava, V.; Cho, W.; Coropceanu, I.; Portner, J.; Diroll, B. T.; Philbin, J. P.; et al. Colloidal Atomic Layer Deposition with Stationary Reactant Phases Enables Precise Synthesis of "Digital" II–VI Nano-Heterostructures with Exquisite Control of Confinement and Strain. *J. Am. Chem. Soc.* **2019**, *141*, 13487–13496.
- (183) Lhuillier, E.; Pedetti, S.; Ithurria, S.; Nadal, B.; Heuclin, H.; Dubertret, B. Two-Dimensional Colloidal Metal Chalcogenides Semiconductors: Synthesis, Spectroscopy, and Applications. *Acc. Chem. Res.* **2015**, *48*, 22–30.
- (184) Li, J. J.; Wang, Y. A.; Guo, W.; Keay, J. C.; Mishima, T. D.; Johnson, M. B.; Peng, X. Large-Scale Synthesis of Nearly Monodisperse CdSe/CdS Core/Shell Nanocrystals Using Air-Stable Reagents via Successive Ion Layer Adsorption and Reaction. *J. Am. Chem. Soc.* **2003**, *125*, 12567–12575.
- (185) Sagar, L. K.; Walravens, W.; Maes, J.; Geiregat, P.; Hens, Z. HgSe/CdE (E = S, Se) Core/Shell Nanocrystals by Colloidal Atomic Layer Deposition. *J. Phys. Chem. C* **2017**, *121*, 13816–13822.
- (186) Robin, A.; Livache, C.; Ithurria, S.; Lacaze, E.; Dubertret, B.; Lhuillier, E. Surface Control of Doping in Self-Doped Nanocrystals. *ACS Appl. Mater. Interfaces* **2016**, *8*, 27122–27128.
- (187) Shen, G.; Guyot-Sionnest, P. HgTe/CdTe and HgSe/CdX (X = S, Se, and Te) Core/Shell Mid-Infrared Quantum Dots. *Chem. Mater.* **2019**, *31*, 286–293.
- (188) Lhuillier, E.; Pedetti, S.; Ithurria, S.; Heuclin, H.; Nadal, B.; Robin, A.; Patriarche, G.; Lequeux, N.; Dubertret, B. Electrolyte-Gated Field Effect Transistor to Probe the Surface Defects and Morphology in Films of Thick CdSe Colloidal Nanoplatelets. *ACS Nano* **2014**, *8*, 3813–3820.
- (189) Chu, A.; Livache, C.; Ithurria, S.; Lhuillier, E. Electronic Structure Robustness and Design Rules for 2D Colloidal Heterostructures. *J. Appl. Phys.* **2018**, *123*, 035701.
- (190) Rossinelli, A. A.; Riedinger, A.; Marqués-Gallego, P.; Knüsel, P. N.; Antolinez, F. V.; Norris, D. J. High-Temperature Growth of Thick-Shell CdSe/CdS Core/Shell Nanoplatelets. *Chem. Commun.* **2017**, *53*, 9938–9941.
- (191) Piveteau, L.; Dirin, D.; Gordon, C. P.; Walder, B. J.; Ong, T.-C.; Emsley, L.; Copéret, C.; Kovalenko, M. V. Colloidal-ALD-Grown Core/Shell CdSe/CdS Nanoplatelets as Seen by DNP Enhanced PASS-PIETA NMR Spectroscopy. *Nano Lett.* **2020**, *20*, 3003–3018.
- (192) Jagtap, A.; Martinez, B.; Goubet, N.; Chu, A.; Livache, C.; Gréboval, C.; Ramade, J.; Amelot, D.; Trousset, P.; Triboulon, A.; et al. Design of a Unipolar Barrier for a Nanocrystal-Based Short-Wave Infrared Photodiode. *ACS Photonics* **2018**, *5*, 4569–4576.
- (193) Yoon, B.; Jeong, J.; Jeong, K. S. Higher Quantum State Transitions in Colloidal Quantum Dot with Heavy Electron Doping. *J. Phys. Chem. C* **2016**, *120*, 22062–22068.
- (194) Kim, J.; Yoon, B.; Kim, J.; Choi, Y.; Kwon, Y.-W.; Park, S. K.; Jeong, K. S. High Electron Mobility of β -HgS Colloidal Quantum Dots with Doubly Occupied Quantum States. *RSC Adv.* **2017**, *7*, 38166–38170.
- (195) Brown, P. R.; Kim, D.; Lunt, R. R.; Zhao, N.; Bawendi, M. G.; Grossman, J. C.; Bulović, V. Energy Level Modification in Lead Sulfide Quantum Dot Thin Films through Ligand Exchange. *ACS Nano* **2014**, *8*, 5863–5872.
- (196) Ganeshan, A. A.; Houtepen, A. J.; Crisp, R. W. Quantum Dot Solar Cells: Small Beginnings Have Large Impacts. *Appl. Sci.* **2018**, *8*, 1867.
- (197) Martinez, B.; Plumont, R.; Gréboval, C.; Rastogi, P.; Prado, Y.; Qu, J.; Chu, A.; Livache, C.; Xu, X. Z.; Cruguel, H.; et al. Azobenzenes as Light-Activatable Carrier Density Switches in Nanocrystals. *J. Phys. Chem. C* **2019**, *123*, 27257–27263.
- (198) Hartley, G. S. The Cis-Form of Azobenzene. *Nature* **1937**, *140*, 281–281.
- (199) Beharry, A. A.; Woolley, G. A. Azobenzene Photoswitches for Biomolecules. *Chem. Soc. Rev.* **2011**, *40*, 4422–4437.
- (200) Kumar, G. S.; Neckers, D. C. Photochemistry of Azobenzene-Containing Polymers. *Chem. Rev.* **1989**, *89*, 1915–1925.
- (201) Samanta, D.; Gemen, J.; Chu, Z.; Diskin-Posner, Y.; Shimon, L. J. W.; Klajn, R. Reversible Photoswitching of Encapsulated Azobenzenes in Water. *Proc. Natl. Acad. Sci. U. S. A.* **2018**, *115*, 9379–9384.
- (202) Credi, A. Artificial Molecular Motors Powered by Light. *Aust. J. Chem.* **2006**, *59*, 157–169.
- (203) Hartley, G. S.; Le Fèvre, R. J. W. 119. The Dipole Moments of Cis- and Trans-Azobenzenes and of Some Related Compounds. *J. Chem. Soc.* **1939**, *1939*, 531–535.
- (204) Martinez, B.; Livache, C.; Meriggiò, E.; Xu, X. Z.; Cruguel, H.; Lacaze, E.; Proust, A.; Ithurria, S.; Silly, M. G.; Cabailh, G.; et al. Polyoxometalate as Control Agent for the Doping in HgSe Self-Doped Nanocrystals. *J. Phys. Chem. C* **2018**, *122*, 26680–26685.
- (205) Huang, J.; Liu, W.; Dolzhnikov, D. S.; Protesescu, L.; Kovalenko, M. V.; Koo, B.; Chattopadhyay, S.; Shenchenko, E. V.; Talapin, D. V. Surface Functionalization of Semiconductor and Oxide Nanocrystals with Small Inorganic Oxoanions (PO_4^{3-} , MoO_4^{2-}) and Polyoxometalate Ligands. *ACS Nano* **2014**, *8*, 9388–9402.
- (206) Liu, H.; Guyot-Sionnest, P. Magnetoresistance of Manganese-Doped Colloidal Quantum Dot Films. *J. Phys. Chem. C* **2015**, *119*, 14797–14804.
- (207) Yang, H.; Wong, E.; Zhao, T.; Lee, J. D.; Xin, H. L.; Chi, M.; Fleury, B.; Tang, H.-Y.; Gaulding, E. A.; Kagan, C. R.; et al. Charge Transport Modulation in PbSe Nanocrystal Solids by AuxAg_{1-x} Nanoparticle Doping. *ACS Nano* **2018**, *12*, 9091–9100.
- (208) Cargnello, M.; Johnston-Peck, A. C.; Diroll, B. T.; Wong, E.; Datta, B.; Damodhar, D.; Doan-Nguyen, V. V. T.; Herzing, A. A.; Kagan, C. R.; Murray, C. B. Substitutional Doping in Nanocrystal Superlattices. *Nature* **2015**, *524*, 450–453.
- (209) Groves, S. H.; Brown, R. N.; Pidgeon, C. R. Interband Magnetoreflection and Band Structure of HgTe. *Phys. Rev.* **1967**, *161*, 779–793.
- (210) Groves, S.; Paul, W. Band Structure of Gray Tin. *Phys. Rev. Lett.* **1963**, *11*, 194–196.
- (211) Bendias, K.; Shamim, S.; Herrmann, O.; Budewitz, A.; Shekhar, P.; Leubner, P.; Kleinlein, J.; Bocquillon, E.; Buhmann, H.; Molenkamp, L. W. High Mobility HgTe Microstructures for Quantum Spin Hall Studies. *Nano Lett.* **2018**, *18*, 4831–4836.
- (212) Moon, C.-Y.; Wei, S.-H. Band Gap of Hg Chalcogenides: Symmetry-Reduction-Induced Band-Gap Opening of Materials with Inverted Band Structures. *Phys. Rev. B: Condens. Matter Mater. Phys.* **2006**, *74*, 045205.
- (213) Madelung, O. *Semiconductors: Data Handbook*; Springer Science & Business Media, 2012.
- (214) Pokatilov, E. P.; Fonoberov, V. A.; Fomin, V. M.; Devreese, J. T. Electron and Hole States in Quantum Dot Quantum Wells within a Spherical Eight-Band Model. *Phys. Rev. B: Condens. Matter Mater. Phys.* **2001**, *64*, 245329.
- (215) Beugeling, W.; Kalesaki, E.; Delerue, C.; Niquet, Y.-M.; Vanmaekelbergh, D.; Smith, C. M. Topological States in Multi-Orbital HgTe Honeycomb Lattices. *Nat. Commun.* **2015**, *6*, 6316.
- (216) Allan, G.; Delerue, C. Tight-Binding Calculations of the Optical Properties of HgTe Nanocrystals. *Phys. Rev. B: Condens. Matter Mater. Phys.* **2012**, *86*, 165437.

- (217) Willardson, R. K.; Beer, A. C. *Semiconductors and Semimetals*; Academic Press, 1977; p 18.
- (218) Al-Otaify, A.; Kershaw, S. V.; Gupta, S.; Rogach, A. L.; Allan, G.; Delerue, C.; Binks, D. J. Multiple Exciton Generation and Ultrafast Exciton Dynamics in HgTe Colloidal Quantum Dots. *Phys. Chem. Chem. Phys.* **2013**, *15*, 16864–16873.
- (219) Livache, C.; Goubet, N.; Martinez, B.; Jagtap, A.; Qu, J.; Ithurria, S.; Silly, M. G.; Dubertret, B.; Lhuillier, E. Band Edge Dynamics and Multiexciton Generation in Narrow Band Gap HgTe Nanocrystals. *ACS Appl. Mater. Interfaces* **2018**, *10*, 11880–11887.
- (220) Evers, W. H.; Goris, B.; Bals, S.; Casavola, M.; de Graaf, J.; van Roij, R.; Dijkstra, M.; Vanmaekelbergh, D. Low-Dimensional Semiconductor Superlattices Formed by Geometric Control over Nanocrystal Attachment. *Nano Lett.* **2013**, *13*, 2317–2323.
- (221) Geuchies, J. J.; van Overbeek, C.; Evers, W. H.; Goris, B.; de Backer, A.; Gantapara, A. P.; Rabouw, F. T.; Hilhorst, J.; Peters, J. L.; Konovalov, O.; et al. In Situ Study of the Formation Mechanism of Two-Dimensional Superlattices from PbSe Nanocrystals. *Nat. Mater.* **2016**, *15*, 1248–1254.
- (222) Boneschanscher, M. P.; Evers, W. H.; Geuchies, J. J.; Altantzis, T.; Goris, B.; Rabouw, F. T.; van Rossum, S. A. P.; van der Zant, H. S. J.; Siebbeles, L. D. A.; Tendeloo, G. V.; et al. Long-Range Orientation and Atomic Attachment of Nanocrystals in 2D Honeycomb Superlattices. *Science* **2014**, *344*, 1377–1380.
- (223) Alimoradi Jazi, M.; Janssen, V. A. E. C.; Evers, W. H.; Tadjine, A.; Delerue, C.; Siebbeles, L. D. A.; van der Zant, H. S. J.; Houtepen, A. J.; Vanmaekelbergh, D. Transport Properties of a Two-Dimensional PbSe Square Superstructure in an Electrolyte-Gated Transistor. *Nano Lett.* **2017**, *17*, 5238–5243.
- (224) Evers, W. H.; Schins, J. M.; Aerts, M.; Kulkarni, A.; Capiod, P.; Berthe, M.; Grandidier, B.; Delerue, C.; van der Zant, H. S. J.; van Overbeek, C.; et al. High Charge Mobility in Two-Dimensional Percolative Networks of PbSe Quantum Dots Connected by Atomic Bonds. *Nat. Commun.* **2015**, *6*, 8195.
- (225) Lu, H.; Carroll, G. M.; Chen, X.; Amarasinghe, D. K.; Neale, N. R.; Miller, E. M.; Sercel, P. C.; Rabuffetti, F. A.; Efros, A. L.; Beard, M. C. N-Type PbSe Quantum Dots via Post-Synthetic Indium Doping. *J. Am. Chem. Soc.* **2018**, *140*, 13753–13763.
- (226) Ramiro, I.; Kundu, B.; Dalmases, M.; Özdemir, O.; Pedrosa, M.; Konstantatos, G. Size- and Temperature-Dependent Intraband Optical Properties of Heavily n-Doped PbS Colloidal Quantum Dot Solid-State Films. *ACS Nano* **2020**, *14*, 7161–7169.
- (227) Ramiro, I.; Özdemir, O.; Christodoulou, S.; Gupta, S.; Dalmases, M.; Torre, I.; Konstantatos, G. Mid- and Long-Wave Infrared Optoelectronics via Intraband Transitions in PbS Colloidal Quantum Dots. *Nano Lett.* **2020**, *20*, 1003–1008.
- (228) Bejaoui, A.; Alonso, M. I.; Garriga, M.; Campoy-Quiles, M.; Goñi, A. R.; Hetsch, F.; Kershaw, S. V.; Rogach, A. L.; To, C. H.; Foo, Y.; et al. Evaluation of the Dielectric Function of Colloidal Cd_{1-x}Hg_xTe Quantum Dot Films by Spectroscopic Ellipsometry. *Appl. Surf. Sci.* **2017**, *421*, 295–300.
- (229) Grynberg, M.; Le Toullec, R.; Balkanski, M. Dielectric Function in HgTe between 8 and 300 K. *Phys. Rev. B* **1974**, *9*, 517–526.
- (230) Szuszkiewicz, W.; Witowski, A. M.; Grynberg, M. The Dynamic Dielectric Function in HgSe and HgTe. *Phys. Status Solidi B* **1978**, *87*, 637–645.
- (231) Houel, J.; Sauvage, S.; Boucaud, P.; Dazzi, A.; Prazeres, R.; Glotin, F.; Ortéga, J.-M.; Miard, A.; Lemaître, A. Ultraweak-Absorption Microscopy of a Single Semiconductor Quantum Dot in the Midinfrared Range. *Phys. Rev. Lett.* **2007**, *99*, 217404.
- (232) Peter, E.; Senellart, P.; Martrou, D.; Lemaître, A.; Hours, J.; Gérard, J. M.; Bloch, J. Exciton-Photon Strong-Coupling Regime for a Single Quantum Dot Embedded in a Microcavity. *Phys. Rev. Lett.* **2005**, *95*, 067401.
- (233) Marzin, J.-Y.; Gérard, J.-M.; Izraël, A.; Barrier, D.; Bastard, G. Photoluminescence of Single InAs Quantum Dots Obtained by Self-Organized Growth on GaAs. *Phys. Rev. Lett.* **1994**, *73*, 716–719.
- (234) Somaschi, N.; Giesz, V.; De Santis, L.; Loredo, J. C.; Almeida, M. P.; Hornecker, G.; Portalupi, S. L.; Grange, T.; Antón, C.; Demory, J.; et al. Near-Optimal Single-Photon Sources in the Solid State. *Nat. Photonics* **2016**, *10*, 340–345.
- (235) Guenzer, C. S.; Bienenstock, A. Temperature Dependence of the HgTe Band Gap. *Phys. Rev. B* **1973**, *8*, 4655–4667.
- (236) Dobrowolska, M.; Mycielski, A.; Dobrowolski, W. Determination of Temperature Dependence of Energy Gap in HgTe by Oscillatory Magnetotransmission Measurements. *Solid State Commun.* **1978**, *27*, 1233–1235.
- (237) Guldner, Y.; Bastard, G.; Voos, M. Calculated Temperature Dependence of the Band Gap of HgTe-CdTe Superlattices. *J. Appl. Phys.* **1985**, *57*, 1403–1405.
- (238) Olkhovets, A.; Hsu, R.-C.; Lipovskii, A.; Wise, F. W. Size-Dependent Temperature Variation of the Energy Gap in Lead-Salt Quantum Dots. *Phys. Rev. Lett.* **1998**, *81*, 3539–3542.
- (239) Ramade, J.; Andriambariariaona, L. M.; Steinmetz, V.; Goubet, N.; Legrand, L.; Barisien, T.; Bernardot, F.; Testelin, C.; Lhuillier, E.; Bramati, A.; et al. Exciton-Phonon Coupling in a CsPbBr₃ Single Nanocrystal. *Appl. Phys. Lett.* **2018**, *112*, 072104.
- (240) Livache, C.; Goubet, N.; Gréboval, C.; Martinez, B.; Ramade, J.; Qu, J.; Triboulin, A.; Cruguel, H.; Baptiste, B.; Klotz, S.; et al. Effect of Pressure on Interband and Intraband Transition of Mercury Chalcogenide Quantum Dots. *J. Phys. Chem. C* **2019**, *123*, 13122–13130.
- (241) Bernevig, B. A.; Hughes, T. L.; Zhang, S.-C. Quantum Spin Hall Effect and Topological Phase Transition in HgTe Quantum Wells. *Science* **2006**, *314*, 1757–1761.
- (242) König, M.; Wiedmann, S.; Brüne, C.; Roth, A.; Buhmann, H.; Molenkamp, L. W.; Qi, X.-L.; Zhang, S.-C. Quantum Spin Hall Insulator State in HgTe Quantum Wells. *Science* **2007**, *318*, 766–770.
- (243) Wu, G. Y.; McGill, T. C. Strain Effects in HgTe-CdTe Superlattices Grown on CdTe Substrates. *Appl. Phys. Lett.* **1985**, *47*, 634–636.
- (244) Brüne, C.; Liu, C. X.; Novik, E. G.; Hankiewicz, E. M.; Buhmann, H.; Chen, Y. L.; Qi, X. L.; Shen, Z. X.; Zhang, S. C.; Molenkamp, L. W. Quantum Hall Effect from the Topological Surface States of Strained Bulk HgTe. *Phys. Rev. Lett.* **2011**, *106*, 126803.
- (245) Wright, N. G.; McMahon, M. I.; Nelmes, R. J.; San-Miguel, A. Crystal Structure of the Cinnabar Phase of HgTe. *Phys. Rev. B: Condens. Matter Mater. Phys.* **1993**, *48*, 13111–13114.
- (246) San-Miguel, A.; Wright, N. G.; McMahon, M. I.; Nelmes, R. J. Pressure Evolution of the Cinnabar Phase of HgTe. *Phys. Rev. B: Condens. Matter Mater. Phys.* **1995**, *51*, 8731–8736.
- (247) Werner, A.; Hochheimer, H. D.; Strössner, K.; Jayaraman, A. High-Pressure X-Ray Diffraction Studies on HgTe and HgS to 20 GPa. *Phys. Rev. B: Condens. Matter Mater. Phys.* **1983**, *28*, 3330–3334.
- (248) Kafalas, J. A.; Gatos, H. C.; Lavine, M. C.; Banus, M. D. High Pressure Phase Transition in Mercury Selenide. *J. Phys. Chem. Solids* **1962**, *23*, 1541–1544.
- (249) Ithurria, S.; Guyot-Sionnest, P.; Mahler, B.; Dubertret, B. Mn²⁺ as a Radial Pressure Gauge in Colloidal Core/Shell Nanocrystals. *Phys. Rev. Lett.* **2007**, *99*, 265501.
- (250) Tolbert, S. H.; Alivisatos, A. P. Size Dependence of a First Order Solid-Solid Phase Transition: The Wurtzite to Rock Salt Transformation in CdSe Nanocrystals. *Science* **1994**, *265*, 373–376.
- (251) Düz, I.; Erdem, I.; Ozdemir, K. S.; Kuzucu, V. First Principles Investigations of HgX (X = S, Se and Te). *Arch. Mater. Sci. Eng.* **2016**, *79*, 5–11.
- (252) Steiner, D.; Dorfs, D.; Banin, U.; Della Sala, F.; Manna, L.; Millo, O. Determination of Band Offsets in Heterostructured Colloidal Nanorods Using Scanning Tunneling Spectroscopy. *Nano Lett.* **2008**, *8*, 2954–2958.
- (253) Diaconescu, B.; Padilha, L. A.; Nagpal, P.; Swartzentruber, B. S.; Klimov, V. I. Measurement of Electronic States of PbS Nanocrystal Quantum Dots Using Scanning Tunneling Spectroscopy: The Role of Parity Selection Rules in Optical Absorption. *Phys. Rev. Lett.* **2013**, *110*, 127406.
- (254) Swart, I.; Liljeroth, P.; Vanmaekelbergh, D. Scanning Probe Microscopy and Spectroscopy of Colloidal Semiconductor Nanocrystals and Assembled Structures. *Chem. Rev.* **2016**, *116*, 11181–11219.

- (255) Liljeroth, P.; Jdira, L.; Overgaag, K.; Grandidier, B.; Speller, S.; Vanmaekelbergh, D. Can Scanning Tunnelling Spectroscopy Measure the Density of States of Semiconductor Quantum Dots? *Phys. Chem. Chem. Phys.* **2006**, *8*, 3845–3850.
- (256) Overgaag, K.; Liljeroth, P.; Grandidier, B.; Vanmaekelbergh, D. Scanning Tunneling Spectroscopy of Individual PbSe Quantum Dots and Molecular Aggregates Stabilized in an Inert Nanocrystal Matrix. *ACS Nano* **2008**, *2*, 600–606.
- (257) Jdira, L.; Liljeroth, P.; Stoffels, E.; Vanmaekelbergh, D.; Speller, S. Size-Dependent Single-Particle Energy Levels and Interparticle Coulomb Interactions in CdSe Quantum Dots Measured by Scanning Tunneling Spectroscopy. *Phys. Rev. B: Condens. Matter Mater. Phys.* **2006**, *73*, 115305.
- (258) Banin, U.; Cao, Y.; Katz, D.; Millo, O. Identification of Atomic-like Electronic States in Indium Arsenide Nanocrystal Quantum Dots. *Nature* **1999**, *400*, 542–544.
- (259) Wang, H.; Lhuillier, E.; Yu, Q.; Zimmers, A.; Dubertret, B.; Ulysse, C.; Aubin, H. Transport in a Single Self-Doped Nanocrystal. *ACS Nano* **2017**, *11*, 1222–1229.
- (260) Janowitz, C.; Orlowski, N.; Manzke, R.; Golacki, Z. On the Band Structure of HgTe and HgSe — View from Photoemission. *J. Alloys Compd.* **2001**, *328*, 84–89.
- (261) Bisquert, J.; Fabregat-Santiago, F.; Mora-Seró, I.; García-Belmonte, G.; Barea, E. M.; Palomares, E. A Review of Recent Results on Electrochemical Determination of the Density of Electronic States of Nanostructured Metal-Oxide Semiconductors and Organic Hole Conductors. *Inorg. Chim. Acta* **2008**, *361*, 684–698.
- (262) Livache, C.; Izquierdo, E.; Martinez, B.; Dufour, M.; Pierucci, D.; Keuleyan, S.; Cruguel, H.; Becerra, L.; Fave, J. L.; Aubin, H.; et al. Charge Dynamics and Optoelectronic Properties in HgTe Colloidal Quantum Wells. *Nano Lett.* **2017**, *17*, 4067–4074.
- (263) Spittel, D.; Poppe, J.; Meerbach, C.; Ziegler, C.; Hickey, S. G.; Eychmüller, A. Absolute Energy Level Positions in CdSe Nanostructures from Potential-Modulated Absorption Spectroscopy (EMAS). *ACS Nano* **2017**, *11*, 12174–12184.
- (264) Haram, S. K.; Quinn, B. M.; Bard, A. J. Electrochemistry of CdS Nanoparticles: A Correlation between Optical and Electrochemical Band Gaps. *J. Am. Chem. Soc.* **2001**, *123*, 8860–8861.
- (265) Sobrova, P.; Ryvolova, M.; Hubalek, J.; Adam, V.; Kizek, R. Voltammetry as a Tool for Characterization of CdTe Quantum Dots. *Int. J. Mol. Sci.* **2013**, *14*, 13497–13510.
- (266) Liu, H.; Keuleyan, S.; Guyot-Sionnest, P. n- and p-Type HgTe Quantum Dot Films. *J. Phys. Chem. C* **2012**, *116*, 1344–1349.
- (267) Kroupa, D. M.; Vörös, M.; Brawand, N. P.; McNichols, B. W.; Miller, E. M.; Gu, J.; Nozik, A. J.; Sellinger, A.; Galli, G.; Beard, M. C. Tuning Colloidal Quantum Dot Band Edge Positions through Solution-Phase Surface Chemistry Modification. *Nat. Commun.* **2017**, *8*, 15257.
- (268) Melnychuk, C.; Guyot-Sionnest, P. Slow Auger Relaxation in HgTe Colloidal Quantum Dots. *J. Phys. Chem. Lett.* **2018**, *9*, 2208–2211.
- (269) Melnychuk, C.; Guyot-Sionnest, P. Auger Suppression in n-Type HgSe Colloidal Quantum Dots. *ACS Nano* **2019**, *13*, 10512–10519.
- (270) Leontiadou, M. A.; Al-Otaify, A.; Kershaw, S. V.; Zhovtiuk, O.; Kalytchuk, S.; Mott, D.; Maenosono, S.; Rogach, A. L.; Binks, D. J. Ultrafast Exciton Dynamics in $\text{Cd}_x\text{Hg}_{(1-x)}\text{Te}$ Alloy Quantum Dots. *Chem. Phys.* **2016**, *469–470*, 25–30.
- (271) Liu, H.; Guyot-Sionnest, P. Photoluminescence Lifetime of Lead Selenide Colloidal Quantum Dots. *J. Phys. Chem. C* **2010**, *114*, 14860–14863.
- (272) Robel, I.; Gresback, R.; Kortshagen, U.; Schaller, R. D.; Klimov, V. I. Universal Size-Dependent Trend in Auger Recombination in Direct-Gap and Indirect-Gap Semiconductor Nanocrystals. *Phys. Rev. Lett.* **2009**, *102*, 177404.
- (273) Klimov, V. I.; McGuire, J. A.; Schaller, R. D.; Rupasov, V. I. Scaling of Multiexciton Lifetimes in Semiconductor Nanocrystals. *Phys. Rev. B: Condens. Matter Mater. Phys.* **2008**, *77*, 195324.
- (274) Neppi, S.; Gessner, O. Time-Resolved X-Ray Photoelectron Spectroscopy Techniques for the Study of Interfacial Charge Dynamics. *J. Electron Spectrosc. Relat. Phenom.* **2015**, *200*, 64–77.
- (275) Spencer, B. F.; Cliffe, M. J.; Graham, D. M.; Hardman, S. J. O.; Seddon, E. A.; Syres, K. L.; Thomas, A. G.; Sirotti, F.; Silly, M. G.; Akhtar, J.; et al. Chemically-Specific Time-Resolved Surface Photovoltage Spectroscopy: Carrier Dynamics at the Interface of Quantum Dots Attached to a Metal Oxide. *Surf. Sci.* **2015**, *641*, 320–325.
- (276) Gréboval, C.; Rastogi, P.; Qu, J.; Chu, A.; Ramade, J.; Khalili, A.; Dabard, C.; Dang, T. H.; Cruguel, H.; Ouerghi, A.; et al. Time Resolved Photoemission to Unveil Electronic Coupling Between Absorbing and Transport Layers in a Quantum Dot Based Solar Cell. *J. Phys. Chem. C* **2020**, *124*, 23400–23409.
- (277) Amelot, D.; Rastogi, P.; Martinez, B.; Gréboval, C.; Livache, C.; Bresciani, F. A.; Qu, J.; Chu, A.; Goyal, M.; Chee, S.-S.; et al. Revealing the Band Structure of FAPI Quantum Dot Film and Its Interfaces with Electron and Hole Transport Layer Using Time Resolved Photoemission. *J. Phys. Chem. C* **2020**, *124*, 3873–3880.
- (278) Martinez, B.; Livache, C.; Goubet, N.; Jagtap, A.; Cruguel, H.; Ouerghi, A.; Lacaze, E.; Silly, M. G.; Lhuillier, E. Probing Charge Carrier Dynamics to Unveil the Role of Surface Ligands in HgTe Narrow Band Gap Nanocrystals. *J. Phys. Chem. C* **2018**, *122*, 859–865.
- (279) Lhuillier, E.; Keuleyan, S.; Rekemeyer, P.; Guyot-Sionnest, P. Thermal Properties of Mid-Infrared Colloidal Quantum Dot Detectors. *J. Appl. Phys.* **2011**, *110*, 033110.
- (280) Fidler, A. F.; Gao, J.; Klimov, V. I. Electron–Hole Exchange Blockade and Memory-Less Recombination in Photoexcited Films of Colloidal Quantum Dots. *Nat. Phys.* **2017**, *13*, 604–610.
- (281) Gao, J.; Fidler, A. F.; Klimov, V. I. Carrier Multiplication Detected through Transient Photocurrent in Device-Grade Films of Lead Selenide Quantum Dots. *Nat. Commun.* **2015**, *6*, 8185.
- (282) Chen, M.; Yu, H.; Kershaw, S. V.; Xu, H.; Gupta, S.; Hetsch, F.; Rogach, A. L.; Zhao, N. Fast, Air-Stable Infrared Photodetectors Based on Spray-Deposited Aqueous HgTe Quantum Dots. *Adv. Funct. Mater.* **2014**, *24*, 53–59.
- (283) Orenstein, J.; Kastner, M. A.; Vaninov, V. Transient Photoconductivity and Photo-Induced Optical Absorption in Amorphous Semiconductors. *Philos. Mag. B* **1982**, *46*, 23–62.
- (284) Main, C.; Nesheva, D. Transient Photocurrent Techniques as a Means of Characterising Amorphous Semiconductors. *J. Optoelectron. Adv. Mater.* **2001**, *3*, 655–664.
- (285) Orenstein, J.; Kastner, M. Photocurrent Transient Spectroscopy: Measurement of the Density of Localized States in $a\text{-As}_2\text{Se}_3$. *Phys. Rev. Lett.* **1981**, *46*, 1421–1424.
- (286) Guyot-Sionnest, P.; Lhuillier, E.; Liu, H. A Mirage Study of CdSe Colloidal Quantum Dot Films, Urbach Tail, and Surface States. *J. Chem. Phys.* **2012**, *137*, 154704.
- (287) Guyot-Sionnest, P. Electrical Transport in Colloidal Quantum Dot Films. *J. Phys. Chem. Lett.* **2012**, *3*, 1169–1175.
- (288) Koktysh, D. S.; Gaponik, N.; Reufer, M.; Crewett, J.; Scherf, U.; Eychmüller, A.; Lupton, J. M.; Rogach, A. L.; Feldmann, J. Near-Infrared Electroluminescence from HgTe Nanocrystals. *ChemPhysChem* **2004**, *5*, 1435–1438.
- (289) O'Connor, É.; O'Riordan, A.; Doyle, H.; Moynihan, S.; Cuddihy, A.; Redmond, G. Near-Infrared Electroluminescent Devices Based on Colloidal HgTe Quantum Dot Arrays. *Appl. Phys. Lett.* **2005**, *86*, 201114.
- (290) Rogach, A. L.; Koktysh, D. S.; Harrison, M.; Kotov, N. A. Layer-by-Layer Assembled Films of HgTe Nanocrystals with Strong Infrared Emission. *Chem. Mater.* **2000**, *12*, 1526–1528.
- (291) Wen, Q.; Kershaw, S. V.; Kalytchuk, S.; Zhovtiuk, O.; Reckmeier, C.; Vasilevskiy, M. I.; Rogach, A. L. Impact of $\text{D}_2\text{O}/\text{H}_2\text{O}$ Solvent Exchange on the Emission of HgTe and CdTe Quantum Dots: Polaron and Energy Transfer Effects. *ACS Nano* **2016**, *10*, 4301–4311.
- (292) Gersten, J.; Nitzan, A. Spectroscopic Properties of Molecules Interacting with Small Dielectric Particles. *J. Chem. Phys.* **1981**, *75*, 1139–1152.
- (293) van Driel, A. F.; Allan, G.; Delerue, C.; Lodahl, P.; Vos, W. L.; Vanmaekelbergh, D. Frequency-Dependent Spontaneous Emission

- Rate from CdSe and CdTe Nanocrystals: Influence of Dark States. *Phys. Rev. Lett.* **2005**, *95*, 236804.
- (294) Klimov, V. I.; Mikhailovsky, A. A.; Xu, S.; Malko, A.; Hollingsworth, J. A.; Leatherdale, C. A.; Eisler, H.-J.; Bawendi, M. G. Optical Gain and Stimulated Emission in Nanocrystal Quantum Dots. *Science* **2000**, *290*, 314–317.
- (295) Lim, J.; Park, Y.-S.; Klimov, V. I. Optical Gain in Colloidal Quantum Dots Achieved with Direct-Current Electrical Pumping. *Nat. Mater.* **2018**, *17*, 42–49.
- (296) Roh, J.; Park, Y.-S.; Lim, J.; Klimov, V. I. Optically Pumped Colloidal-Quantum-Dot Lasing in LED-like Devices with an Integrated Optical Cavity. *Nat. Commun.* **2020**, *11*, 271.
- (297) Geiregat, P.; Van Thourhout, D.; Hens, Z. A Bright Future for Colloidal Quantum Dot Lasers. *NPG Asia Mater.* **2019**, *11*, 41.
- (298) Fan, F.; Voznyy, O.; Sabatini, R. P.; Bicanic, K. T.; Adachi, M. M.; McBride, J. R.; Reid, K. R.; Park, Y.-S.; Li, X.; Jain, A.; et al. Continuous-Wave Lasing in Colloidal Quantum Dot Solids Enabled by Facet-Selective Epitaxy. *Nature* **2017**, *544*, 75–79.
- (299) Olk, P.; Buchler, B. C.; Sandoghdar, V.; Gaponik, N.; Eychmüller, A.; Rogach, A. L. Subwavelength Emitters in the Near-Infrared Based on Mercury Telluride Nanocrystals. *Appl. Phys. Lett.* **2004**, *84*, 4732–4734.
- (300) Shopova, S. I.; Farca, G.; Rosenberger, A. T.; Wickramanayake, W. M. S.; Kotov, N. A. Microsphere Whispering-Gallery-Mode Laser Using HgTe Quantum Dots. *Appl. Phys. Lett.* **2004**, *85*, 6101–6103.
- (301) Guyot-Sionnest, P.; Roberts, J. A. Background Limited Mid-Infrared Photodetection with Photovoltaic HgTe Colloidal Quantum Dots. *Appl. Phys. Lett.* **2015**, *107*, 253104.
- (302) Jagtap, A.; Goubet, N.; Livache, C.; Chu, A.; Martinez, B.; Gréboval, C.; Qu, J.; Dandeu, E.; Becerra, L.; Witkowski, N.; et al. Short Wave Infrared Devices Based on HgTe Nanocrystals with Air Stable Performances. *J. Phys. Chem. C* **2018**, *122*, 14979–14985.
- (303) Ackerman, M. M.; Tang, X.; Guyot-Sionnest, P. Fast and Sensitive Colloidal Quantum Dot Mid-Wave Infrared Photodetectors. *ACS Nano* **2018**, *12*, 7264–7271.
- (304) Tang, X.; Ackerman, M. M.; Shen, G.; Guyot-Sionnest, P. Towards Infrared Electronic Eyes: Flexible Colloidal Quantum Dot Photovoltaic Detectors Enhanced by Resonant Cavity. *Small* **2019**, *15*, 1804920.
- (305) Pradhan, S.; Di Stasio, F.; Bi, Y.; Gupta, S.; Christodoulou, S.; Stavrinidis, A.; Konstantatos, G. High-Efficiency Colloidal Quantum Dot Infrared Light-Emitting Diodes via Engineering at the Supra-Nanocrystalline Level. *Nat. Nanotechnol.* **2019**, *14*, 72–79.
- (306) Chuang, C.-H. M.; Brown, P. R.; Bulović, V.; Bawendi, M. G. Improved Performance and Stability in Quantum Dot Solar Cells through Band Alignment Engineering. *Nat. Mater.* **2014**, *13*, 796–801.
- (307) Roither, J.; Kovalenko, M. V.; Pichler, S.; Schwarzl, T.; Heiss, W. Nanocrystal-Based Microcavity Light-Emitting Devices Operating in the Telecommunication Wavelength Range. *Appl. Phys. Lett.* **2005**, *86*, 241104.
- (308) Wang, C.; Roither, J.; Kirschschlager, R.; Kovalenko, M. V.; Brehm, M.; Fromherz, T.; Kan, Q.; Tan, P.; Liu, J.; Chen, H.; et al. Enhanced Infrared Emission from Colloidal HgTe Nanocrystal Quantum Dots on Silicon-on-Insulator Photonic Crystals. *Appl. Phys. Lett.* **2009**, *95*, 053107.
- (309) Chen, T.; Reich, K. V.; Kramer, N. J.; Fu, H.; Kortshagen, U. R.; Shklovskii, B. I. Metal–Insulator Transition in Films of Doped Semiconductor Nanocrystals. *Nat. Mater.* **2016**, *15*, 299–303.
- (310) Fu, H.; Reich, K. V.; Shklovskii, B. I. Hopping Conductivity and Insulator-Metal Transition in Films of Touching Semiconductor Nanocrystals. *Phys. Rev. B: Condens. Matter Mater. Phys.* **2016**, *93*, 125430.
- (311) Mott, N. F. Metal-Insulator Transition. *Rev. Mod. Phys.* **1968**, *40*, 677–683.
- (312) Moreira, H.; Yu, Q.; Nadal, B.; Bresson, B.; Rosticher, M.; Lequeux, N.; Zimmers, A.; Aubin, H. Electron Cotunneling Transport in Gold Nanocrystal Arrays. *Phys. Rev. Lett.* **2011**, *107*, 176803.
- (313) Liu, Y.; Gibbs, M.; Puthussery, J.; Gaik, S.; Ihly, R.; Hillhouse, H. W.; Law, M. Dependence of Carrier Mobility on Nanocrystal Size and Ligand Length in PbSe Nanocrystal Solids. *Nano Lett.* **2010**, *10*, 1960–1969.
- (314) Kim, H.; Cho, K.; Kim, D.-W.; Lee, H.-R.; Kim, S. Bottom- and Top-Gate Field-Effect Thin-Film Transistors with p Channels of Sintered HgTe Nanocrystals. *Appl. Phys. Lett.* **2006**, *89*, 173107.
- (315) Kim, D.-W.; Jang, J.; Kim, H.; Cho, K.; Kim, S. Electrical Characteristics of HgTe Nanocrystal-Based Thin Film Transistors Fabricated on Flexible Plastic Substrates. *Thin Solid Films* **2008**, *516*, 7715–7719.
- (316) Lan, X.; Chen, M.; Hudson, M. H.; Kamysbayev, V.; Wang, Y.; Guyot-Sionnest, P.; Talapin, D. V. Quantum Dot Solids Showing State-Resolved Band-like Transport. *Nat. Mater.* **2020**, *19*, 323–329.
- (317) Chen, M.; Shen, G.; Guyot-Sionnest, P. State-Resolved Mobility of $1 \text{ cm}^2/(\text{Vs})$ with HgSe Quantum Dot Films. *J. Phys. Chem. Lett.* **2020**, *11*, 2303–2307.
- (318) Gréboval, C.; Noumbe, U.; Goubet, N.; Livache, C.; Ramade, J.; Qu, J.; Chu, A.; Martinez, B.; Prado, Y.; Ithurria, S.; et al. Field-Effect Transistor and Photo-Transistor of Narrow-Band-Gap Nanocrystal Arrays Using Ionic Glasses. *Nano Lett.* **2019**, *19*, 3981–3986.
- (319) Kovalenko, M. V.; Schaller, R. D.; Jarzab, D.; Loi, M. A.; Talapin, D. V. Inorganically Functionalized PbS–CdS Colloidal Nanocrystals: Integration into Amorphous Chalcogenide Glass and Luminescent Properties. *J. Am. Chem. Soc.* **2012**, *134*, 2457–2460.
- (320) Yakunin, S.; Dirin, D. N.; Protesescu, L.; Sytnyk, M.; Tollabimazraehno, S.; Humer, M.; Hackl, F.; Fromherz, T.; Bodnarchuk, M. I.; Kovalenko, M. V.; et al. High Infrared Photoconductivity in Films of Arsenic-Sulfide-Encapsulated Lead-Sulfide Nanocrystals. *ACS Nano* **2014**, *8*, 12883–12894.
- (321) Lambert, K.; Dendooven, J.; Detavernier, C.; Hens, Z. Embedding Quantum Dot Monolayers in Al_2O_3 Using Atomic Layer Deposition. *Chem. Mater.* **2011**, *23*, 126–128.
- (322) Devloo-Casier, K.; Geiregat, P.; Ludwig, K. F.; van Stiphout, K.; Vantomme, A.; Hens, Z.; Detavernier, C.; Dendooven, J. A Case Study of ALD Encapsulation of Quantum Dots: Embedding Supported CdSe/CdS/ZnS Quantum Dots in a ZnO Matrix. *J. Phys. Chem. C* **2016**, *120*, 18039–18045.
- (323) Pourret, A.; Guyot-Sionnest, P.; Elam, J. W. Atomic Layer Deposition of ZnO in Quantum Dot Thin Films. *Adv. Mater.* **2009**, *21*, 232–235.
- (324) Liu, Y.; Tolentino, J.; Gibbs, M.; Ihly, R.; Perkins, C. L.; Liu, Y.; Crawford, N.; Hemminger, J. C.; Law, M. PbSe Quantum Dot Field-Effect Transistors with Air-Stable Electron Mobilities above $7 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. *Nano Lett.* **2013**, *13*, 1578–1587.
- (325) Lhuillier, E.; Keuleyan, S.; Guyot-Sionnest, P. Transport Properties of Mid-Infrared Colloidal Quantum Dot Films. In *Quantum Dots and Nanostructures: Synthesis, Characterization, and Modeling IX*; International Society for Optics and Photonics, 2012; Vol. 8271, p 827109.
- (326) Baibich, M. N.; Broto, J. M.; Fert, A.; Van Dau, F. N.; Petroff, F.; Etienne, P.; Creuzet, G.; Friederich, A.; Chazelas, J. Giant Magnetoresistance of (001)Fe/(001)Cr Magnetic Superlattices. *Phys. Rev. Lett.* **1988**, *61*, 2472–2475.
- (327) Cryer, M. E.; Fiedler, H.; Halpert, J. E. Photo-Electrosensitive Memristor Using Oxygen Doping in HgTe Nanocrystal Films. *ACS Appl. Mater. Interfaces* **2018**, *10*, 18927–18934.
- (328) Jang, J.; Liu, W.; Son, J. S.; Talapin, D. V. Temperature-Dependent Hall and Field-Effect Mobility in Strongly Coupled All-Inorganic Nanocrystal Arrays. *Nano Lett.* **2014**, *14*, 653–662.
- (329) Yun, J.; Cho, K.; Kim, S. Flexible Logic Circuits Composed of Chalcogenide-Nanocrystal-Based Thin Film Transistors. *Nanotechnology* **2010**, *21*, 235204.
- (330) Chen, M.; Lu, H.; Abdelazim, N. M.; Zhu, Y.; Wang, Z.; Ren, W.; Kershaw, S. V.; Rogach, A. L.; Zhao, N. Mercury Telluride Quantum Dot Based Phototransistor Enabling High-Sensitivity Room-Temperature Photodetection at 2000 nm. *ACS Nano* **2017**, *11*, 5614–5622.
- (331) Kang, M. S.; Sahu, A.; Norris, D. J.; Frisbie, C. D. Size-Dependent Electrical Transport in CdSe Nanocrystal Thin Films. *Nano Lett.* **2010**, *10*, 3727–3732.

- (332) Wang, C.; Shim, M.; Guyot-Sionnest, P. Electrochromic Nanocrystal Quantum Dots. *Science* **2001**, *291*, 2390–2392.
- (333) Vanmaekelbergh, D.; Liljeroth, P. Electron-Conducting Quantum Dot Solids: Novel Materials Based on Colloidal Semiconductor Nanocrystals. *Chem. Soc. Rev.* **2005**, *34*, 299–312.
- (334) Puntambekar, A.; Wang, Q.; Miller, L.; Smieszek, N.; Chakrapani, V. Electrochemical Charging of CdSe Quantum Dots: Effects of Adsorption versus Intercalation. *ACS Nano* **2016**, *10*, 10988–10999.
- (335) Biscaras, J.; Chen, Z.; Paradisi, A.; Shukla, A. Onset of Two-Dimensional Superconductivity in Space Charge Doped Few-Layer Molybdenum Disulfide. *Nat. Commun.* **2015**, *6*, 8826.
- (336) Wu, C.-L.; Yuan, H.; Li, Y.; Gong, Y.; Hwang, H. Y.; Cui, Y. Gate-Induced Metal–Insulator Transition in MoS₂ by Solid Superionic Conductor LaF₃. *Nano Lett.* **2018**, *18*, 2387–2392.
- (337) Rastogi, P.; Chu, A.; Greboval, C.; Qu, J.; Noumbe, U. N.; Chee, S.-S.; Goyal, M.; Khalili, A.; Xu, X. Z.; Cruguel, H.; et al. Pushing Absorption of Perovskite Nanocrystals into the Infrared. *Nano Lett.* **2020**, *20*, 3999–4006.
- (338) Noumbé, U. N.; Gréboval, C.; Livache, C.; Brûlé, T.; Doudin, B.; Ouerghi, A.; Lhuillier, E.; Dayen, J.-F. Ionic Glass–Gated 2D Material–Based Phototransistor: MoSe₂ over LaF₃ as Case Study. *Adv. Funct. Mater.* **2019**, *29*, 1902723.
- (339) Gréboval, C.; Chu, A.; Magalhaes, D. V.; Ramade, J.; Qu, J.; Rastogi, P.; Khalili, A.; Chee, S.-S.; Aubin, A.; Vincent, G.; et al. Ferroelectric Gating of Narrow Band-Gap Nanocrystal Arrays with Enhanced Light Matter Coupling. *ACS Photonics* **2021**, *8*, 259–268.
- (340) Biscaras, J.; Hurand, S.; Feuillet-Palma, C.; Rastogi, A.; Budhani, R. C.; Reyren, N.; Lesne, E.; Lesueur, J.; Bergeal, N. Limit of the Electrostatic Doping in Two-Dimensional Electron Gases of La_xO₃ (X = Al, Ti)/SrTiO₃. *Sci. Rep.* **2015**, *4*, 6788.
- (341) Caviglia, A. D.; Gariglio, S.; Reyren, N.; Jaccard, D.; Schneider, T.; Gabay, M.; Thiel, S.; Hammerl, G.; Mannhart, J.; Triscone, J.-M. Electric Field Control of the LaAlO₃/SrTiO₃ Interface Ground State. *Nature* **2008**, *456*, 624–627.
- (342) Biscaras, J.; Bergeal, N.; Hurand, S.; Grossetête, C.; Rastogi, A.; Budhani, R. C.; LeBoeuf, D.; Proust, C.; Lesueur, J.; Lesueur, Two-Dimensional Superconducting Phase in LaTiO₃/SrTiO₃ Heterostructures Induced by High-Mobility Carrier Doping. *Phys. Rev. Lett.* **2012**, *108*, 247004.
- (343) Yu, D.; Wang, C.; Guyot-Sionnest, P. N-Type Conducting CdSe Nanocrystal Solids. *Science* **2003**, *300*, 1277–1280.
- (344) Yu, Q.; Cui, L.; Lequeux, N.; Zimmers, A.; Ulysse, C.; Rebuttin, V.; Pinna, N.; Aubin, H. In-Vacuum Projection of Nanoparticles for On-Chip Tunneling Spectroscopy. *ACS Nano* **2013**, *7*, 1487–1494.
- (345) Yu, Q.; Mottaghizadeh, A.; Wang, H.; Ulysse, C.; Zimmers, A.; Rebuttin, V.; Pinna, N.; Aubin, H. Verwey Transition in Single Magnetite Nanoparticles. *Phys. Rev. B: Condens. Matter Mater. Phys.* **2014**, *90*, 075122.
- (346) Wang, H.; Lhuillier, E.; Yu, Q.; Mottaghizadeh, A.; Ulysse, C.; Zimmers, A.; Descamps-Mandine, A.; Dubertret, B.; Aubin, H. Effects of Electron-Phonon Interactions on the Electron Tunneling Spectrum of PbS Quantum Dots. *Phys. Rev. B: Condens. Matter Mater. Phys.* **2015**, *92*, 041403.
- (347) Gravrand, O.; Wlassow, J.; Bonnefond, L. A Calibration Method for the Measurement of IR Detector Spectral Responses Using a FTIR Spectrometer Equipped with a DTGS Reference Cell. In *High Energy, Optical, and Infrared Detectors for Astronomy VI*; International Society for Optics and Photonics, 2014; Vol. 9154, p 91542O.
- (348) Liu, H.; Lhuillier, E.; Guyot-Sionnest, P. 1/f Noise in Semiconductor and Metal Nanocrystal Solids. *J. Appl. Phys.* **2014**, *115*, 154309.
- (349) Lai, Y.; Li, H.; Kim, D. K.; Diroll, B. T.; Murray, C. B.; Kagan, C. R. Low-Frequency (1/f) Noise in Nanocrystal Field-Effect Transistors. *ACS Nano* **2014**, *8*, 9664–9672.
- (350) Hooge, F. N. 1/f Noise Sources. *IEEE Trans. Electron Devices* **1994**, *41*, 1926–1935.
- (351) Downs, C.; Vandervelde, T. E. Progress in Infrared Photodetectors Since 2000. *Sensors* **2013**, *13*, 5054–5098.
- (352) Konstantatos, G.; Sargent, E. H. PbS Colloidal Quantum Dot Photoconductive Photodetectors: Transport, Traps, and Gain. *Appl. Phys. Lett.* **2007**, *91*, 173505.
- (353) Prins, F.; Buscema, M.; Seldenthuis, J. S.; Etaki, S.; Buchs, G.; Barkelid, M.; Zwillaer, V.; Gao, Y.; Houtepen, A. J.; Siebbeles, L. D. A.; et al. Fast and Efficient Photodetection in Nanoscale Quantum-Dot Junctions. *Nano Lett.* **2012**, *12*, 5740–5743.
- (354) Dong, Y.; Chen, M.; Yiu, W. K.; Zhu, Q.; Zhou, G.; Kershaw, S. V.; Ke, N.; Wong, C. P.; Rogach, A. L.; Zhao, N. Solution Processed Hybrid Polymer: HgTe Quantum Dot Phototransistor with High Sensitivity and Fast Infrared Response up to 2400 nm at Room Temperature. *Adv. Sci.* **2020**, *7*, 2000068.
- (355) Schornbaum, J.; Zakharko, Y.; Held, M.; Thiemann, S.; Gannott, F.; Zaumseil, J. Light-Emitting Quantum Dot Transistors: Emission at High Charge Carrier Densities. *Nano Lett.* **2015**, *15*, 1822–1828.
- (356) Hotta, S.; Yamao, T.; Bisri, S. Z.; Takenobu, T.; Iwasa, Y. Organic Single-Crystal Light-Emitting Field-Effect Transistors. *J. Mater. Chem. C* **2014**, *2*, 965–980.
- (357) Shulga, A. G.; Kahmann, S.; Dirin, D. N.; Graf, A.; Zaumseil, J.; Kovalenko, M. V.; Loi, M. A. Electroluminescence Generation in PbS Quantum Dot Light-Emitting Field-Effect Transistors with Solid-State Gating. *ACS Nano* **2018**, *12*, 12805–12813.
- (358) Kahmann, S.; Shulga, A.; Loi, M. A. Quantum Dot Light-Emitting Transistors—Powerful Research Tools and Their Future Applications. *Adv. Funct. Mater.* **2020**, *30*, 1904174.
- (359) Bisri, S. Z.; Takenobu, T.; Sawabe, K.; Tsuda, S.; Yomogida, Y.; Yamao, T.; Hotta, S.; Adachi, C.; Iwasa, Y. p-i-n Homojunction in Organic Light-Emitting Transistors. *Adv. Mater.* **2011**, *23*, 2753–2758.
- (360) Liu, C.-F.; Liu, X.; Lai, W.-Y.; Huang, W. Organic Light-Emitting Field-Effect Transistors: Device Geometries and Fabrication Techniques. *Adv. Mater.* **2018**, *30*, 1802466.
- (361) He, P.; Jiang, C.; Lan, L.; Sun, S.; Li, Y.; Gao, P.; Zhang, P.; Dai, X.; Wang, J.; Peng, J.; et al. High-Performance, Solution-Processed Quantum Dot Light-Emitting Field-Effect Transistors with a Scandium-Incorporated Indium Oxide Semiconductor. *ACS Nano* **2018**, *12*, 4624–4629.
- (362) Kagan, C. R. Flexible Colloidal Nanocrystal Electronics. *Chem. Soc. Rev.* **2019**, *48*, 1626–1641.
- (363) Jang, J.; Cho, K.; Lee, S. H.; Kim, S. Transparent and Flexible Thin-Film Transistors with Channel Layers Composed of Sintered HgTe Nanocrystals. *Nanotechnology* **2008**, *19*, 015204.
- (364) Hao, Q.; Tang, X.; Cheng, Y.; Hu, Y. Development of Flexible and Curved Infrared Detectors with HgTe Colloidal Quantum Dots. *Infrared Phys. Technol.* **2020**, *108*, 103344.
- (365) Diroll, B. T.; Chen, M.; Corpceanu, I.; Williams, K. R.; Talapin, D. V.; Guyot-Sionnest, P.; Schaller, R. D. Polarized Near-Infrared Intersubband Absorptions in CdSe Colloidal Quantum Wells. *Nat. Commun.* **2019**, *10*, 4511.
- (366) Guyot-Sionnest, P.; Hines, M. A. Intraband Transitions in Semiconductor Nanocrystals. *Appl. Phys. Lett.* **1998**, *72*, 686–688.
- (367) Gendron, L.; Carras, M.; Huynh, A.; Ortiz, V.; Koeniguer, C.; Berger, V. Quantum Cascade Photodetector. *Appl. Phys. Lett.* **2004**, *85*, 2824–2826.
- (368) Lhuillier, E.; Ribet-Mohamed, I.; Rosencher, E.; Patriarche, G.; Buffaz, A.; Berger, V.; Carras, M. Interface Roughness Transport in Terahertz Quantum Cascade Detectors. *Appl. Phys. Lett.* **2010**, *96*, 061111.
- (369) Delga, A. Quantum Cascade Detectors: A Review. In *Mid-infrared Optoelectronics*; Tournié, E., Cerutti, L., Eds.; Woodhead Publishing Series in Electronic and Optical Materials; Woodhead Publishing, 2020; Chapter 8, pp 337–377.
- (370) Konstantatos, G.; Badioli, M.; Gaudreau, L.; Osmond, J.; Bernechea, M.; de Arquer, F. P. G.; Gatti, F.; Koppens, F. H. L. Hybrid Graphene–Quantum Dot Phototransistors with Ultrahigh Gain. *Nat. Nanotechnol.* **2012**, *7*, 363–368.
- (371) Sun, Z.; Liu, Z.; Li, J.; Tai, G.; Lau, S.-P.; Yan, F. Infrared Photodetectors Based on CVD-Grown Graphene and PbS Quantum Dots with Ultrahigh Responsivity. *Adv. Mater.* **2012**, *24*, 5878–5883.

- (372) Jariwala, D.; Marks, T. J.; Hersam, M. C. Mixed-Dimensional van Der Waals Heterostructures. *Nat. Mater.* **2017**, *16*, 170–181.
- (373) Grotevent, M. J.; Hail, C. U.; Yakunin, S.; Dirin, D. N.; Thodkar, K.; Borin Barin, G.; Guyot-Sionnest, P.; Calame, M.; Poulikakos, D.; Kovalenko, M. V.; Shorubalko, I.; et al. Nanoprinted Quantum Dot–Graphene Photodetectors. *Adv. Opt. Mater.* **2019**, *7*, 1900019.
- (374) Robin, A.; Lhuillier, E.; Xu, X. Z.; Ithurria, S.; Aubin, H.; Ouerghi, A.; Dubertret, B. Engineering the Charge Transfer in All 2D Graphene-Nanoplatelets Heterostructure Photodetectors. *Sci. Rep.* **2016**, *6*, 24909.
- (375) Tang, X.; Lai, K. W. C. Graphene/HgTe Quantum-Dot Photodetectors with Gate-Tunable Infrared Response. *ACS Appl. Nano Mater.* **2019**, *2*, 6701–6706.
- (376) Tang, X.; Wu, G. F.; Lai, K. W. C. Twisted Graphene-Assisted Photocarrier Transfer from HgSe Colloidal Quantum Dots into Silicon with Enhanced Collection and Transport Efficiency. *Appl. Phys. Lett.* **2017**, *110*, 241104.
- (377) Huo, N.; Gupta, S.; Konstantatos, G. MoS₂–HgTe Quantum Dot Hybrid Photodetectors beyond 2 μm. *Adv. Mater.* **2017**, *29*, 1606576.
- (378) Balandin, A. A. Low-Frequency 1/f Noise in Graphene Devices. *Nat. Nanotechnol.* **2013**, *8*, 549–555.
- (379) Nam, M.; Kim, S.; Kim, S.; Jeong, S.; Kim, S. W.; Lee, K. Near-Infrared-Sensitive Bulk Heterojunction Solar Cells Using Nanostructured Hybrid Composites of HgTe Quantum Dots and a Low-Bandgap Polymer. *Sol. Energy Mater. Sol. Cells* **2014**, *126*, 163–169.
- (380) Im, S. H.; Kim, H.; Kim, S. W.; Kim, S.-W.; Seok, S. Il. Efficient HgTe Colloidal Quantum Dot-Sensitized near-Infrared Photovoltaic Cells. *Nanoscale* **2012**, *4*, 1581.
- (381) Sun, L.; Shen, L.; Meng, F.; Xu, P.; Guo, W.; Ruan, S. The Short Circuit Current Improvement in P3HT:PCBM Based Polymer Solar Cell by Introducing PSBTBT as Additional Electron Donor. *J. Nanosci. Nanotechnol.* **2014**, *14*, 3446–3449.
- (382) Grätzel, M. Dye-Sensitized Solar Cells. *J. Photochem. Photobiol., C* **2003**, *4*, 145–153.
- (383) Baxter, J. B.; Aydil, E. S. Nanowire-Based Dye-Sensitized Solar Cells. *Appl. Phys. Lett.* **2005**, *86*, 053114.
- (384) Savich, G. R.; Pedrazzani, J. R.; Sidor, D. E.; Maimon, S.; Wicks, G. W. Dark Current Filtering in Unipolar Barrier Infrared Detectors. *Appl. Phys. Lett.* **2011**, *99*, 121112.
- (385) Savich, G. R.; Pedrazzani, J. R.; Sidor, D. E.; Wicks, G. W. Benefits and Limitations of Unipolar Barriers in Infrared Photodetectors. *Infrared Phys. Technol.* **2013**, *59*, 152–155.
- (386) Dolzhnikov, D. S.; Zhang, H.; Jang, J.; Son, J. S.; Panthani, M. G.; Shibata, T.; Chattopadhyay, S.; Talapin, D. V. Composition-Matched Molecular “Solders” for Semiconductors. *Science* **2015**, *347*, 425–428.
- (387) Palaferri, D.; Todorov, Y.; Bigoli, A.; Mottaghizadeh, A.; Gacemi, D.; Calabrese, A.; Vasanello, A.; Li, L.; Davies, A. G.; Linfield, E. H.; et al. Room-Temperature Nine-Mm-Wavelength Photodetectors and GHz-Frequency Heterodyne Receivers. *Nature* **2018**, *556*, 85–88.
- (388) Zhu, B.; Chen, M.; Zhu, Q.; Zhou, G.; Abdelazim, N. M.; Zhou, W.; Kershaw, S. V.; Rogach, A. L.; Zhao, N.; Tsang, H. K. Integrated Plasmonic Infrared Photodetector Based on Colloidal HgTe Quantum Dots. *Adv. Mater. Technol.* **2019**, *4*, 1900354.
- (389) Chen, M.; Shao, L.; Kershaw, S. V.; Yu, H.; Wang, J.; Rogach, A. L.; Zhao, N. Photocurrent Enhancement of HgTe Quantum Dot Photodiodes by Plasmonic Gold Nanorod Structures. *ACS Nano* **2014**, *8*, 8208–8216.
- (390) Tang, X.; Chen, M.; Ackerman, M. M.; Melnychuk, C.; Guyot-Sionnest, P. Direct Imprinting of Quasi-3D Nanophotonic Structures into Colloidal Quantum-Dot Devices. *Adv. Mater.* **2020**, *32*, 1906590.
- (391) Chen, H.-L.; Cattoni, A.; De Lépinau, R.; Walker, A. W.; Höhn, O.; Lackner, D.; Siefer, G.; Faustini, M.; Vandamme, N.; Goffard, J.; et al. A 19.9%-Efficient Ultrathin Solar Cell Based on a 205-nm-Thick GaAs Absorber and a Silver Nanostructured Back Mirror. *Nat. Energy* **2019**, *4*, 761–767.
- (392) Rogalski, A. Dual-Band Infrared Detectors. In *Photodetectors: Materials and Devices V*; International Society for Optics and Photonics, 2000; Vol. 3948, pp 17–30.
- (393) Choi, J. J.; Wenger, W. N.; Hoffman, R. S.; Lim, Y.-F.; Luria, J.; Jasieniak, J.; Marohn, J. A.; Hanrath, T. Solution-Processed Nanocrystal Quantum Dot Tandem Solar Cells. *Adv. Mater.* **2011**, *23*, 3144–3148.
- (394) Péré-Laperne, N.; Rubaldo, L.; Kerlain, A.; Carrère, E.; Dargent, L.; Taalat, R.; Berthoz, J. 10 μm Pitch Design of HgCdTe Diode Array in Sofradir. In *Quantum Sensing and Nanophotonic Devices XII*; International Society for Optics and Photonics, 2015; Vol. 9370, p 937022.
- (395) Livache, C.; Martinez, B.; Goubet, N.; Ramade, J.; Lhuillier, E. Road Map for Nanocrystal Based Infrared Photodetectors. *Front. Chem.* **2018**, *6*, 575.
- (396) Na, J.; Park, K.; Kim, J. T.; Choi, W. K.; Song, Y.-W. Air-Stable Few-Layer Black Phosphorus Phototransistor for near-Infrared Detection. *Nanotechnology* **2017**, *28*, 085201.
- (397) Amani, M.; Regan, E.; Bullock, J.; Ahn, G. H.; Javey, A. Mid-Wave Infrared Photoconductors Based on Black Phosphorus-Arsenic Alloys. *ACS Nano* **2017**, *11*, 11724–11731.
- (398) Huang, L.; Tan, W. C.; Wang, L.; Dong, B.; Lee, C.; Ang, K.-W. Infrared Black Phosphorus Phototransistor with Tunable Responsivity and Low Noise Equivalent Power. *ACS Appl. Mater. Interfaces* **2017**, *9*, 36130–36136.
- (399) Suess, R. J.; Leong, E.; Garrett, J. L.; Zhou, T.; Salem, R.; Munday, J. N.; Murphy, T. E.; Mittendorff, M. Mid-Infrared Time-Resolved Photoconduction in Black Phosphorus. *2D Mater.* **2016**, *3*, 041006.
- (400) Guo, Q.; Pospischil, A.; Bhuiyan, M.; Jiang, H.; Tian, H.; Farmer, D.; Deng, B.; Li, C.; Han, S.-J.; Wang, H.; et al. Black Phosphorus Mid-Infrared Photodetectors with High Gain. *Nano Lett.* **2016**, *16*, 4648–4655.
- (401) Wu, D.; Wang, Y.; Zeng, L.; Jia, C.; Wu, E.; Xu, T.; Shi, Z.; Tian, Y.; Li, X.; Tsang, Y. H. Design of 2D Layered PtSe₂ Heterojunction for the High-Performance, Room-Temperature, Broadband, Infrared Photodetector. *ACS Photonics* **2018**, *5*, 3820–3827.
- (402) Yu, X.; Yu, P.; Wu, D.; Singh, B.; Zeng, Q.; Lin, H.; Zhou, W.; Lin, J.; Suenaga, K.; Liu, Z.; Wang, Q. J. Atomically Thin Noble Metal Dichalcogenide: A Broadband Mid-Infrared Semiconductor. *Nat. Commun.* **2018**, *9*, 1545.
- (403) Long, M.; Wang, Y.; Wang, P.; Zhou, X.; Xia, H.; Luo, C.; Huang, S.; Zhang, G.; Yan, H.; Fan, Z.; et al. Palladium Diselenide Long-Wavelength Infrared Photodetector with High Sensitivity and Stability. *ACS Nano* **2019**, *13*, 2511–2519.
- (404) Luo, L.-B.; Wang, D.; Xie, C.; Hu, J.-G.; Zhao, X.-Y.; Liang, F.-X. PdSe₂ Multilayer on Germanium Nanocones Array with Light Trapping Effect for Sensitive Infrared Photodetector and Image Sensing Application. *Adv. Funct. Mater.* **2019**, *29*, 1900849.
- (405) Sahu, A.; Khare, A.; Deng, D. D.; Norris, D. J. Quantum Confinement in Silver Selenide Semiconductor Nanocrystals. *Chem. Commun.* **2012**, *48*, S458–S460.
- (406) Sahu, A.; Qi, L.; Kang, M. S.; Deng, D.; Norris, D. J. Facile Synthesis of Silver Chalcogenide (Ag₂E; E = Se, S, Te) Semiconductor Nanocrystals. *J. Am. Chem. Soc.* **2011**, *133*, 6509–6512.
- (407) Richner, P.; Galliker, P.; Lendenmann, T.; Kress, S. J. P.; Kim, D. K.; Norris, D. J.; Poulikakos, D. Full-Spectrum Flexible Color Printing at the Diffraction Limit. *ACS Photonics* **2016**, *3*, 754–757.
- (408) Richner, P.; Kress, S. J. P.; Norris, D. J.; Poulikakos, D. Charge Effects and Nanoparticle Pattern Formation in Electrohydrodynamic NanoDrip Printing of Colloids. *Nanoscale* **2016**, *8*, 6028–6034.
- (409) Rinnerbauer, V.; Kovalenko, M.; Lavchiv, V.; Kocher, G.; Roither, J.; Heiss, W.; Hingerl, K. Spectroscopic Ellipsometry of Layer by Layer Deposited Colloidal HgTe Nanocrystals Exhibiting Quantum Confinement. *Phys. E* **2006**, *32*, 104–107.
- (410) Rinnerbauer, V.; Hingerl, K.; Kovalenko, M.; Heiss, W. Quantum Confinement in Layer-by-Layer Deposited Colloidal HgTe Nanocrystals Determined by Spectroscopic Ellipsometry. *Appl. Surf. Sci.* **2007**, *254*, 291–294.