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Low-dimensional II–VI semiconductor nanostructures of ternary alloys and transition metal ion doping: synthesis, optical properties and applications

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Light matter interactions in spin-controlled devices, which are also known as dilute magnetic semiconductors, have gained significant attention in the past few years. The strong electron–phonon interaction and outstanding excitonic behavior of II–VI semiconductors are advantages over III–V semiconductors, especially in low-dimensional systems. Herein, we comprehensively review the spin-exchange interactions induced by transition metal ion doping through sp–d exchange, which enhances the possibility of spin-based photonics in binary and ternary alloys and transition metal ion-doped II–VI semiconductors. For one-dimensional structures, few magnetic ions can ferromagnetically couple to form exciton magnetic polarons, resulting in coherent spin-polarized emission, which opens a new horizon for the development of tunable bandgaps, active waveguides, tunable emissions and lasing at the nanoscale based on different structures, compositions, morphologies and excitation conditions. Photonic crystals supply significant information about photon–exciton interactions in coupled optical microcavities. Lastly, we extend the discussion to the most recent advancements of one-dimensional structures for device applications and their possible prospectives in future technology.

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1. Introduction

Incorporation of transition metal (TM) ions in II–VI semiconductors and their ternary alloys induces spin-carrier injection and spin–spin coupling due to their p–d hybridization (involving anions and TM ions) and has been extensively used for dilute magnetic semiconductor (DMS) applications. DMSs are an emerging class of semiconductors that exhibit both semiconductor behavior and ferromagnetic response. For their practical application, DMSs must show ferromagnetic response at room or above room temperature according to the theoretical prediction of ferromagnetic display in wide bandgap semiconductors by Dielt *et al.*¹ Extensive research on II–VI-based DMSs over the last two decades has led to the construction of DMS-incorporated structures such as quantum wells and superlattices. The exchange interaction between the (s, p) bands of the host lattice and d-band of magnetic ions is responsible for the

magneto-optical and magneto-electrical effect at room temperature. The d–d transition peaks of transition metal ions in II–VI semiconductors are influenced not only by the exciton recombination process but also by spin-related couplings.² Based on the recent advancements in DMSs, antiferromagnetic (AFM) coupled ions have been observed at room temperature, which show the d–d intra-band transition peaks at a higher energy level than FM-coupled ions. Spin-related micro-photoluminescence (PL) response and theoretical studies revealed a red-shift in the d–d transition peak of the Mn(II) ions in FM systems, enabling bosonic lasers and other spintronic applications. Polarons are quasi-particles that interact in the quantum state coupled with the spin of localized magnetic ions in DMSs, which are called magnetic polarons. Magnetic polarons are subdivided into bound magnetic polarons (BMPs), free magnetic polarons (FMPs) and excitonic magnetic polarons (EMPs). EMPs are self-organized magnetic quasi-particles formed by excitons in DMSs. At high excitation power, EMPs merge due to the overlapping tails of their carrier wave function and form a long cluster of collective magnetic polarons, creating a large ferromagnetic domain. The large collective response of the exciton magnetic polaron state pronounced the bosonic behavior.^{3–5} Spintronics is an advanced device technology that utilizes quantum spin states of electrons and controlled charge states (spin up and spin down) for the fabrication of magnetic memory devices or switching applications. In this case, the integration of electronics,

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magnetism and spin in single devices is an extremely important research topic for the development of spintronic devices. An important task related to spintronic devices such as MRAM, spin LED and spin FET is to effectively and efficiently enable the spin injection within materials to control the spin transport and detection within semiconductor devices.^{6–8}

II–VI semiconductors especially zinc chalcogenides (ZnO, ZnS, ZnSe, and ZnTe) and cadmium chalcogenides (CdS, CdSe and CdTe) have been intensively studied in the field of one-dimensional (1D) nanostructures due to their superior optical properties.^{9–12} Particularly, 1D nanostructures with a size in the range of a few nanometers (lateral dimensions between 1 to 100 nm) have attracted significant attention over the past three decades due to their numerous fascinating properties, which can be tuned by changing their shape, size and growth conditions.^{13–16} Accordingly, it is important to investigate the various optical phenomena such as electron–exciton interaction, electron–phonon interaction, and other emissions such as donor–acceptor pair exciton and anti-site emission in 1D nanostructures to understand their crystal quality and sensitivity to light. Zinc and cadmium chalcogenides possess bandgaps in the spectral range of 2.25 eV to 3.37 eV (visible to UV) and 1.4 eV to 2.42 eV (NIR to visible), respectively, and their direct bandgap together with quantum confinement make them suitable for future optoelectronic applications. Furthermore, the bandgap of ternary alloys of II–VI semiconductors can be tuned based on their stoichiometric ratio, resulting in a response to multi-spectrum wavelengths. Thus, utilizing the quantum confinement effect in 1D nanostructures of ternary alloys has attracted significant attention for bandgap tuning in technologically motivated research based on photon absorption or emission, which are synthesized *via* physical deposition methods such as chemical vapor deposition.^{16–18}

1D structures (scaled down to the submicron range) have shown unique properties in extensive and novel applications. The 1D structure of nanowires (NWs) and carbon nanotubes (NTs) has attracted great interest in curiosity-driven and technology-motivated research, which were originally called “whiskers” but more recently known as nanobelts (NBs).¹⁹ As the size of structures is scaled down from the bulk to 1D (scaled down to submicron range), the exchange interaction between spin and carriers becomes predominant, and consequently their optical and electrical properties can be modified using an external magnetic field, which is commonly applied in spintronics, optical imaging and fields.²⁰ In the last decade, high-quality II–VI ternary alloy 1D nanostructures have been prepared for the fabrication of tunable nano-lasers, LEDs, solar cells and photodetectors with elaborate control and reproducibility achieved by various research groups.^{21–23} Therefore, the current progress in 1D nanostructure ternary alloys of II–VI semiconductors has strengthened the expectation of their pivotal role in the future energy conversion and biological applications (Fig. 1).^{16,24} Different prototypes have the capability to assemble and integrate individual structures into functional devices on a large scale. In this case, 1D hybrid nanostructures are highly appropriate in various optoelectronic applications such as type-I and type-II junctions.

Specifically, type-I junctions exhibit high fluorescence and light emission yield (LEDs and lasers), whereas type-II junctions facilitate the formation of e–h pairs with improved energy conversion efficiency.^{21,25}

In this review, we explore the curiosity-driven/achievement and technologically motivated research in the past decade related to 1D micro/nanostructures of II–VI semiconductors and their application in the device industry. In the introduction, we present a general background on 1D nanostructures related to II–VI semiconductors and their ternary alloys and TM doping in II–VI materials and their ternary alloys. The incorporation of TM results in spin-carrier injection, spin–spin coupling among TM ions and p–d hybridization, which endow direct semiconductors with multitudinous potential in photonic and optoelectronic applications. EMPs, BMP and localized EMPs are striking areas of research on DMS, which need to be explored in 1D systems. Finally, we discuss the future perspectives of 1D nanostructures of II–VI semiconductors and their ternary alloys with and without doping, and their potential for new device applications.

2. Synthesis of 1D nanostructures from II–VI semiconductors and their alloys

Various research groups have synthesized 1D nanostructure II–VI semiconductors employing different techniques. In this case, each synthesis mechanism is crucial for the growth of 1D nanostructures that have good reproducibility, crystallinity, and most importantly cost-effectiveness. Therefore, the investigation of the chemistry and physics of nanostructures can reasonably facilitate the control of the growth pattern, crystal structure and compositional complexity of materials for their practical applications toward novel technologies.^{26,27} Lieber and co-workers reported the application of NW in biochemical sensors, extracellular and intracellular electrical sensors, molecule delivery, injectable electronics, biosynthesis and optical neuromodulation for brain science. Moreover, the importance of NW structures such as axial, radial/coaxial modulated, branched and kink structures are next-generation building blocks for bioelectronics, photonics and electronic circuits. The high surface-to-volume ratio, fast signal response, high temporal resolution and minimum energy consumption of nanostructures highlight their importance for future applications.^{28–30}

II–VI 1D nanostructure ternary alloys of cadmium and zinc chalcogenides have been widely used in various applications due to their tunable bandgaps behavior and excellent transport properties. The bandgap of binary alloys can be tuned by changing their size; meanwhile, the bandgap of ternary alloys can be adjusted by changing their composition.^{31–33} TM ion doping in binary and ternary II–VI alloys can tune their bandgap together with magnetic ion inner conversion and energy transfer within the crystal symmetry of the host II–VI materials. Therefore, light-matter interaction provides fascinating information and tunable optical emission from ferromagnetic semiconductors, which usually arises from the carrier–spin interaction, spin–spin coupling and sp–d hybridization between the anions and dopant ions.^{5,34,35}



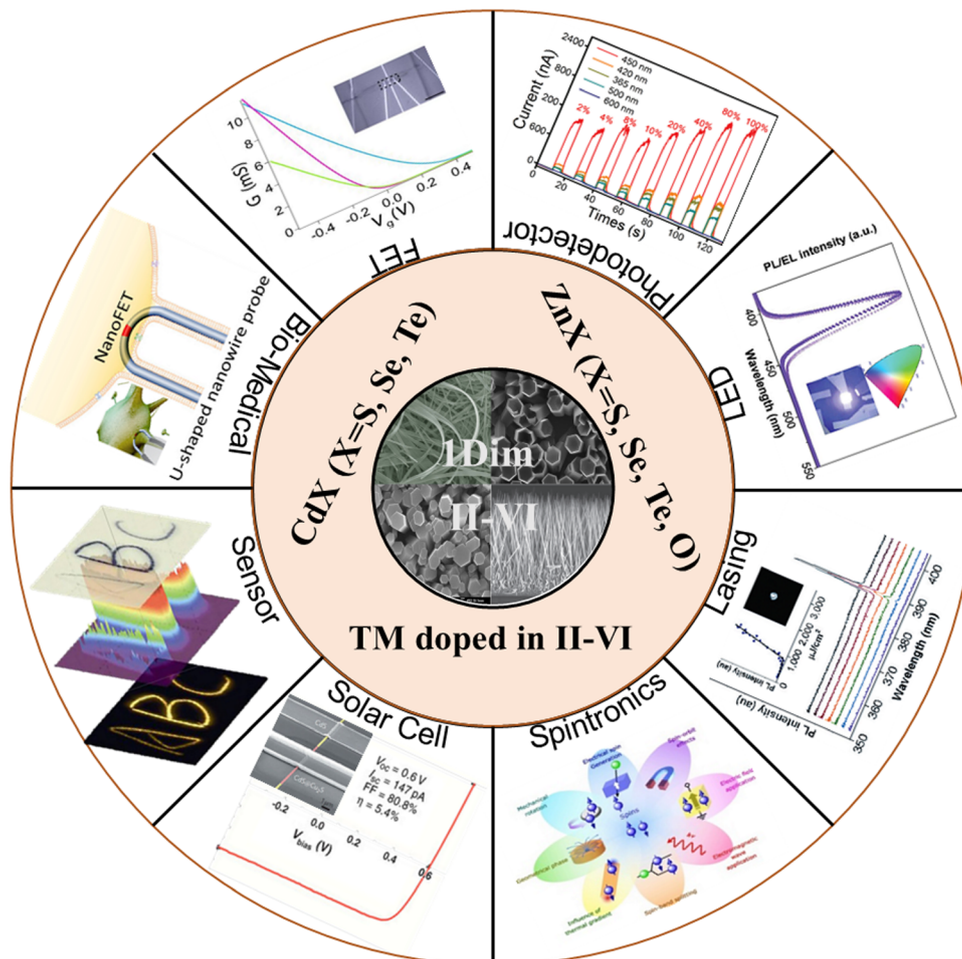


Fig. 1 Graphical display of II–VI semiconductors.

2.1 Growth mechanism

The synthesis of 1D nanostructures from II–VI semiconductors has been demonstrated by nucleation growth *via* a vapor phase mechanism and solution process. The merits of the vapor phase mechanism include the production of high-quality II–VI crystal materials due to the high-temperature growth process. Wagner and co-workers introduced the vapor–liquid–solid approach, which is currently established and widely employed to grow 1D nanostructures for different applications together with device integration.³⁶ Meanwhile, the solution process is a relatively cost-effective method to prepare 1D nanostructures at a lower temperature; however, it compromises their crystallinity. In this section, we present in detail the elaborate growth mechanisms and experimental techniques employed to prepare high-quality 1D structures from II–VI semiconductors and the recent developments in ternary alloys for the production of high-quality and critical stoichiometry materials for future application.^{37–39}

2.1.1 Vapor phase growth. Vapor phase growth involves two main mechanisms, as follows: (i) nucleation growth and (ii) absence of nucleation growth, which is also known as the vapor liquid solid (VLS) and vapor solid (VS) mechanism. In the case

of nucleation growth, a catalyst initializes the growth mechanism and acts as the seed during the growth, in which the host material is vaporized in the presence of a source gas and reacts with the catalyst (as the seed) to form a 1D nanostructure. In this case, the shape, size and morphology can be controlled by controlling the catalyst shape and size as a function of the temperature.^{39–41}

2.1.1.1 Vapor–liquid–solid (VLS) growth. VLS growth is the most extensively adopted process to synthesize high-crystalline, flexible and large-scale 1D nanostructures. The kinetics of VLS is based on the alloying formation of the catalyst and host material, leading to the formation of single or branched-shaped homo/heterogeneous nanostructures on a single substrate. However, controlling the kinetics of VLS is challenging under conventional conditions. Therefore, two-zone and three-zone temperature furnaces are employed to analyze the fast kinetics or *in situ* techniques such as transmission electron microscopy (TEM) to investigate the growth kinetics in real time.^{40–42} The shape, size, crystal structure and morphology of the desired material depend on the formation of a solid–liquid alloy or adsorption of the liquid–solid interface in supersaturated form,



which is affected by numerous parameters such as pressure, deposition temperature and time, carrier gas flow and catalyst droplet size.^{43,44} Moreover, the polarity of II–VI semiconductors can be determined by dumbbell analysis *via* high angle annular dark field (HAADF) scanning transmission electron microscopy (STEM) imaging for high-medium atomic number and low atomic number NWs obtained from the VLS mechanism. This change in polarity in NWs leads to twinning or polymorphism at elevated temperature. Therefore, HAADF and STEM provide suitable information about the complex hetero-structures and hetero-interfaces between polar and nonpolar semiconductors.^{45–47} Self-catalytic growth in VLS is a special case in which the self-constituent metallic element serves as the catalyst for the formation of the host compound, for example, Cd metallic particles are used to grow cadmium chalcogenides. However, Van der epitaxial growth in VLS also supports the self-catalytic growth of micro/nanostructures based on II–VI materials on a mica substrate, as reported by various groups. The growth of high-quality micro/nanostructures is directly correlated with the substrate temperature, source gas, flow rate, evaporation temperature of the host material and growth time.^{47–50}

2.1.1.2 Vapor–solid (VS) growth. The vapor–solid mechanism is a spontaneous condensation process without the use of a catalyst or metal droplet. The spontaneous condensation of vapor into a solid is induced by a decrease in the Gibbs free energy of crystallization or decrease in super-saturation.^{45,49} The reaction temperature in the VS mechanism is slightly higher than that in VLS, which accumulates anion–cation adatoms on the substrate. The quality of the nanostructures obtained *via* the VS mechanism directly depends on the growth temperature, time and super-saturation ratio to realize the high diffusivity of the adatoms. The VS mechanism is associated with the radial thickening (formation of NB) and tapering of NWs at high super-saturation at elevated temperature. In the case of II–VI semiconductors, there are numerous reports on the growth of wurtzite phase crystal structures as NBs under the appropriate ambient conditions.^{50–52}

2.1.2 Solution process. The solution process is also widely used to grow 1D colloidal NWs at low temperature in comparison with the VLS or VS mechanism. It exhibits numerous advantages over high temperature growth methods such as controlled diameter, surface passivation, solubility and large production. The diameter can be controlled in the solution process either with help of surface-capping ligands or surfactant-free method. In the surfactant method, monodisperse metal catalyst nanoparticles (NPs) having a diameter of a few nanometers help to grow colloidal NWs, which is protected by surface-capping ligands around the nanostructure. Alternatively, surfactant-free nanostructures are prepared in squalene (as the solvent) and semiconductor precursors, where the semiconductor precursor thermally decomposes into nanodroplets, subsequently assisting the growth of NWs.^{25,35} Similarly, organometallic (metal with a low melting point) precursors are dissolved in an organic solvent at elevated temperature and the formed liquid–metal droplets (few nanometers) act as a catalyst to induce the nucleation process, followed by the growth of 1D structures of II–VI ternary alloys.^{53–55}

2.1.2.1 Solution–liquid–solid (SLS) growth. SLS is used to synthesize crystalline structures below 200 °C in the presence of a catalyst (Bi, In, Sn and Ga). Metallic catalysts are used to prepare colloidal structures with various compositions and structures (such as axial heterojunction and core/shell). SLS growth is similar to VLS growth but differ by its low temperature in a controlled environment. In SLS, the influencing parameter is temperature, which controls the diameter of the rods and wires in the range of 2–20 nm diameter.^{25,34} Moreover, the preparation of various hetero-structures of type-I and type-II II–VI semiconductors has been reported using the solution growth mechanism. In the case of type-I (CdS/CdSe), excitons recombine in the core of CdSe and CdS, serving as a concentrator to improve the light efficiency. Alternatively, in type-II (CdSe/CdTe), they serve as two localized charge carriers, exhibiting Coulomb interaction and exchange interaction of interface-related exciton recombination.^{54,56–58}

2.2 Growth extension

2.2.1 Single nanowire array. Single NW arrays have been achieved by epitaxial growth on a crystalline substrate. The crystalline planes of the substrate effectively control the orientation of NWs on a large scale, which is very important for device fabrication. II–VI NWs grown epitaxially on the surface of a substrate with the lattice mismatch of less than 10% show a tremendous photonic response. The building blocks for the formation of NWs are the hetero-interface between the host material and catalyst particles, facilitating the fast growth of the structure perpendicular/vertical to the substrate surface together with large tolerance to lattice mismatch.⁵⁹ The epitaxial growth of ZnO NW arrays was first reported by Yang and co-workers, which have been extensively reported in a broad range of electronic and photonic applications.^{60,61} Moreover, the fast growth of NW arrays enable lattice relaxation in lateral dimensions, leading to the formation of a large critical height prior to defect nucleation, as reported for lattice mismatched epitaxial films. In this case, a high growth temperature is suitable for the growth of high-crystalline materials and epitaxial growth *via* the formation of covalent bonds between 1D structures and the substrate, thus requiring the corresponding in-plane parameters to overcome the crystalline defects. van der Waals interaction or epitaxial growth provides a strong chemical bond between heterojunctions with minimum impact on their lattice matching. Thus, various II–VI nanostructures have been successfully grown using the van der Waals epitaxial method, exhibiting strong luminescence behavior.⁴⁶

2.2.2 Heterostructure nanowires. Heterostructure nanostructures are classified as (i) axial and (ii) radial structures. Axial heterostructures have two different materials with the same diameter, whereas radial structures have two different materials with different diameters, which are also known as core/shell nanostructures and first reported in 2002.^{60,61} Heterostructure or heterojunction NWs have various advantages in bioelectronics, solar cells and photonic devices. Lieber and co-workers also reported that the formation of coaxial multishell NW materials depends on the degree of freedom (DoF) of their synthetic modulation and tuned properties, leading to advances in nanophotonics.^{62–64} Moreover, branched nanostructures with



heterogeneous or homogeneous junctions possess a more complex design but open a new horizon in various energy-related applications. Branched NWs with homo or heterogeneous junctions provide a higher degree of complexity to facilitate the design of interconnected hierarchical nanostructures with rich electrical, optical, and chemical properties, especially in energy applications.^{65–67} Wang and co-worker reported the synthesis of heterojunction/hyper-branched Si and GaN NWs.⁶⁸ Dai and co-worker reported the preparation of CdSe/CdS core/shell NWs *via* a two-step process and CVD.⁶⁹ Pan reported the one-step thermal evaporation of multiple sources to synthesize Si/CdS_xSe_{1-x} core/shell NWs.^{70–73} Moreover, the composition of the CdS and CdSe precursors in 1D nanostructure CdS_xSe_{1-x} NWs can alter their length *via* CVD together with composition grading and formation of lateral and branched heterostructures of CdS_xSe_{1-x}. In addition, Buhro reported the growth of ZnSe_xTe_{1-x} NWs by a solution process with a composition gradient. Specifically, they employed a three-segment heterostructure having ZnSe and ZnTe localized at the terminal ends of NWs for the growth of the ZnSe_xTe_{1-x} NWs.⁷⁴

2.2.3 Composition gradient/transition metal ion doping.

The composition gradient in 1D II–VI semiconductors exhibits a broad range of tunable emissions in a single structure, which is difficult to attain *via* planar epitaxial growth. The axial direction growth of NWs is important inducing a bandgap gradient *via* the vapor phase growth mechanism.⁷⁵ The composition gradient and doping profile can be controlled by the concentration or stoichiometry ratio of the precursors and substrate temperature zone, as reported by Zhang and others.^{75–78} Similarly, TM ion doping in II–VI materials make them suitable for future applications such as spintronics.^{79–82} The origin of ferromagnetism in semiconductors especially II–VI semiconductors is related to the spin exchange interaction between TM ion impurities and carriers available at the conduction or valence band through sp–d exchange interaction. The available electrons in the outer level (3d and 4s) of TMs are of great interest to explain the possible carrier–spin interaction. Electron spin is a quantum phenomenon coupled with the magnetic moment and degree of spin, which carries information regarding spin up or spin down. In Mn, its 3d⁵ shell is half-filled and 4s² electron bonds to s–p³. According to Hund's rule, the 5d-electrons have parallel spin and opposite spin, requiring a considerably large energy. Coulomb potential, carrier–spin (electron–spin) interaction and Mn–Mn (spin–spin) interaction are possible interactions reported in DMS by different groups. Zou and co-workers reported the doping of various II–VI materials 1D nanostructures with TM ions for DMS application, which will be discussed in detail in the next section.

3. Fundamental optical properties of II–VI nanostructures

A wide range of direct bandgap materials is classed as II–VI semiconductors, which normally exhibit high optical absorption and emissions within the visible range. Cadmium and zinc

chalcogenides are promising potential II–VI materials in the field of 1D nanostructures, which have been extensively studied over the last two decades for various optical applications due to their high quantum efficiency and light sensitivity. II–VI semiconductors possess wurtzite-type, zinc blende and diamond-like structures.^{83–85} In the case of wurtzite-type structures, their valence bands arise from p-states having lower symmetry than diamond and zinc blende like-structures. Inversely, the minimum of the conduction band is Γ_7 symmetry with an s-state. The valence band is split into three sub-bands due to the spin-orbit coupling and crystal field; A-exciton (Γ_9 or heavy hole), B-exciton (Γ_7 or light hole), and C-exciton (Γ_7 or split-off (SO) band). To investigate the optoelectronic or photoelectrical behavior of II–VI semiconductors, it is necessary to develop a deep understanding of the band structure-related exciton transitions. In this section, we highlight the recent progress in zinc and cadmium chalcogenide binary alloys, ternary alloys and TM ion doping in 1D nanostructure-related electronic and optical transitions.

3.1 TM ion doping in cadmium chalcogenides (CdX; X = S, Se, or Te)

3.1.1 CdS. Cadmium sulfide is an attractive II–VI semiconductor material with a bandgap of 2.4 eV in the visible range, and thus has been extensively used in various optoelectronic applications. The 1D nanostructure of CdS exhibits strong luminescence, high mobility and excellent transport behavior in comparison with its bulk counterpart.^{86–89} Given that II–VI compound semiconductors are traditionally optically active, the exploration of their luminescence properties is essential for various optical applications. Excitons are quasi-bosonic particles that freely move in the crystal lattice and describe the bound state of an electron–hole (e–h) pair with electrostatic Coulombic force interactions. Therefore, the carrier–carrier, carrier–exciton and carrier–phonon interaction exhibit superior features under an excitation wavelength.^{90–92} In comparison to their bulk counterparts, nanostructures have a relatively higher exciton–polariton coupling strength. Leite *et al.* reported that the temperature-dependent multiple phonons in CdS under different excitation powers exhibit strong electron–lattice interaction.⁹³ In addition, the typical luminescence process in ternary alloys and TM ion-doped II–VI materials provides suitable information to understand the transition state of electron relaxation to the ground state.^{94,95}

3.1.1.1 Transition metal ion-related luminescence behavior. TM doping in 1D CdS is quite interesting for DMS application. This is because the d–d transition between the ground-state ⁶A₁ and first-excited ⁴T₁ state of the ferromagnetic (FM) and anti-ferromagnetic (AFM) coupled spin ions in the host matrix make it useful in various spin-related applications.^{95–97} Ge *et al.* reported the coherent bosonic lasing of EMP at 520 nm at 232.393 $\mu\text{J cm}^{-2}$ for Ni-doped CdS synthesized by the VLS mechanism. 1D CdS with TM ion doping was excited at different wavelengths (405 nm and 532 nm) to understand the Mn²⁺ transition in the host CdS crystal symmetry.⁹⁸ The



EMP emission from the Ni(*n*)-doped CdS and Mn-doped CdS 1D nanostructures synthesized by the VLS mechanism has been reported by various groups. Moreover, Zhang *et al.* also reported that Sn-doped CdS exhibited a low threshold value for an NW laser due to plasmon polaritons.^{99,100} The formation of an LEMP emission in CdS is related to the formation of Ni clusters even at a low doping concentration. A high doping concentration (up to 2–4%) showed that the Ni₂ cluster emission in CdS is related to LEMP, which is consistent with the theoretical *ab initio* calculation using the VASP software. The localized and delocalized states interact with the d-bands, showing LEMP due to the delocalized charge transfer effect of coupled Ni ions surrounded by I and S ions. The LEMP decays *via* the efficient hopping mechanism due to the high Ni doping, exhibiting the corresponding behavior. Interestingly, the FM coupling of the Ni(*n*) pair has a stable energy of 121 meV, which is lower than that of the AFM Ni pair, indicating that the incorporation of the Ni ion in the host lattice modifies its band structure. According to the DOS, the d-band is occupied above the Fermi level, which is close to the conduction band minimum. The strong hybridization of the d-band with the valence band dominates the exciton and exciton–spin interaction. However, in the case of AFM, the metallic states of the d-band are dominant and distributed cross at the Fermi level. Another prominent effect known as the charge transfer effect has also been reported, where the spin-down state contacts with the continuum conduction band and the d-states can extend the band edge to a lower energy. The charge transfer effect in the continuum state combined with FM coupling will lead to a second band near the highest d-level, which is called bound EMP or BEMP. Therefore, the aggregation of Ni clusters and FM coupling of the d–d transition energy states can exhibit LEMP and BEMP in 1D NBs. The LEMP can form BEMP as FM ion coupled with TM aggregate and doping formed charge transfer at low energy side due to M–L cluster in CdS lattice to produce single-mode lasing and EMP lasing as excited by fs-laser. High excitation density of coherent exciton may lead to the formation of coherent EMP or CEMP for lasing. The spatial and energy difference between two EMPs and the BEMP (LEMP) level have a much longer lifetime, lowest energy and limited space. Therefore, the CBEMP state occurs at a smaller threshold to produce lasing with fs pulse pumping. When the EMP threshold value is achieved, EMP produces lasing through the formation of CEMP with a coherent emission. Therefore, the bosonic nature favors the formation of CEMP, LEMP and CBEMPs in the host lattice to produce the EMP condensate for single-mode lasing, which spontaneously coupled with spin will produce multi-exciton emission. This new concept related to exciton-magnetic polarons can tune the intrinsic microscopic interactions in strongly correlated compounds related to the size and spin ion-doping ratio in DMS. Therefore, the excitonic nature will produce multi-spin-coupled polaronic excitonic states, which may lead to new magnetic and/or spin-related optical properties.⁵

Fig. 2a shows the electronic band structure of pure CdS and Fig. 2b reveals the density of states for single Cd, S and CdS through the sp hybridization of the occupied and unoccupied

available states.^{95,101} Fig. 2c shows the Mn atom doping in CdS, where theoretical studies predict that one Mn replaces Cd due to the smaller ionic radius of the former, resulting in a spin distribution iso-sphere and spin polarization. The Mn-related d–d transition occurs when the TM ion is doped in the crystals and the magnetic coupling within the lattice influences its location. The binding energy of the Mn-3d state for the magnetic ion impurity in semiconductors can optimize the Hubbard parameter ($U = 2.8$ eV) for a $2 \times 2 \times 2$ supercell, showing a strong correlation with the experimental data for the d–d transition in Mn-doped CdS.^{102,103} Fig. 2c shows the spin-polarized electronic band structure of single Mn-doped CdS calculated along the high symmetry directions. In the case of two Mn ion contributions, leading to 5.6% dopant impurity in the supercell, the energy difference between the antiferromagnetic and ferromagnetic states is 26 meV. The simulation results show the favorable antiferromagnetic state for the long-range dopant in the CdS nanostructure. The optical absorption of the electronic transition for the occupied and unoccupied states is shown in Fig. 2d for pure and doped CdS. In the case of pure CdS, its absorption edge occurs at 2.43 eV, which is generated between S-2p and Cd-2s with the valence and conduction band. According to the comparison with Mn-doped CdS, the doped sample has a weak band edge peak intensity, which is shifted toward a lower energy value. In the case of single Mn, its related emission is located at 2.20 eV, which is close to the experimental value of 2.13 eV for the d–d transition, while the double Mn ion peak is located at 2.38 eV. The d–d transition of double Mn is larger than that of the single Mn ion owing to the AFM coupling of the Mn ions in the CdS system. Therefore, the d–d transition occurs between the ⁶A₁ ground state and first excited ⁴T₁ state for the FM and AFM coupled Mn ions. In the case of AFM coupling, one spin-up Mn ion interacts with the empty spin-up state, exhibiting a blue shift in comparison with FM coupling, as observed in the luminescence spectra of other II–VI semiconductors.^{103–106}

Fig. 2g shows the PL spectra of 1D belt-like structures of both undoped and Mn(*n*)-doped CdS and Mn(*n*)-doped synthesized by the VLS mechanism. The undoped CdS shows a strong near-band edge emission at 508 nm and no emission peak after 525 nm.^{5,102} In the case of the Mn-doped samples, their luminescence spectra show emission bands in the range of 570–900 nm, besides their bandgap emission, as shown in Fig. 2g. The appearance of multiple emission peaks in their emission spectra is due to the d–d transition in the ferromagnetic (MnS)_n cluster. The emission peaks at 510 nm and 518 nm in the doped sample are due to EMP, as also reported for a nanocrystal (NC) by different research groups.^{104–106} Therefore, the tunable emissions with different Mn ions in 1D CdS NBs make them interesting for various DMS applications. These emissions are attributed to the interaction between the free excitons and ferromagnetic cloud, as previously reported in other II–VI materials.⁹⁷ The yellow band emission at 572 nm is attributed to Mn²⁺, which has been reported to be located in the range of 576–598 nm and is assigned to the d–d transition between the ⁴T₁ excited state and the ⁶A₁ ground state of Mn²⁺. Besides the band edge, other emission peaks also appear independently under a 532 nm excitation source below the band



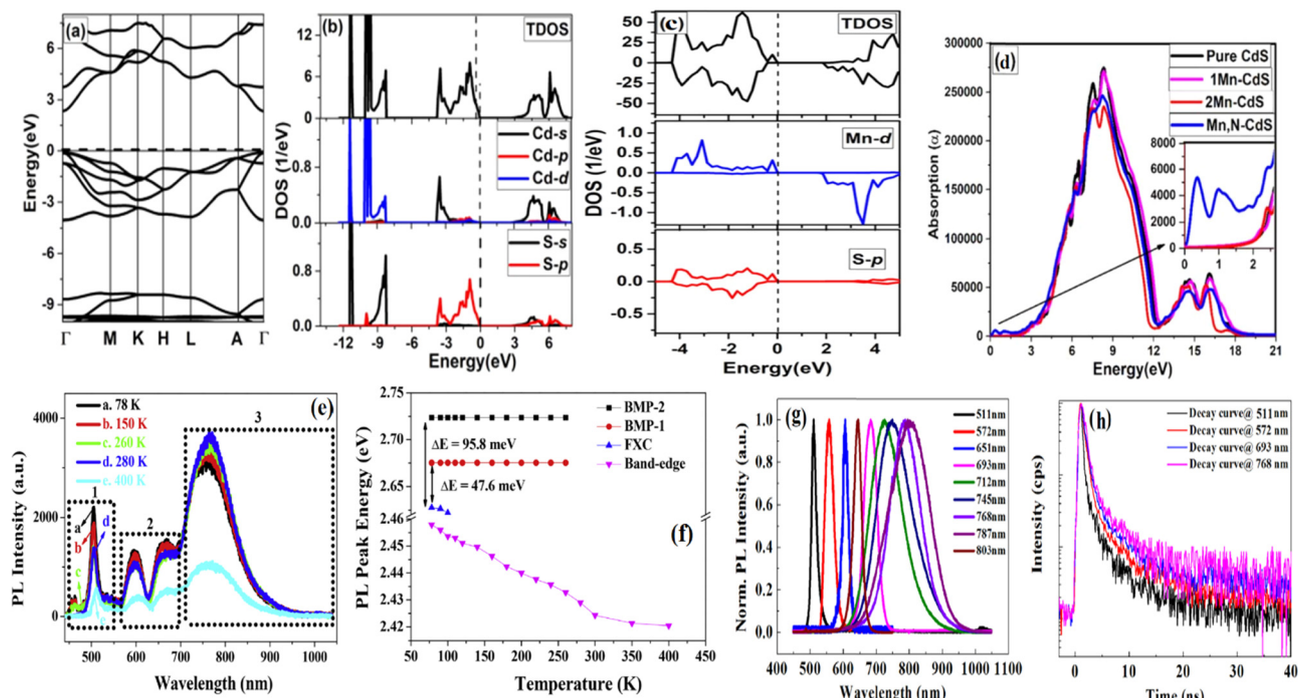


Fig. 2 (a) Calculated band structure and (b) TDOS and PDOS of pure CdS. (c) TDOS and PDOS of Mn-d, S-p and N-p for FM configurations in $\text{Cd}_{34}\text{Mn}_2\text{S}_{36}$. (d) Calculated optical absorption coefficient for pure CdS and Mn-doped CdS with and without N co-doping.⁹⁵ (e) Temperature-dependent micro-PL spectra in the bright-field optical image of an individual Mn-doped CdS nanosheet. (f) Temperature-dependent PL peak positions of the band-edge for CdS (purple), FXC (blue), BMP-1 (red), and BMP-2 (black) in the Mn(II)-doped CdS nanosheet. (f) AFM- and FM-coupled Mn ions in CdS.¹⁰⁷ (g) PL spectra of the obtained single undoped CdS and Mn^{2+} -doped CdS belt at 300 K. The black line is from the pure CdS belt and the other lines are from the emission of CdS doped with different Mn ions. (h) PL lifetime decay plots for the band-edge emission together with Mn^{2+} -related emissions at room temperature in an Mn-doped CdS single belt excited at 405 nm.¹⁰²

edge with the presence of a near band edge emission at 515 nm upon 405 nm excitation. This tunable emission owing to the Mn interaction with the host material has rarely been reported for DMS structures.^{8,103} Thus, the ferromagnetic exchange in this (MnS)_n cluster has strong coherent coupling between magnetic ions within the lattice, which may lead to the formation of localized exciton magneton polarons because of the p-d hybridization and FM coupling.^{108–110} Fig. 2h shows the PL decay curves for 1D CdS with TM ion doping excited at different wavelengths (405 nm and 532 nm) to understand the Mn^{2+} transition with the host CdS crystal symmetry.¹⁰² According to the PL decay curve, we can understand that the small change with a few nanoseconds (ns) smaller than the free exciton lifetime indicates that it is related to the combination of free excitons and EMP in Mn-doped CdS and the large change is due to the presence of inhomogeneous states (defects) in the structure. However, the steady changes in lifetime in the same direction indicate that the emission spectra in the red to near IR region are related to the Mn ion and no defect-related states are involved in the bands. A further increase in excitation power will reduce the lifetime due to the interaction of more spin states and large coherent coupling.^{111–113} Therefore, this lifetime in the range of 1 ns reveals the reflection of spin-spin interaction during lattice relaxation, as reported in other II-VI semiconductors. Fig. 2e shows the luminescence spectra of an Mn-doped CdS nanosheet, highlighting the three different emission spectra in the range of (1) 440–550 nm, (2) 560–680 nm

and (3) 700–1050 nm. The emission spectra in the range of 440 nm to 550 nm (green emission) at 78 K can be subdivided into triplet emissions owing to the splitting of the valence band into three sub-bands. According to the crystal field symmetry, the spin-orbital interaction of excited electrons can occupy three exciton energies [A-exciton (2.5535 eV), B-exciton (2.5675 eV) and C-exciton (2.629 eV)].^{107,114,115} Fig. 3f confirms the existence of a band edge emission, free exciton emission and spin-exciton interaction at low temperature. Moreover, the band-edge peak emission as a function of temperature is mainly observed in wide bandgap semiconductors.

Ge *et al.* reported the coherent bosonic lasing of EMP at 520 nm at $232.393 \mu\text{J cm}^{-2}$ for Ni-doped CdS synthesized *via* the VLS mechanism.⁹⁸ Fig. 3a and b show the room temperature PL spectra of doped and undoped CdS and a low temperature for an Ni concentration of 1.54% in CdS NB, respectively. In the VLS mechanism, the S vacancies are normally generated during the growth of a 1D system in the presence of hydrogen in the source gas. In the case of Ni^{2+} -doped CdS, the dopant atom substitutes the Cd ion at a higher concentration but this may not happen for a low doping concentration.^{114,115,117} Another important discussion is the crystal symmetry of the host material and dopant ion interaction with the crystal symmetry. Under luminescence, the transition of the carrier excitation and de-excitation/recombination will provide detailed information about the spin-carrier interactions in semiconductors. Therefore, the EMP-related peak in Ni-doped CdS



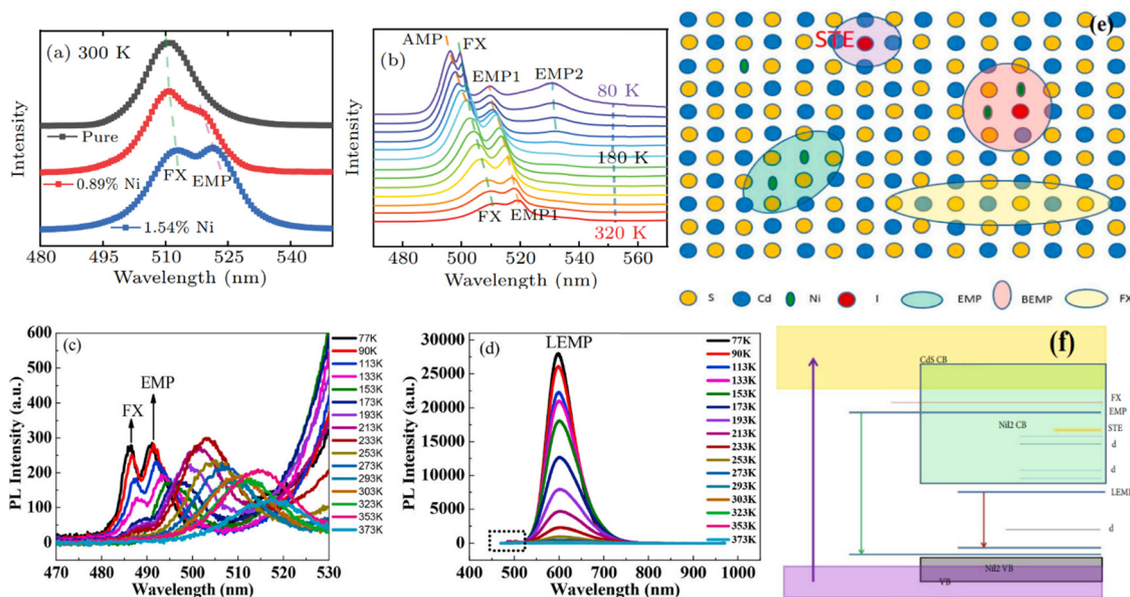


Fig. 3 (a) Micro-PL spectra of CdS NB at different Ni concentrations in the range of 480–550 nm at RT. (b) Temperature-dependent micro-PL spectra of CdS NB with Ni concentration of 1.54% in the range of 470–570 nm.⁹⁸ (c) Temperature dependence of PL of single Co-doped CdS NB excited with CW laser at 405 nm. (d) Magnified selected spectrum in the range of 470–530 nm, as marked by a rectangle.⁵ (e) Spatial distribution diagram of the different excited states in NiI-doped CdS lattice. (f) Electronic structure of CdS:NiI₂ NB, in which the nanocluster of NiI₂ is incorporated.¹¹⁶

will become dominant with an increase in the dopant concentration, which is directly correlated with the coupling of the dopant cation in the crystal symmetry. In the case of the pure CdS belt-like structure, the redshift in its PL spectra is owing to deficiency of S ion during its growth, which may lead to the formation of donor levels in the conduction band. Consequently, the dopant ion incorporated in the CdS lattice will induce more defects within the host lattice, which may produce strong exchange interaction with the spin magnetic ions with native defect states.^{115,117,118} This strong exchange interaction produces a broad EMP emission or redshift at room temperature. Ge and co-workers reported an increase in the intensity of the free exciton and EMP emission ratio. The lattice shrinkage may weaken the electron–phonon interaction at low temperature. At 80 K, different peaks appear in the emission spectrum, corresponding to free excitons, EMPs and antiferromagnetic magnetic polarons of Ni²⁺ pairs. Moreover, the bosonic lasing of EMPs condensates out of complicated states upon excitation by a femtosecond (fs) laser, as reported in other II–VI 1D nanostructures.^{3,117} The spectra ranging from the red to near-IR region originate from the FM coupling with the Mn clusters become prominent at a temperature lower than 260 K owing to the localized EMP, as observed in Mn-doped CdS. The same behavior of EMP formation has been observed in ZnSe Co-doped NWs. However, the lowest transition level for Co-doped CdS exhibits p–d hybridization through the high 3d state, which normally possesses a longer lifetime.¹¹⁸ The low-temperature luminescence spectra of Co-doped CdS reveal strong free exciton, EMP and LEMP emissions, as shown in Fig. 3c and d. The TM in the host material also results in a red shift in the band edge and LEMP emissions due to the high level of d–d transitions from ⁴T₁(P) to ⁴A₂(F) of the Co ion.^{5,119–121} When the temperature is lower than room temperature, the slight blue shift in the intra-shell d–d

transition emission in the emission spectra implies that the d–d transition in Co-doped CdS is less localized than that in the Mn-doped CdS lattice. Based on the VLS mechanism, various studies highlighted the successful incorporation of TM ions in 1D CdS NWs and NBs, which exhibit room temperature ferromagnetism and anti-ferromagnetism. Ni-doped CdS crystals also exhibited a broad EMP emission at 600 nm, which is attributed to the d–d transition of ³T₁(P) to ³T₁(F) of the intrinsic Ni(II) ions in sulfide. Gaussian fitting of the broad EMP emission based on experimental data for Ni(II) doped in CdS revealed that the cubic crystal field may split due to the reduced symmetry information; therefore, the broad emission in Ni-doped CdS is due to its two-level emission, which is difficult to distinguish in comparison with the Mn-doped CdS nanostructure synthesized *via* the VLS mechanism.^{107,109,113}

Fig. 3e shows single Ni and I and coupled Ni + I doping in the host lattice, resulting in an inhomogeneous DMS structure. Fig. 3f shows the ion doping-related electronic structure of CdS:NiI₂ NB. The EMP emission observed very far from the CdS:NiI₂ band edge emission owing to the high magnetic ion concentration or defect formation due to F or Cl during growth *via* the VLS mechanism in the presence of hydrogen in the source gas. Meanwhile, Br or I ions are heavy and may be co-doped with Ni in the host lattice, exhibiting strong lasing phenomena, which will be detailed discuss in the lasing section. The co-dopant ion in the CdS lattice may lead to the formation of exciton complexes or acceptor/bound excitons or self-trap excitons, as shown in Fig. 3e, which have different exciton spatial distributions within the lattice.^{114,120}

High-density EMPs can polarize and combine with the free excitons in 1D NB of CdS:Co, leading to coherent emission behavior at room temperature. EMP condensation in NB happens due to the following reasons: (1) spin coupled with excitons and phonons; (2) spin ion aggregation with FM coupling; (3) EMP is



more stable than free excitons in materials having a higher exciton binding energy than thermal energy, which are coherently coupled through FM exchange interaction; and (4) the 2LO phonon along the *c*-axis in a hexagonal structure has a large transition probability and strong polarizability to interact with excitons, favoring the EMP state in TM-ion doped semiconductors. Due to its weak interactive force, highly stable spin–exciton–phonon (LO phonon) polarization may exhibit a bipolaron emission in the PL spectrum at high temperatures. Moreover, the much lighter effective mass of excitons than electrons in the CdS:Co semiconductor results in their strong interaction with EMP condensation at a critical temperature. The steady-state spectrum exhibits a two-band emission near the bandgap, which is related to the free excitons and EMP at room temperature, as shown in Fig. 3c, for 1D NB of Co-doped CdS. The second peak (518 nm) is very close to the 2LO A1 phonon, which indicates that EMP formation close to the band edge requires phonon polarization in the same direction. Therefore, the orientation-dependent EMP collect free excitons to condense and produce a lasing emission even in the perpendicular direction. This coherent EMP condensation resulting in lasing in II–VI 1D structures needs to be studied under ns laser excitation for a better understanding of the exciton–exciton, exciton–electron, exciton–phonon and exciton–spin interactions on variable timescales.^{100,122} The spectral line width and intensity are strongly dependent on the excitation power of Co-doped CdS. In the case of the threshold power of 7.4 μJ, the linear increase in intensity is due to the inhomogeneous distribution of magnetic ions in the belt. Alternatively, for a threshold power of 16 μJ, the lasing line width and intensity are not very wide due to the exciton–carrier or exciton–phonon interaction, which may involve lasing in single structures. This random lasing in 1D structures depends on the exposed area, which is helpful to understand the collective EMP lasing phenomena in doped wurtzite microstructures.^{100,123} The collective EMP condensation favored by the 1D longitudinal polarization and FM coupling of magnetic ions can be detected at room temperature for a better understanding of the magnetic exciton BEC in DMS nanostructures. The exciton magnetic polaron states in 1D NB and NW appear due to the exchange interaction between the spin–carrier interactions, resulting in the formation of EMPs. The overlapping tails of the carrier wavefunctions and spins result in large cluster formations under a high excitation density and respond as collective magnetic polarons to pronounce the bosonic nature in NWs and NBs. Therefore, the collective EMP condensate due to coherent light in nanostructures shows photo-magnetism behavior, which increases the possibility of modulating single-mode lasing by FM coupled with exciton in DMS. The large EMP coherence space and optical or magnetic modulation in the 1D nano/microstructure of wide bandgap semiconductors need further study for their nanophotonic application in future devices.⁵

3.1.2 CdSe. Cadmium selenide (CdSe) possesses a direct bandgap (~1.74 eV), and thus exhibits n-type conductivity and astonishing electronic and optoelectronic properties similar to other II–VI materials.^{10,83} Its wurtzite structure is more stable than its zinc blende and rock salt (normally obtained at high

pressure).^{124–127} Deng *et al.* successfully prepared zinc blende NC using low cost and green reagents. The preparation of 1D tube CdSe nanostructures including NW, NB and NT has been reported using both physical and solution synthesis mechanisms.¹²⁷ Dai *et al.* successfully controlled the shape of CdSe NCs during their synthesis and demonstrated that elongated nanorods (NRs) possess many interesting optical properties.¹²⁸ Kuno *et al.* revealed that the nanocrystallite emission originating from the surface-related defects is due to its large surface-to-volume ratio.¹²⁹ However, its bandgap luminescence arises from its optically inactive fine structure state or dark excitons. Meanwhile, Pan *et al.* reported an enhancement in the luminescence spectra of CdSe NW and NBs synthesized *via* the VLS mechanism.¹³⁰ The growth of 1D nanostructures is directed by well-confined structures, resulting in high optical gain under high power, which are suitable for ultrafine red-color lasing. In the case of 1D CdSe, the excited electrons in its conduction band interact with phonons and rapidly decay, which is in contrast with QDs of CdSe due to the phonon bottleneck effect.^{131,132} Mirov and co-workers successfully doped TM ions in CdSe and other II–VI semiconductors to realize mid-IR lasers.¹³³ The related ternary alloy dopants occupied the tetrahedral sites in the cubic structure of the II–VI semiconductors instead of the octahedral sites in the crystal. The tetrahedral sites in the crystal provide a smaller crystal field splitting and shift the emission to the IR region due to the presence of the dopant ions.

3.1.2.1 Transition metal ion-related luminescence behavior. To understand the lasing phenomena in the CdSe matrix, it is important to determine its spatial recombination through spectroscopic measurements. CdSe NC has a poor band edge emission, which is attributed to two main reasons, as follows: (i) the optically inactive state at the bottom of its conduction band, which is known as dark exciton and (ii) transition from defects to the valence band. Interestingly, in the case of 1D nanostructure CdSe synthesized *via* the VLS mechanism, it exhibits strong luminescence at high excitation, which may increase the population rate at the lowest conduction band level. Alternatively, high crystallinity was achieved in the hexagonal CdSe nanostructure obtained *via* the VLS mechanism, which exhibited the red color lasing line of individual crystalline CdSe NB, as reported by Pan.^{130,134,135} Meanwhile, the size-dependent and lateral thickness of the CdSe nano-platelets tuned the LO–phonon corresponding to low exciton–phonon coupling, large oscillator strength and energy spacing of both. The optical transition energy related to the exciton–phonon interaction in CdSe was controlled by its lateral size and thickness by Ahtsheim.¹³¹ CdSe platelet-like structures are attractive for optoelectronics and enable control of the exciton energy states based on their thickness; however, their lateral dimension disparity in LO–phonon coupling is independent of their thickness and lateral size.

Another major key feature of II–VI materials is that heavy anions in the crystal field offer a low optical phonon energy cutoff value and reduce the efficiency of non-radiative decay. This decrease in efficiency provides the potential to realize a high fluorescence yield at room temperature, as shown in



Fig. 4a.¹³³ Yu *et al.* synthesized CdSe nanoplatelets and nano-ribbons doped with Mn^{2+} ions using the solution process mechanism, as shown in Fig. 4b.¹³⁶ TM ion doping in CdSe in low-dimensional systems (NWs, nanoribbon, *etc.*) results in unique magneto-optical behavior, as characterized by various spectroscopic analyses. Mn^{2+} doping in CdSe results in strong carrier-induced ferromagnetism due to the sp-d exchange interaction between the magnetic ion and electronic state of CdSe. Thus, quantum confinement at the nanoscale provides dominant carrier-spin exchange interactions and opens new horizons in the field of DMS.^{137–141} Mikulec and Erwin reported that the incorporation of Mn^{2+} in ZnSe and CdS is comparatively more convenient than in the CdSe matrix. The probability of Mn ion doping in the CdSe matrix is 50% for its bulk, whereas in its NC, it is still challenging.^{142,143} The surface adsorption of magnetic ion impurities during the nucleation process at the nanoscale results in the formation of clusters,

which known as magic-sized clusters suitable for uniform doping in the host crystal. In the case of nanoscale or 1D nanostructures, a reduction in their dimensions may increase the possibility of quantum confinement and provide a low probability of impurity ions adsorbed within the magic-sized cluster. Yu *et al.* successfully doped Mn^{2+} ions in CdSe NC, as shown in Fig. 4b.^{136,144} Magnetic ion doping in magic-sized cluster formations during the nucleation stage of growth provides a uniform doping concentration in the host semiconductor. Cu^{2+} , Ag^{2+} and other TM ions have been successfully doped in NC, exhibiting a red-shifted broadband emission at room temperature. Pan *et al.* prepared hexagonal CdSe NW- and NB-like structures *via* the CVD method, which exhibited red-color lasing within individual nanostructures.¹³⁰ Chen *et al.* also prepared a 1D CdSe nanostructure *via* the CVD method, which exhibited strong excitonic behavior at different temperatures, as shown in Fig. 4c–e.¹⁴⁵ Its temperature dependent absorption

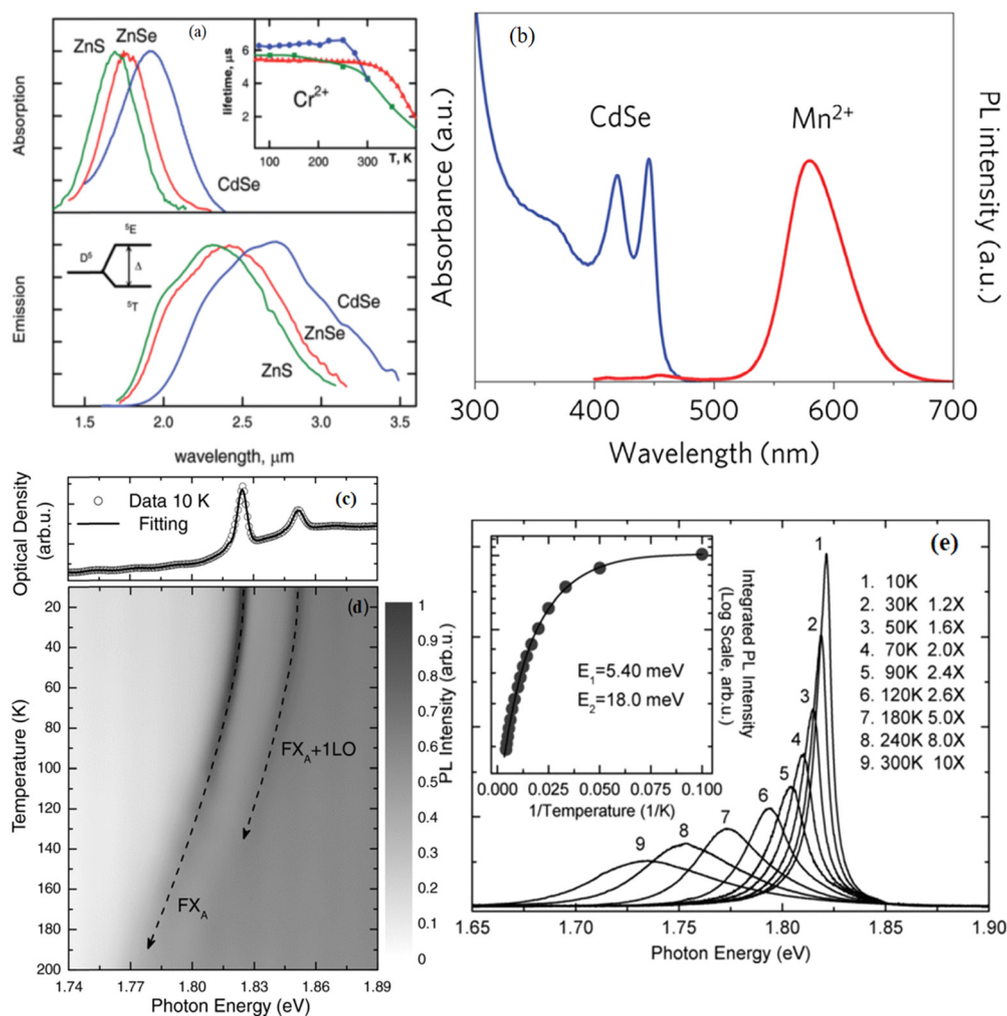


Fig. 4 (a) Normalized room temperature absorption and emission-gain spectra of Cr^{2+} ions in ZnS, ZnSe, and CdSe crystals. Inset shows the temperature dependence of Cr^{2+} ion lifetime in ZnS (circle), ZnSe (triangle), and CdSe (square) crystals; Copyright © 2010, Wiley InterScience.¹³³ (b) Absorption (blue line) and PL (red line) spectra of Mn-doped CdSe NPLs; Copyright © 2009, Springer Nature.¹³⁶ (c) Low temperature (10 K) free-exciton absorption of CdSe NWs. The solid curve is the multi-Gaussian fitting of the absorption spectrum. (d) Temperature-dependent mapping of the absorption spectrum. (e) Free-exciton emission of CdSe NWs at various temperatures. The inset plots the dependence of the integrated PL intensity on the reciprocal of temperature, and the solid curve is the Arrhenius fit with two activation energies, Copyright © 2011, Wiley.¹⁴⁵



spectra at 10 K show that the strong emission spectra of the exciton peak (1.8248 eV) and free exciton peak coincide with the LO phonon (1.8515 eV) of CdSe NWs. Due to lattice expansion, the low temperature mapping of CdSe NWs strongly exhibits the exciton–phonon interaction. According to the Bose–Einstein approximation, the exciton–phonon interaction function of temperature and low temperature variation is very helpful in analyzing the thermal expansion and exciton–phonon interaction due to bandgap shrinkage. Similarly, the temperature-dependent luminescence spectra show an increase in intensity at low temperature and the inset image in Fig. 4e shows the difference in activation energy (5.40 eV and 18 meV), as calculated using the dual activation energy model, where the energy of 18 meV is comparable with the bulk exciton energy.^{145,146} According to the crystal field theory, the ground state of the Cr²⁺ ion is 5d to the upper levels in the visible spectrum. The tetrahedral site of the crystal field (T_d) splits 5d into triplet (⁵T₂) and doublet (⁵E) states. In the case of Cr²⁺, the triplet state allows the transition between levels and the allowed spin state, whereas in the case of Fe²⁺, the doublet state allows the transition in the host matrix. Therefore, the Cr²⁺ ions can undergo a transition from ⁵E (5d) to ⁵T₂ (5d), which is promising for laser application. The Jahn–Teller mechanism and spin–orbital coupling are also responsible for the energy splitting. Numerous reports have shown that Cr²⁺-doped CdSe exhibits direct lasing at room temperature. Similarly, Chen *et al.* showed that a change in the excitation density will enhance the spontaneous emission; therefore, multiple peaks appear at 715 nm and 725 nm (above the threshold value of 0.296 MW cm⁻²) at room temperature. These multimode random lasing in a single structure of CdSe NW in the NIR region make it suitable for NIR optoelectronic applications such as sensors, lasers and energy conversion devices.^{133,147,148}

Solution-processed colloidal QDs exhibit exciton and bi-exciton lasing at room temperature. Their spectral symmetric PL is temporally integrating the streak image over the nanosecond fitted by the pseudo-Voigt function for low and high fluence provide information related to their exciton and bi-exciton transition. The giant oscillator strength near the band edge emission has strong coherent phasing strength, as correlated with a single atom in the host matrix. Therefore, ultrafast dynamics spectroscopy provides detailed information of the carrier and decay kinetics in the picosecond (ps) domain. The bi-exciton lifetime of the CdSe matrix is dominated by non-radiative Auger recombination up to 10 ns, which is almost the two order of magnitude larger than the measured radiative lifetime. The streak camera images of a CdSe CQ well excited with 1 kHz, 70 fs laser pulses and fluence of 20 μJ cm⁻² with the ultrafast time decay measurement value about ~10 ps unambiguously assigned the peak to the stimulated emission, providing evidence that the intensity-dependent spectra originated from the bi-exciton transition within the respective energy position at an ultralow threshold fluence.¹⁴⁹

3.1.3 CdTe. Similar to other II–VI semiconductors, CdTe exhibits strong luminescence behavior and is an attractive

material for various optoelectronic applications such as solar cells, photodetectors, and lasing. CdTe is a direct bandgap (1.45 eV) semiconductor with inherent p-type conductivity due to the presence of Cd vacancies, whereas CdS and CdSe show n-type conductivity.^{150–152} Tang *et al.* reported the spontaneous organization of CdTe NWs from NPs due to strong dipole–dipole interactions.¹⁵² Neretina *et al.* synthesized vertically aligned CdTe NWs through a catalytically driven growth method.¹⁵³ Hiesinger *et al.* studied both n-type and p-type CdTe excitation spectra with exciton luminescence at 2 K.¹⁵⁴ In the case of n-CdTe, two peak lines appeared at 1.59375 eV and 1.59320 eV, corresponding to an exciton bound to a neutral donor (D⁰, X). The emission peaks at 1.5919 eV and 1.5893 eV originated from an exciton bound to an ionized donor (D⁺, X) and exciton bound to a neutral acceptor (A⁰, X), respectively. Alternatively, in p-type CdTe, all these emission lines appeared stronger with two other emission lines at 1.5896 eV and 1.5898 eV corresponding to an exciton bound to a neutral acceptor (A⁰, X). In the case of n-CdTe, the weak bands at 1.596 to 1.598 eV are assigned to the spin-triplet exciton, which is also called the lower branch polariton (LBP) and upper polariton branch (UPB) of free excitons, respectively. The higher energy emission at 1.6031 eV is related to the *n* = 2 excited state of free excitons, whereas the low energy onset showed the 1LO phonon sideband at 1.5747 eV. At 20.6 K, the 8LO phonon peaks appeared with a background, whereas at 1.8 K, the exciton coincides with the low energy threshold field to increase the value at a low temperature. In the case of p-type CdTe, the LO phonon peaks appear with lower intensity than n-CdTe with two additional peaks. As the temperature increased below 20 K, these two additional peaks due to neutral acceptor (A⁰, X) disappeared between the oscillatory structures.

3.1.3.1 Transition metal ion-related luminescence behavior.

Various researchers have extensively reported TM ion doping in CdTe for DMS and optoelectronics applications such as solar cells. TM ion doping in II–VI materials is related to the 4d element effect on both optical and electrical properties. Kurchatov and others research reported the successful incorporation of TM ions within the II–VI host matrix, which exhibited IR lasing emission at room temperature.^{155–157} Lafuente-Sampietro *et al.* reported the resonant optical control of the spin (single Cr atom) in CdTe/ZnTe QDs.^{158,159} At 5 K, the Cr spin relaxation was observed by a strong resonant laser field, which is responsible for the Cr atom tuning optical Stark effect. The circular polarized PL spectra of the magnetic anisotropy of the Cr spin were induced by biaxial strain. The exchange interaction between the spin atom and exciton further splits the Cr energy level due to the high polarizability of the crystal field. To understand the carrier dynamics in CdTe/ZnTe QDs doped with Cr atoms, a pump probe experiment was performed to observe the relaxation time and non-equilibrium distribution of the Cr spin population with a circularly polarized resonant pump pulse. The peak intensity map of Cr doped in CdTe/ZnTe QDs showed the low energy bright exciton state and dark exciton. The Cr ion interaction with phonons in the



system shows strong spin–phonon coupling at low temperature, which is an important step forward to control the TM ion coherent spin for future spintronic devices.^{158,160} Similarly, the Cr spin is coupled with excitons near the valence band and shows nano-magnetic behavior due to the optical pumping of electron–Cr and hole–Cr. Resonance and non-resonance optical pumping can be directly used as spin memory (write and erase, respectively). Therefore, the spin channel can be studied with polarized emission, which responds to the relaxation channel of the Cr ion in II–VI semiconductors. Therefore, the optical pumping- and power-dependent PL intensity-related temperature effect is directly affected by the hole–Cr spin near the valence band and electron–Cr local generation of phonons.

Optical pumping shows the positively charged exciton and relaxation of Mn-spin exchange coupled with confined hole spin in CdTe. This spin can be observed under the optical injection of spin-polarized carriers in the time domain of a few tens of ns. Interestingly, the localized spin state can be controlled by the charged state of strain-free QDs. Thus, the magnetic anisotropy of an atom can be controlled by an electrical field, making it suitable for miniaturized data storage application for fine structure and strain-free QDs. This fine structure shows the spin splitting of the Mn ion in the energy range of a few meV coupled with heavy-hole spin near the valence band of CdTe. Pumping and a magnetic field influence the Mn-electron dynamics in the excited state and Mn-hole in the ground state. In the case of zero magnetic field, the spin-relaxation time of the Mn-hole is shorter due to valence band mixing.^{157,159–161} PL time delay measurements can further explain the dynamics of charge carrier interaction with the optically active injection of spin polarized photo-carriers. Similarly, Mn doped in CdTe QD shows a spin flip under resonant excitation, which is mainly controlled by the carrier–spin interaction. This coherent dynamics is directly observed in the ultrafast time domain and localizes the spin ion in the CdTe matrix.^{157,159–161} The Bose–Einstein condensation (BEC) of the exciton–polariton-related phase transition is related to the quantum effect in the CdTe/CdMgTe microcavity at 5 K.¹⁶² The spontaneous quantum degeneracy of polaritons in the CdTe microcavity excited by non-resonant pumping significantly narrows the polariton emission line width. The condensation of exciton polaritons in CdTe-based microcavity condensation is not standard BEC due to leakage in the cavity. The polaritons are non-interacting as bosons but do not follow the conservation law above some critical density at a specific temperature. Accordingly, the spatial coherence and macroscopic polarization through condensation can be used as a polariton-laser in CdTe. Thus, for wide bandgap semiconductors such as ZnO and GaN, the polariton condensation builds macroscopic polarization and spatial coherence for polariton lasers with an increase in temperature.^{162–165}

3.1.4 Ternary alloy of cadmium chalcogenides (CdX; X=S, Se, Te). The excitonic emissions of II–VI semiconductors can be easily observed at room temperature due to the fact that their exciton binding energy is higher than thermal energy. Therefore, many researchers have established optically pumped

stimulated emissions of II–VI semiconductors and their alloys 1D nanostructures. Secondly, the low threshold within low dimension II–VI semiconductors stimulates the excitonic emission, which is a prerequisite for lasing at room temperature. Thirdly, the formation of an optical cavity in 1D nanostructures (whispering gallery or Fabry–Perot mode lasing) enables lasing action to be achieved.^{166–169} In the case of ternary alloys, they have advantages over binary semiconductors including wavelength/bandgap tunability of multi-color lasing, as reported for both Zn and Cd chalcogenides. Tunable emissions from the UV to visible to NIR regions from Zn and Cd chalcogenide 1D structures have been demonstrated *via* different synthetic mechanisms such as VLS, VS, and solution process.^{77,170–172} Moreover, TM ion doping in the II–VI semiconductor matrix also results in lasing phenomena; therefore, lasing phenomena are often associated with exciton–exciton interaction, electron–hole plasma and exciton–polariton near the bandgap. CdS doped with NiI resulted in the formation of (NiS)_x and NiI₂ clusters in 1D NB, exhibiting a dual lasing line in a single structure at room temperature. Another important factor is the exposed excitation area, which shows the lasing threshold in a single structure in all directions, as observed in CdSe nanostructures.^{116,122,173–175}

A Cd-based ternary alloy CdSe_xS_{1–x} NW with a homogenous composition was synthesized through a solution–liquid–solid process by Yang and co-workers.^{58,176} Ternary alloy nanostructures have attracted significant attention owing to their continuous tunable bandgap, making them suitable for various electrical and optical applications such as photodetectors, solar cell, and FETs. Rosenthal *et al.* reported the ternary NC of CdSe_xS_{1–x} having different sizes and compositions; their results indicated that the surface states are strongly correlated with the relaxation process of photo-excited carriers, as studied by ultrafast fluorescence upconversion spectroscopy.^{176–178} Colloidal CdSe_xS_{1–x} NWs exhibit strong emission, which can be tuned in the range of 508 nm to 628 nm with a narrow spectral width together with a high quantum yield. Therefore, the tunable and polarized emission of CdSe_xS_{1–x} NWs provides greater insight to understand the exciton dynamics for various optical applications, as shown in Fig. 5a–d. Fig. 5a and c show the absorption and PL spectra of CdSe_xS_{1–x} alloy NWs, respectively, where the variation in the composition gradient shows a redshift in the bandgap tuning with an increase in the Se ratio. A similar study was conducted for CdSe_xS_{1–x} alloy NWs synthesized *via* CVD, NCs and quantum dots (QDs).^{170,179–182} At low temperature PL, the defect-related emission of different compositions was suppressed except for CdSe and CdSe_{0.25}S_{0.75}, indicating that the formation of alloy CdSe_xS_{1–x} NWs can be feasible for high-quality polarized LEDs, as shown in Fig. 5f. The spectral broadening of the emission can be related with (i) inhomogeneous broadening, (ii) optical phonon interaction and (iii) acoustic phonon interaction at low temperature. Alternatively, the bandwidth broadening can be related with (i) acoustic phonon scattering, (ii) longitudinal optical (LO) phonon scattering and (iii) exciton surface scattering. Below 130 K, for CdSe and CdSe_{0.25}S_{0.75}, the acoustic phonons contribute significantly (dotted line comes from inhomogeneous broadening and acoustic phonon scattering)



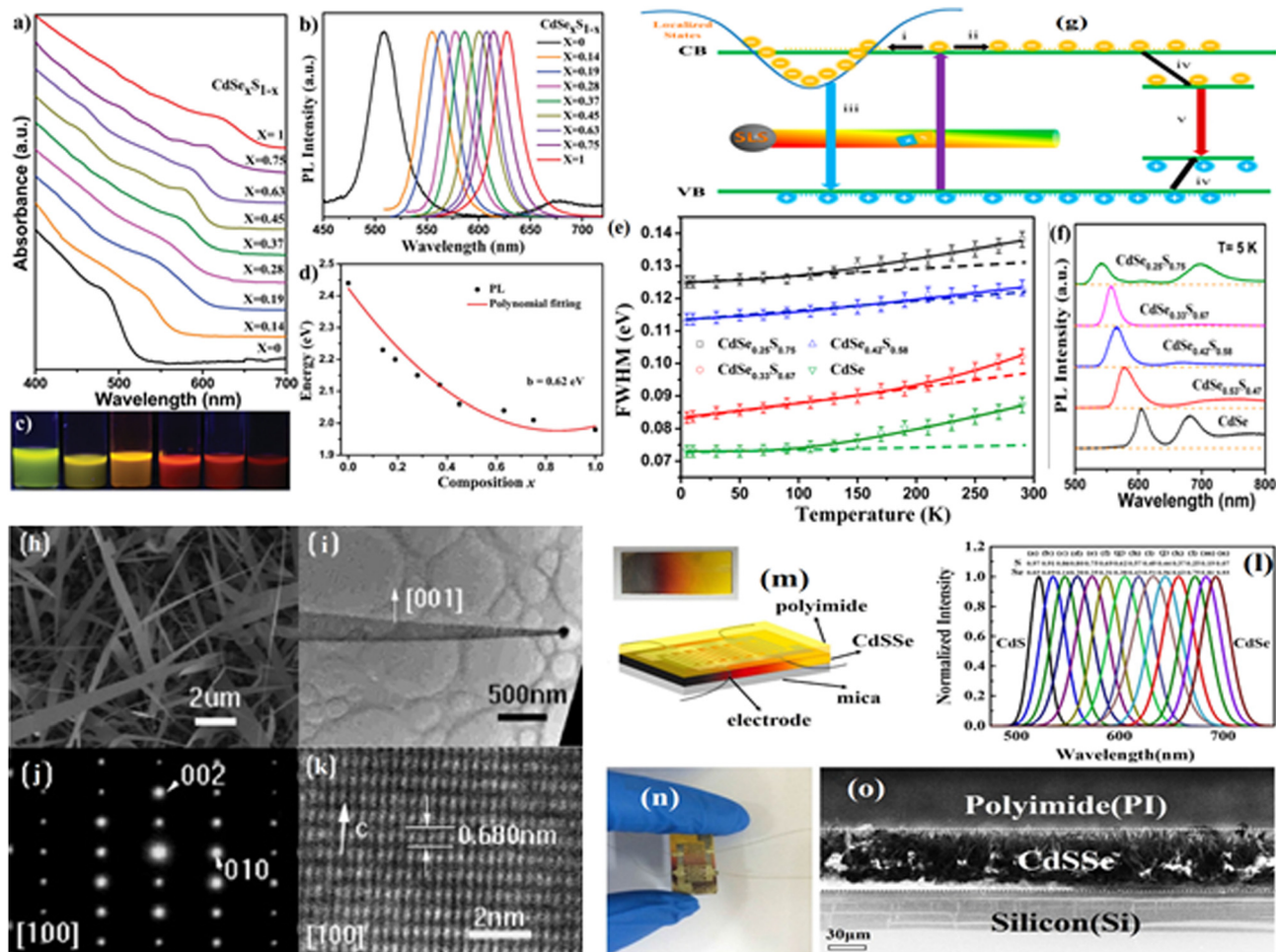


Fig. 5 (a) UV-vis absorption spectra of $\text{CdSe}_x\text{Se}_{1-x}$ prepared via a solution process and (b) PL spectra of $\text{CdSe}_x\text{Se}_{1-x}$ NWs with different Se contents. (c) Photographs of typical emission colors from the obtained alloyed NWs under UV 365 nm irradiation. (d) Energy gap–composition correlation of $\text{CdSe}_x\text{Se}_{1-x}$ NWs, where the data points represent the value extracted from their respective PL.¹⁷⁶ (e) Solid lines were fit using Varshni's law and the temperature-dependent exciton–phonon model, while the dotted line represents the inhomogeneous broadening and the acoustic phonon scattering. (f) PL spectra of varying compositions of alloyed NWs at 5 K. (g) Schematic of the kinetic model of exciton recombination in NWs.⁵⁸ (h) SEM morphology of ternary $\text{CdSe}_x\text{Se}_{1-x}$ NBs synthesized by CVD; (i) TEM image; and (j) and (k) selected area electron diffraction (SAED) pattern and HRTEM image, respectively.⁷⁰ (l) PL spectra measured at different composition along the length direction of a $\text{CdSe}_x\text{Se}_{1-x}$ NW. (m) Real-color photograph of $\text{CdSe}_{1-x}\text{Se}_x$ chip and diagram of the sweat sensor and (n) real image of the as-prepared sweat sensor. (o) Cross-sectional SEM image of PI-coated $\text{CdSe}_x\text{Se}_{1-x}$ NW chip.¹⁸⁸

but above 130 K, the LO phonon and surface phonon contribute more effectively and are responsible for the bandwidth broadening, as shown in Fig. 5e. According to their luminescence spectra, we can understand the recombination dynamics of photo-generated carriers with the help of a schematic diagram (Fig. 5g). This process can be described as follows: (i) photo-induced excitons can localize in the valleys, (ii) radiative recombination, (iii) generation of photoinduced excitons and (iv) surface defects and (v) trapped electrons recombine with trapped holes to produce defect-related emissions. However, both the compositional gradient of alloys and nanoscale dimensions strongly influence the exciton recombination due to the surface-induced quantum confinement, as reported in different studies.^{183–186} Therefore, PL decay measurement becomes an effective tool to analyze the recombination process at a very low excitation power. The preparation of $\text{CdSe}_x\text{Se}_{1-x}$ ternary alloys via the CVD route was reported by Pan and other research groups,

as shown in Fig. 5h–k.^{75,77,179} Liu's and co-workers also reported the preparation of $\text{CdSe}_x\text{S}_{1-x}$ alloy NWs for various applications such as sensors, thermoelectrics (Fig. 5l–o), wave-guides and photodetectors in recent years.^{187–189} Besides ternary alloys of II–VI semiconductors or heterostructures, radial core/shell nanostructures are also important for bandgap engineering, which have attracted attention due to their special structural characteristics and miscellaneous applications such as field-effect transistors, solar cells, biomedical applications, and memory devices.^{25,69–71,190} In the case of type-II heterostructures of CdSeTe, wurtzite (CdSe) and zinc blende cubic (CdSe) structures can be achieved by controlling the temperature and ligand. Moreover, the Se and Te contents in heterostructures tune the bandgap-related emission, while the diameter and shape of tetrapods affect the emission linewidth.¹⁹¹ Low-threshold lasing for CdTe at the surface of core of NW or NC exhibited a strong lasing emission line. Colloidal type-II (CdSe/CdTe) heterostructures have minimal



overlap of e-h wave function at the interface, which intensifies the amplified spontaneous emission with a high density of charge transfer and exhibits strong lasing at a lower threshold value.¹⁹² Interfacial type-II heterostructures lengthen the Auger recombination lifetime, which supports continued lasing, and the decrease in oscillator strength between electrons and holes will intensify the optical gain for LED and laser applications. Transient absorptions show the multiple oscillation at a high pump power assigned to acoustic phonon normally observed in single metal nanostructures with a quality factor of 9–10. The inhomogeneity of CdTe NWs lead to different charge carriers and charge trapping, which is related to the surface and pump power.¹⁹³

Pan *et al.* synthesized an axial NW-like structure of CdS/CdSSe *via* the VLS mechanism. The pumping fluence exhibited two broad emissions at 520 nm (green emission) and 600 nm (red emission) at a low pump power of 29 $\mu\text{J cm}^{-2}$, corresponding to spontaneous emissions. Both emissions showed a sharp bandwidth of 0.4 nm at a high power, demonstrating the stimulated emission in the heterostructure. Moreover, the slight blue shift at a power corresponding to electron-hole plasma formation at the surface contributed to exciton-phonon scattering and red shift due to the bandgap renormalization effect.^{75,77} Guo *et al.* synthesized a high-quality ternary alloy of CdS_{1-x}Se_x nanotriplets for broadband tunable single-mode lasing and high integration optical circuits and photonics communication.¹⁷³ Various factors such as the compositional variation and diameter of the tripod-related cavity had a prominent effect on tuning the band gap of CdS_{1-x}Se_x and single-mode lasing. Therefore, their tunable wavelength has numerous applications such as wavelength-converted devices, nanophotonic lasers and optical communication on signal chips.^{123,194,195} Yang *et al.* also reported the bandgap tuning of colloidal NWs of CdSe_xS_{1-x} synthesized *via* a solution process. At low temperature, their PL spectrum showed a surface defect-related emission, corresponding to the excitons and localization states based on the compositional ratio of the NWs. Therefore, the greater number of carriers localized in these states, which do not move to the surface, possible enhanced the radiative recombination in the ternary alloy.

The CdS/SnS₂ superlattice microwire synthesized *via* CVD showed multiple emissions, which were modulated by the excitons and photons in the 1D system. The physical deposition method successfully achieves strong lasing for multicolor emission at a low threshold value. Novel multi-peak emissions with controllable periods were observed for the first time due to the contribution from the 1D photonic crystal and periodical exciton confinement.^{58,176} Sn doping in CdS not only tuned the bandgap emission but also changed the defect-related emission line. Peng *et al.* also observed the strong electron-phonon coupling interaction in the optical micro-cavity of CdS-Sn, which is associated with the interaction, trapping and recombination of carriers in 1D nanostructures.¹⁹⁶ Tian *et al.* showed the effect of quantum confinement on the polarization anisotropy emission in CdS-Sn micro-cone-like structures. The polarization ratio achieved reached 60%, providing suitable information about the quantum confinement and optical confinement in nanostructures related to the dielectric contrast

upon the incorporation of Sn in the CdS matrix. Also, the decrease in the cross-section radius of the nanostructures was attributed to Sn doping and the reduction in the carrier lifetime. Moreover, the polarization ratio of the magnetic ion or spin ion corresponding to dielectrics is beneficial to study spintronic devices at room temperature.^{123,197,198} Bao *et al.* reported the single-mode laser of a CdS NW channel on an SiN waveguide chip with high efficiency of up to 58% by evanescent coupling. The composition gradient demonstrated various emission colors ranging from green to red, while the lasing emission wavelength ranged from 520 nm to 738 nm, providing broad wavelength tripod lasers.¹⁹⁹ Li *et al.* reported the multiphoton pump lasing of bi-excitons from a solution of colloidal CdSe/CdS nanoplatelets.²⁰⁰ The colloidal solution showed the optimal lateral size, which minimized the ultralow threshold for lasing at room temperature. CdTe alloys with other optical active materials such as CdSe as a type-II semiconductor tetrapod structure or ZnTe exhibit a strong optical response and are useful in various optoelectronic applications. 1D NW of CdS/CdSTe/CdTe showed high optical responsivity due to the n-type and p-type characteristics of CdS and CdTe, respectively. CdSTe acts as the depletion layer, which is similar to intrinsic silicon acting as sandwich in a PIN photodetector. Light illumination will transfer the photoelectrons from CdTe to CdS and transfer holes from CdS to CdSTe simultaneously, and then to the CdTe shell layer. The CdSTe layer reduces the carrier recombination probability and increases the photocurrent, making it suitable for photodetection in a wide spectral range.^{201,202}

Fig. 6a and b show the lasing in CdS NBs co-doped with 0.1–0.68% Co(II) and prepared *via* CVD. The two-band emission near the band-edge is located at 508–510 nm (FX) and at 518–525 nm for the EMP at room temperature. YAG:Nd (355 nm using a 2 ns laser) showed a lasing line at 518 nm (Fig. 6a). Fig. 6b shows the line width and intensity as a function of excitation power. Beyond the threshold power of 7.4 μJ , a super-linear increase in the emission intensity can be seen. Regardless of the inhomogeneous distribution of the dopant ions in NBs, a single lasing mode always occurs at the EMP location. Over 6 μJ , the exciton-carrier or phonon interactions start and become involved in the lasing process. The Co(II) ions with ferromagnetic coupling provide significant carriers and driving force to complete this collective emission process, as observed in other II-VI semiconductors. Other II-VI semiconductors doped with TM ions and excited by a fs laser exhibit an EMP lasing line, whereas CdS:Co NBs show lasing with both ns and fs lasers. This difference indicates that the EMP in the latter should have a larger coherence length and coherence time because ns laser excitation can cover more microscopic interactions, such as exciton-exciton, exciton-phonon, exciton-carrier, and exciton-spin, inside the excitation zone, while these interactions usually exist in variable time scales.³

The type-II interface of ZnTe/CdSe reveals a significant blue shift toward a lower energy band, whereas the high-energy band is related with CdSe. The relatively small lattice difference between two semiconductor core-shell nanostructures with a high surface-to-volume ratio is suitable for photovoltaic applications.^{171,190,203}



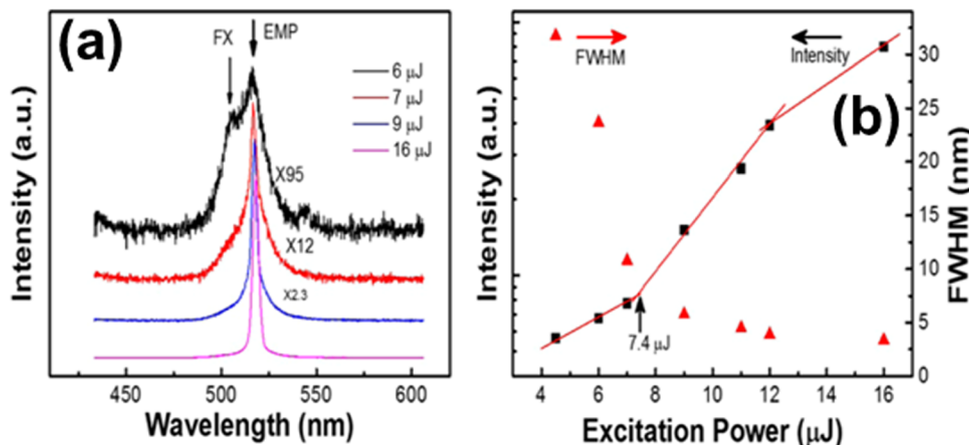


Fig. 6 (a) PL spectra from spontaneous emission to stimulated emission when excited by a nanosecond pulse laser (355 nm). The PL emission intensity and FWHM were extracted and plotted as a function of pumping power. (b) When the excitation intensity was below the threshold, a broad spontaneous emission band was observed, which showed a linear increase with excitation power, while when the pumping power was larger than the threshold, the emission intensity showed a super-linear increase with the excitation power. Moreover, the width at half-maximum (FWHM) became narrower an increase in the pumping power. Copyright © 2016, ACS Publications.³

Wang *et al.* demonstrated that CdSe/Cds/ZnS QDs with a spherical optical cavity propagated the light around their circumference due to the ZnS shell at room temperature.²⁰⁴ The ratio of S:Se in CdSeS NW could be controlled by the exciton dynamics together with bandgap tuning, as measured by fs transient absorption spectroscopy. The lifetime measurement showed a shorter wavelength with an increase in the Se ratio, while an increase in the S ratio revealed a large redshift in the steady-state absorption. The short time of about 10 ps with an increase in the Se ratio also revealed the photo-bleaching recovery from 8.4% to 57.7%, making CdSSe 1D NWs suitable for photovoltaic and optoelectronic applications with good control of both their optical and excited state properties. The homogeneity of an alloy depends on the phonon frequencies of the atomic mass in the unit cell and lattice configuration. A change in the S:Se ratio resulted in a linear shift in the Raman bands of the LO phonon frequency to a higher or lower wavenumber. The Raman bands of CdS and CdSe show the homogeneous distribution of Se and S atoms in QDs together with new Raman bands between the LO phonon frequency, showing the breaking of the translation invariance in the lattice. The additional vibrational peaks show the presence of an isotope mixture near the center of the optical mode, which is correlated with the linear frequency-mass dependency in the crystal system.^{170,182,183,205–207} In the case of QDs, the size-dependence broadening of their Raman lines due to phonon confinement, while the weak confinement and narrow bandwidth distribution result in a sharp Raman line due to the uniform or homogeneous distribution of the precursors.

3.2 TM ion doping in zinc chalcogenides (ZnX; X = O, S, Se, or Te)

3.2.1 ZnSe. ZnSe is an important II–VI semiconductor used in various optoelectronic applications such as nonlinear optical devices, LEDs, FETs, and panel displays. The 1D nanostructure of ZnSe has become very attractive due to its unique optical, electrical and thermal properties due to the reduction in its

dimensions. 1D nanostructures of ZnSe have been grown *via* the vapor–liquid–solid mechanism and solution process method to attain different types of 1D structures such as NWs, NRs, NBs, NTs, core/shell structures, bi-axial structures, tetrapods and alloys. Various researchers reported the 1D growth of ZnSe using various physical and chemical mechanisms.^{21,208–210} Li *et al.* reported the size-dependent catalytic growth of periodically twinned ZnSe NWs *via* CVD, whereas oriented arrays of ZnSe NRs were obtained by Zhang *et al.* using MOCVD.^{210,211} Wang *et al.* reported the epitaxial growth of 1D ZnSe NWs *via* MBE on a GaAs substrate.²¹² The ZnSe zinc blende QDs were converted into hexagonal wurtzite 1D NWs through the oriented attachment of NC. TM ion doping in 1D ZnSe also resulted in an excellent optical response, which has been extensively studied by various research groups.^{213–216}

3.2.1.1 Transition metal ion-related luminescence behavior.

Fig. 7 shows the PL spectra of different TM ion-doped (Mn^{2+} , Ni^{2+} , Co^{2+} , and Fe^{3+}) ZnSe nanostructures.^{20,117,213,216} Mn-doped ZnSe nanostructures have been synthesized *via* the VLS mechanism and solution process, as reported by many researchers.^{20,217–219} Fig. 7a shows the PL spectrum of Mn-doped ZnSe nanoribbons, where the peak at 461 nm is assigned to the near band edge emission at room temperature. At 77 K, the peaks split into two bands, originating from the free excitons and EMP. The emission peaks at 534 nm and 646 nm correspond to the low energy side and high energy side of the d–d transition (D1) originating from ${}^4\text{T}_1({}^4\text{G}) \rightarrow {}^6\text{A}_1({}^6\text{s})$ in Mn-doped ZnSe. The emission peak at 585 nm is related to the d–d transition. The d–d transition of Mn in ZnSe has been extensively reported in other II–VI semiconductors. In the case of ZnSe as the host matrix, the peaks at 534 nm and 650 nm corresponding to Mn can be attributed to the following: (i) Mn–Mn ions aggregate to form AFM coupling, (ii) self-activation of $\text{V}_{\text{Zn}}\text{–Cl}_{\text{Se}}$ complexes due to Mn doping, (iii) direct coupling of FM Mn–Mn pairs and (iv) generation of surface trap states. Therefore, structural relaxation is involved in the optical emission of the



Mn–Mn interaction pair in the ZnSe host matrix, crystal field and crystal lattice distortion to show the both FM and AFM pairs in the same zone. Therefore, low temperature PL provides sufficient information about the origin of the emission spectrum related to the Mn–Mn interactions in the host material. The emission peaks at 533 nm, 580 nm and 664 nm at 77 K are ascribed to the ZnSe emission related to the d–d transition and their small redshift is due to the sp–d hybridization effect. Fig. 7b shows the PL spectrum of Fe³⁺-doped ZnSe nanoribbons synthesized *via* the VLS mechanism, exhibiting three emission peaks at 467 nm, 553 nm and 630 nm.²¹³ The peak at 467 nm is assigned as the band edge peak, as observed in Mn-doped ZnSe (461 nm; Fig. 7a). The emission peaks at 553 nm and 630 nm correspond to the second (D1) and third (D2) emission peaks related to the Fe³⁺ d–d transition, respectively. Moreover, the emission peak at around 467 nm under different excitations shows a redshift with an increase in intensity due to the following reasons: (i) bound excitons, (ii) EMP interactions and (iii) e–h plasma formation at the surface. In the case of low power, the bound exciton peak is dominant, while a further increase in power resulted in a red shift in the peak and increase in its intensity due to EMP. With a further increase in power, the emission peak reached 493 nm due to e–h plasma formation at the surface of the nanoribbon, which is related with excess charge of Fe³⁺. Therefore, low temperature luminescence and lifetime measurements provide suitable information about the emission spectra in the host matrix, as also studied by different groups.^{220–224}

Fig. 7c shows the emission spectra of Ni-doped ZnSe at different temperatures.¹¹⁷ At 300 K, the two emission peaks appear at 463 nm and 471 nm, which are assigned to the bound excitons and EMP, as discussed for Mn²⁺- and Fe³⁺-doped ZnSe.^{225–228} At low temperature, the shoulder peak at 453 nm appears due to the incorporation of Ni ions; however, the combination of free excitons with AFM coupling with Ni ions or clusters resulting in the formation of this peak has rarely been reported in magnetically doped semiconductors.^{214,229–231} Hou reported that the incorporation of TM in ZnSe is the origin of its ferromagnetism, which may lead to a red shift in its emission spectrum. However, AFM coupling may lead to a blue

shift in its emission spectrum when coupled with a d–d transition. Normally AFM coupling appears at a high doping ratio with bound excitons but is rarely reported with a low doping. Other studies on Ni-doped ZnSe also revealed the importance of DMS for future spintronics.^{232–234} Fig. 7d shows the emission spectra of Co-doped ZnSe, which was prepared by Zou *et al.* *via* the VLS mechanism, exhibiting anomalous non-linear optical behavior due to the incorporation of magnetic ions in the ZnSe nanoribbon-like structure. The PL emission spectra were recorded with a change in the Co doping concentration from 0.14% to 1.07%. It was observed that as the doping concentration increased, the intensity of the peak at 555 nm was enhanced due to Co–Co coupling. Thereby, the intra-shell with high d–d transition was responsible for the FM coupling of the TM ion pair, which may produce a nonlinear optical absorption. Moreover, the intensity ratio of the band edge emission and d–d transition can tune the visual emission under different doping ratios, as normally reported in NC.^{235–238}

Lasing in pure ZnSe is attributed to its near-band edge emission, which is sharp due to the coherent transition without thermal distribution. Similarly, the spectra of Mn-doped ZnSe also exhibit lasing under different pump powers and the relationship between the excitation power and intensity was also presented. As the excitation power increased, the intensity ratio also increased in the nanostructure, exhibiting lasing in the doped sample. Normally, II–VI semiconductors doped with TM ions exhibit lasing in the NIR region.^{133,155,239–241} In addition, various studies reported that lasing is owing to the exciton–polariton within the microcavity formed in the structure during its growth or wurtzite structure.^{91,162} In the case of Mn-doped ZnSe NW, the two peaks in its emission spectrum at 466.8 nm and 472.5 nm are assigned to the bound exciton band edge emission and phonon-assisted exciton transition, respectively. A phonon-assisted exciton transition is possible when the excitons and phonons are coherent space or density of states and the same polarization may lead to the generation of a spontaneous polariton–exciton-related emission. In the case of the belt-like structure, the polariton–exciton emission has been reported at 475 nm with a broad

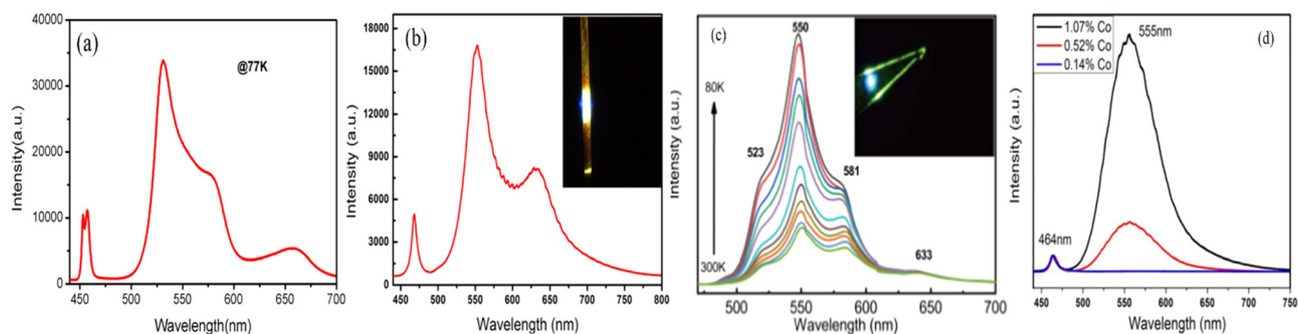


Fig. 7 Transition metal ion doping in ZnSe matrix. (a) Temperature-dependent PL spectrum of Mn-doped ZnSe nanoribbons excited with a CW laser (405 nm) at 77 K, Copyright © 2020, IOP Publishing.²⁰ (b) PL spectrum of an individual Fe-doped ZnSe nanoribbon with a 405 nm laser upon excitation. The inset in the top right corner is the PL image of the nanoribbon in the dark field observation. Copyright © 2017, Applied Science.²¹³ (c) Temperature dependence of the near-band-edge luminescence peak (FXs, EMPs) of Ni₁₂-doped ZnSe NBs from 300 K to 80 K at relatively high (1.2%) doping concentration, Copyright © 2021, IOP publishing.¹¹⁷ (d) PL emission spectra measured at room temperature with different concentrations of Co-doped ZnSe NRs, Copyright © 2021, IOP Publishing.²¹⁶



emission at 650 nm related to the trapped-state emission.²⁴² In the case of ZnSe:Ni nanoribbons, the free exciton lasing emission was dominant at a low excitation power, and as the excitation power increased, the EMP peak began to anomalously increase and became sharp. This anomalous behavior in doped ZnSe:Ni is owing to the nonlinear EMP-related lasing interaction with spin, LO phonons and excitons polarized in the same direction.^{117,230}

3.2.2 ZnS. ZnS is a wide bandgap semiconductor, which normally exists as cubic zinc blende (3.72 eV) and hexagonal wurtzite (3.77 eV) phases at room temperature. The wide bandgap and optical properties of ZnS make it a potential candidate for various applications such as flat panel displays, LEDs, sensors, and lasers. 1D nanostructures of ZnS have been synthesized *via* both physical and chemical methods, which have been extensively used in various applications.^{243–245} Meng and co-workers synthesized ultrafine ZnS NW *via* vapor phase growth in the presence of Au as a catalyst.²⁴⁶ By controlling the substrate temperature, the gold thickness defined the average diameter of ZnS NW, ranging from 10–20 nm. High-density 1D ZnS NWs were grown on anodic alumina oxide (AAO) templates by Ding *et al.* and their luminescence spectra revealed that the stimulated emission originated from the narrow resonant cavity modes.²⁴⁷ Wang *et al.* explained in detail the different morphologies of 1D ZnS nanostructures prepared using the vapor phase mechanism.^{204,248,249} Moreover, solution-processed ZnS nanostructures have been applied in various light emission devices such as flat panels, LEDs and biosensors.^{18,21,245,249} TM ion doping in ZnS nanostructures has been reported by various groups *via* both chemical and physical methods.^{250–253} ZnS structures show high transmission in the visible spectrum and large excitonic binding energy of 40 meV, making them significant materials for photonic and optoelectronic applications.²¹ Xiong's group reported the synthesis of high-crystalline wurtzite phase ZnS NWs *via* the VLS mechanism.²⁵¹ At 10 K, the bandgap emission became dominant with a weaker defect luminescence. In the wurtzite structure, the spin-orbital interaction and splitting of the crystal field near the valence band distribute the exciton energies into three bands (named A (FXA), B (FXB) and C (FXC)), having the split energy among them of 89 meV (FXC–FXB) and 57 meV (FXB–FXA), respectively. In the luminescence spectra, FXA and FXB were clearly observed at room and low temperature, respectively, whereas FXC was not observed at room temperature due to its fast decay time. The excitation intensity clearly showed the change in relative intensity of the exciton- and defect-related emission due to the difference in their density of states. Therefore, FXA and FXB showed a red-shift (8 meV and 12 meV) with an increase in excitation, respectively, but defects were insensitive to the photon flux and were saturated. This shift in exciton emission is due to the laser heating effect in the 1D nanostructure; thereby, the absence of a change with laser excitation is related to the shallow level emission rather than the shallow donor-bound excitons.^{254,255} In the case of heterojunction ZnS/CdS QDs, the trap carriers across the barrier exhibited the activation energy of 18 and 300 meV, which is responsible for lasing due to the ZnS shell acting as a spherical optical cavity, making the transition from linear to nonlinear tendency possible at room temperature.

The temperature gradient showed that (i) the single structure has less impurities or defects (such as Zn vacancies and interstitial S) and (ii) the thermal broadening of the exciton peaks represents the exciton-phonon interaction. Above 120 K, the inhomogeneous broadening is owing to the acoustic-phonon interaction and below 120 K, the LO phonon is responsible for the inhomogeneous broadening (Bose-Einstein condensation fitting). The exciton transition in the ZnS matrix and coupling strength of exciton-LO phonon were directly determined from the broadening of the exciton peak energy and width. These stable and high crystalline NWs are promising materials for the fabrication of high-efficient LEDs and lasers in the UV range.²⁵¹

3.2.2.1 Transition metal ion-related luminescence behavior.

The total energy was calculated for both the spin and non-spin polarized states of ZnS NWs doped with Cr ions, as shown in Fig. 8(a)–(c). Cr(II) interacts with its nearest neighbor S atom by replacing a Zn ion. The p-d hybridization between Cr and S comes from the d-state and the p-states show a small magnetization. The spin density of states or distribution of Cr ions in ZnS NWs shows that the large contribution of the dopant ion to the total magnetization in the opposite direction favors the AFM interaction. According to the Bader charge (BC) analysis of the bond length and charge transfer, the doped NWs are stable due to the stronger Coulomb interaction between the S and Cr atoms than pure ZnS NWs. Moreover, the spin-down shows its semiconducting nature, whereas the spin-up shows that the half-metallic ferromagnetic semiconductor seems to be a good candidate for DMS spintronic applications. The FM state is stabilized through the spin-up of the Cr ion, which is partially filled due to the energy gain from the double-exchange mechanism, but the super-exchange mechanism has no energy gain. The spin-down of the Cr ion shows its semiconducting properties or absence of magnetism behavior. The exchange spin splitting of the d-orbital of Cr ions in the FM state is larger than that in the AFM state due to the double-exchange mechanism. Moreover, the intrinsic vacancy can be overcome by co-doping with non-metals such as N, C, and I at the anion site. The co-doping with N atoms contributes one hole to the valence band, which lowers the Fermi level close to the valence band. According to calculations, the total energy of co-doping at the surface is lower than that in the bulk, indicating that N prefers the surface of NW. In addition, the N-p states are asymmetrical in nature, which are involved in magnetization and favor the AFM coupling of the d-state with p-state of the N and S ions. Therefore, the co-doped atoms are involved in magnetization and the spin density is concentrated around the Cr and N atoms with a lower density around the S atom.

Fig. 8(a)–(c) show the absorption spectra of pure ZnS, the surface configuration of Cr ion and co-doped samples, which show high absorption in the near IR and UV-Vis regions. The surface-incorporated Cr ion showed the d-d transition energy of about 0.95 and 1.53 eV in the near IR region (Fig. 8b). The absorption peak at 2.42 eV corresponds to the transition between the d-state and p-orbital in the valence band. The peak located at 3.05 eV is attributed to the transition between



the interband transition and d-states of the Cr(II) ion. Therefore, the Cr(II) ion reduces the fundamental bandgap energy from 3.49 to 3.20 eV and the slight redshift appears through the sp-d exchange interaction of the host and the localized d-electrons of the transition metal ion. Moreover, the co-doped sample showed a sharp peak at around 1.32–1.85 eV, which is assigned to the defect states of the co-doped samples generated by the d-d transition of the Cr spin ion (Fig. 8c). Therefore, the single Cr ion and co-doping of ZnS NWs decreased the bandgap and e-h pair formation under illumination increased the absorption intensity. According to the optical absorption coefficient, the d-d transitions for the FM and AFM configurations using the electronic structures of the near and far configurations are shown in Fig. 8(a)–(c). To understand the magnetic coupling, we need to understand the optical absorption of the FM and AFM in the Cr-coupled ion system. In the case of the FM configuration, the occupied d-states are below the Fermi level, whereas the unoccupied d-states are above the conduction band with the d-d transitions peak appearing at 0.79 eV and 1.14 eV, respectively. In the case of the AFM configuration, the d-d transition energies are 0.94 eV, 1.32 eV, and 1.53 eV for a single Cr ion in the host lattice. Meanwhile, the AFM and FM configurations are blue- and redshifted in the luminescence spectra, respectively, in various other II-VI semiconductors such as ZnS and ZnSe.^{258,259} It is clear that the optical band gap and d-d transition peaks appear at the same energy level in both the FM and AFM configurations due to the paramagnetic behaviors in the far configuration. The peaks related

to the bandgaps in both FM and AFM appear at 3.23 eV, while the peaks related to the d-band-to-d-band transitions appear at 0.90 and 1.52 eV in both coupled spin systems, respectively.²⁵⁶

TM ion doping in ZnS has been studied extensively to control its optical and electrical properties; therefore, it is of special interest in nanoscale spintronic applications. Various researchers successfully incorporated TM ions (Mn, Ni, Fe, Co, Cu, *etc.*) in ZnS using different synthesis mechanisms.^{260–266} Among them, Mn doping in ZnS has been studied widely due to its phosphorescence and spin-related application. Radovanovic *et al.* used the CVD route to synthesize Mn-doped II-VI and III-V semiconductors.⁹⁷ The calculations reported by Liu *et al.* showed that the strong d-d transition in ZnS as the host matrix is owing to the formation of Mn ion clusters, and thereby the regular distribution of Mn in the host matrix exhibits room temperature ferromagnetism.²⁵⁰ The s, p and d orbital density of states of S, Zn and Mn for Mn@DOS-ZnS in the ferromagnetic ground states and spin exchange within the d orbitals of the Mn ion in ZnS. The calculation results show the ferromagnetic exchange and half-metallic exchange at the Fermi level for the d-orbital, making it very interesting for DMS applications. The half-metallic exchange of the d-orbital present in the electronic structure shows ferromagnetic double exchange interaction and super exchange anti-ferromagnetic interaction at room temperature. Fernandez *et al.* studied the low temperature absorption and luminescence of a ZnS epilayer prepared on a GaAs substrate, where both experimental and theoretical calculation provided suitable information

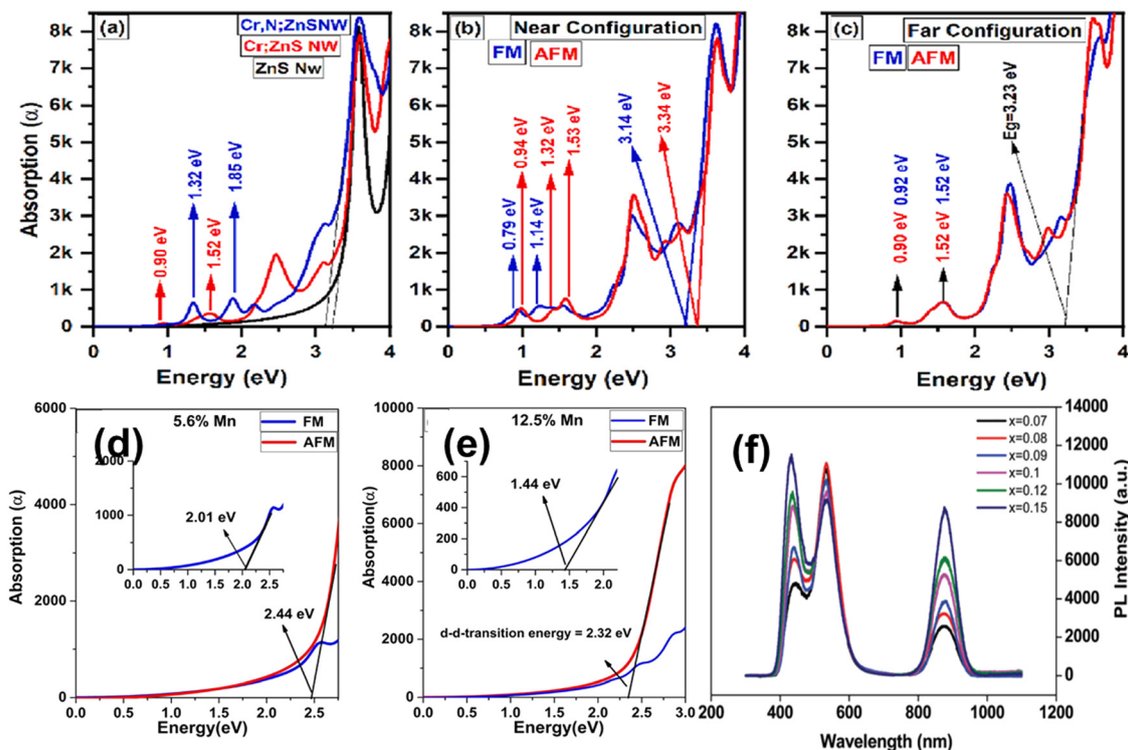


Fig. 8 Optical absorption of (a) pure ZnS nanowire, single Cr(II) ion-doped wire, and single Cr and single N co-doped wires, (b) FM and AFM systems in near configuration, and (c) FM and AFM in far configurations, Copyright © 2023, Elsevier.²⁵⁶ Absorption spectra of FM and AFM coupled Mn ion-doped systems with 5.6% (d) and 12.5% (e) concentration of Mn dopants, Copyright © 2023 IOP, Publishing.²⁵⁷ (f) Room-temperature PL spectra of Zn_{1-x}Mn_xS nanobelts, Copyright © 2017, The Royal Society of Chemistry.²⁵⁸



on the temperature-dependent heavy holes, light holes and split-off band together with accurate information on the temperature-dependent eigen-states, broadening and strain-related parameters, respectively.²⁵⁵ Radiative recombination results in a strong injection rate due to inelastic exciton-polariton scattering. The emission peak at 470 nm is related to the excitonic transition in the UV range from the valence band to the conduction band. Alternatively, the peak at 340 nm corresponds to the defect-related emission and that at 590 nm assigned to the Mn transition. Mn exhibits an excitonic transition band due to charge transfer at 355 nm ($\text{Mn}^{2+} 3d \rightarrow \text{C.B.}$), together with a series of d-d transitions including ${}^6\text{A}_1(6s) \rightarrow {}^4\text{E}(4d)$, ${}^4\text{T}_2(4d)$, $({}^4\text{A}_1, {}^4\text{E})({}^4\text{G})$, ${}^4\text{T}_2({}^4\text{G})$ and ${}^4\text{T}_1({}^4\text{G})$ bands. The transition from ${}^4\text{T}_1 \rightarrow {}^6\text{A}_1$ is the lowest excited state for the Mn transition within ZnS.^{258,264,267}

Fig. 8(d) and (e) show the optical absorption behavior of Mn-doped ZnS with different Mn concentrations. In comparison with pure ZnS, Mn-doping and co-doping resulted in several absorption edges due to the excitonic effect in ZnS. Secondly, the Mn doping resulted in a slight shift toward a lower energy together with a d-d transition at 2.19 eV. In the case of a single Mn ion in ZnS, an electron flip from the spin up to down state with the energy at 2.73 eV was observed. According to the electronic transition, the 3d-state of Mn occupies the top of the valence band and the 4s-state of Zn occupies the bottom of the conduction band. In the case of co-doping (C, Mn), the corresponding transition between 3d (Mn) and 2p (C) near the Fermi level is half metallic. The d-d transition for the Mn-Mn pairs in FM and AFM coupling is obtained from the electronic structure. In the FM configuration, the 3d-states of Mn are fully occupied in the up-spin and the 3d-states are empty in the spin-down. Also, a one-electron spin flip is observed in ZnS with the optical d-d transition energy of 2.01 eV for 5.25% Mn concentration and 1.44 eV for 12.5% Mn concentration in the FM configuration. The spin of one electron is flipped and the transition is from the up-spin to down-spin state. The experimental result for ZnS:Mn NB²⁵⁸ was smaller than that of the single Mn ion-doped system (2.19 eV). In AFM coupling, the filled up-spin state of one Mn couples with the empty up-spin state of another Mn ion. Therefore, the d-d transition for 5.25% Mn concentration is 2.44 eV and that for 12.5% Mn is 2.32 eV in the AFM configuration, supporting the experimental result. Thus, the d-d transition for FM coupled Mn doping is always smaller than that for AFM coupled Mn ion doping, which can appear simultaneously in Mn-doped ZnS NB and ZnSe NB in their luminescence spectra.^{257,258,268,269}

The VLS growth mechanism for the growth of 1D ZnS nanostructures may lead to the formation of twin or stacking faults during the growth of NWs and NB- and NR-like structures.^{244,249,255} Therefore, the defect-related emission spectra of different morphologies are different due to the surface relaxation or surface passivation in different nanostructures. Mn doping in the ZnS matrix with different morphologies also results in different Mn-related peaks, as discussed in detail above for other II-VI materials. Kamran *et al.* showed that the emission from the localized Mn ion requires no contribution from surface capping.²⁵⁸ Fig. 8f shows that Mn-doped ZnS NBs exhibited a green emission due to the dopant together with additional peaks in the range of 420 to 460 nm and 560 to 890 nm.

High-temperature growth *via* CVD normally leads to the formation of stacking faults in ZnS, which show emission peaks in the range of 520 to 550 nm, suggesting the formation of AFM related with Mn-Mn pair formation. In the case of ZnS:Mn NB, the dopant ion in the surface has no contribution, whereas defects can participate in AFM Mn-Mn pair formation with a slight blue shift compared with the emission of a single Mn ion in the ZnS lattice. However, QDs show a strong surface effect, directly influencing the exciton stability and luminescence. In the case of 3% Mn in ZnS NB, its emission band decomposed into four emission peaks located at 452.4, 535.5, 579.8, and 649.3 nm, reflecting the presence of spin-exciton interactions and magnetic couplings. The blue emission located at 452.4 nm corresponds to the anti-ferromagnetic coupling of 4 Mn ions. The green emission located at 535.5 nm shows the interaction of defects with the anti-ferromagnetic coupling of 2 Mn ions. Alternatively, a yellow emission line at 579.8 nm due to the d-d transition (${}^4\text{T}_1({}^4\text{G})\text{-}{}^6\text{A}_1({}^6\text{S})$) of the Mn^{2+} ion was also reported in ZnS:Mn QDs. The orange emission at 649.3 nm corresponds to the ferromagnetic Mn-Mn pair, as reported for CdS:Mn. Moreover, the calculations for ZnS:Mn show that the red- and blue-shifts of the d-d transition correspond to the FM and AFM coupling in the MnS cluster, respectively. Further, an increase in the Mn concentration to up to 5% showed an emission in the near IR region at 877.6 nm. Due to the high growth temperature in CVD, the Mn ions aggregate with a high number, shifting the yellow emission to the red or near IR region, as observed for a doping concentration above 6% due to FM coupling. The blue shift shows the AFM aggregation of the $(\text{Mn})_n$ cluster of different energies in the host lattice. In Mn-doped ZnS NB, the incorporation of 0.2% Mn resulted in the clustering or aggregation of two Mn ions. With an increase in the concentration of Mn ions, multiple emission lines or more Mn aggregates were formed in the host lattice, showing AFM coupling at 535 nm and cluster formation at 460 nm. According to the experimental results, an increase in Mn ion concentration and the growth time can both increase the aggregation of Mn ions in the ZnS lattice with stable clusters, reflecting the origin of ferromagnetism in the ZnS:Mn crystal.

Moreover, the presence of defects due to TM ion doping may lead to metallic behavior at the surface of the nanostructure and the formation of Mn clusters may cause AFM coupling.^{250,266} Therefore, the emission peaks located at 452 nm, 579 nm and 877 nm in the spectra are assigned as the AFM coupling, d-d transition and Mn cluster formation in the NIR region, respectively, as shown in Fig. 8f. Under different composition gradients, the intensity and peak positions in the emission spectra explain possible transition states in the host matrix. Li and co-workers reported the appearance of the Mn-related defect emission at 585 nm for ZnS NRs.²⁶⁷ Kang *et al.* reported that Mn/Fe- and Co-doped ZnS synthesized *via* CVD showed room temperature ferromagnetism.²⁷⁰ Wang *et al.* fabricated an Mn-doped ZnS sea urchin-like structure *via* the solvothermal method, and also assigned the emission peak at 587 nm to the Mn^{2+} ion in the host matrix.²⁷¹ Therefore, TM ion doping in the ZnS matrix can tune its electrical and optical properties for spintronic applications.



3.2.3 ZnTe. In many respects, the inherent p-type conductivity of ZnTe makes it the most attractive wide bandgap material in the Zn-family and its superior optical properties are due to its direct bandgap (2.26 eV) and Bohr exciton radius (6.2 nm), showing potential for various optoelectronic applications.^{272–276} Its PL emission spectrum at 13 K shows the splitting of a single peak into multiple peaks, providing suitable information about the host matrix-related exciton emission. The emission peak at 525 nm is related to the band edge emission without Y-emission (600–700 nm) for high-quality nanostructures obtained *via* the VLS mechanism. Moreover, the Y-emission also shows a strong intensity with the incorporation of impurities in the ZnTe matrix due to the presence of the strain effect. The degeneracy of the conduction and valence bands is 2-fold (Γ_{6c}) and 4-fold (Γ_{8b}) in zinc blende ZnTe, respectively. The large spin–orbital splitting at the topmost valence band is assigned to three exciton states, whereas strain may split the exciton in the topmost valence band into free exciton heavy holes and light holes. A ZnTe epilayer showed the free exciton heavy hole and light hole at 2.379 and 2.375 eV at 2 K, respectively.^{277–279} Utama *et al.* reported the presence of free exciton heavy hole and light hole at 2.377 and 2.372 eV, respectively. However, the bound excitons became dominant at low temperature and their emission peak observed at 2.375 eV due to the neutral acceptor caused by defects or extrinsic impurities. In the case of room temperature free excitons, their emission was more prominent than that of bound excitons, where the emission of the bound excitons decreased with an increase in temperature due to the ionization effect. In this case, the emission at 2.368 eV (intrinsic shallow acceptor levels; Zn vacancy) is lower than that of free exciton heavy hole at low temperature due to the following reasons: (i) absence of elemental doping and (ii) less stoichiometric-related defects during VLS growth. ZnTe is a polar semiconductor that exhibits strong exciton–phonon interaction at low temperature due to resonance phenomena. The LO phonons are coupled with free exciton light holes, leading to the emission of higher-order LO phonons in the ZnTe matrix, which has also been reported for CdS. Thus, the decay rate of excitons is highly enhanced due to the hot-exciton process.²⁸⁰ The band edge emission of ZnTe is in the visible spectrum (green emission) without the incorporation of any impurities, making it suitable for the fabrication of visible optoelectronic devices such as LEDs and lasers.^{274,281,282} The unique physical properties of ZnTe make it interesting for the fabrication of monolithic integrated optical circuits of signal transmission and processing, exhibiting the best electro-optical figure of merits among semiconductors, terahertz generation and detection together with very favorable photodetection properties in various device applications such as p-type field effect transistors (FETs).^{281–283} Wide bandgap semiconductors such as ZnTe have strong exciton-related emission in comparison with narrow bandgap semiconductors such as CdSe and CdTe, in which a band edge emission often arises from separate carriers after photo-excitation. Therefore, the carrier effect in ZnTe has also been reported by researchers, which may cause a redshift in its emission spectrum as a function of the incident intensity related with the surface plasmon effect.^{284–286} These phenomena may depend on the growth conditions, synthetic mechanism and quality of final product.

3.2.3.1 Transition metal ion-related luminescence behavior. The TM ions in II–VI semiconductors (based on tellurides) are well-known diluted magnetic semiconductors (DMS), where the substitution of the cations in the host semiconductor by TM ions is appealing for future spintronic devices. Most research groups used the molecular beam epitaxy (MBE) technique to substitute TM ions (such as Mn and Cr) into ZnTe for spintronic applications.^{6,79,105,287} Other methods have also been employed for TM ion doping in ZnTe, as reported by various groups.^{2,284,285,288–290} PL is an effective and non-destructive technique to determine the bandgap and other related information such as impurities in the host matrix to find the origin of the transition in the spectra. Fig. 9c shows the lasing in a single belt-like structure of Fe-doped ZnTe prepared *via* the VLS mechanism at room temperature. The room temperature PL spectra show the NIR emission for Fe-doped ZnTe. Angle-resolved measurements revealed the formation of a cavity in the microstructure due to the presence of Zn vacancy, which exhibits strong luminescence in the NIR region at room temperature. Moreover, fs laser excitation exhibits the near band edge lasing but no emission can be seen in the NIR region as observed by continuous laser excitation. This indicates that the exciton-defect-related emission is due to the slow/fast coherent recombination process occurring in a single structure.²⁹⁰ Moreover, according to the electronic configuration of Ni doped in the ZnTe matrix, Ni^{2+} is tetrahedrally coordinated by four anions (T_d symmetry point). The T_{2g} orbital of $3d^8$ of Ni^{2+} interacts with the p-orbital valence band to form strong sp–d hybridization, which breaks the optical selection rule and shows a d–d transition in the emission spectrum. Moreover, the PL emission spectrum reveals a redshift due to the following reasons: (i) radiative recombination of excitons or photo-induced carriers, (ii) bandgap renormalization effect, (iii) carrier–carrier scattering and (iv) e–h plasma formation at the surface under a high laser power.^{285,291} Another factor that causes a redshift is the structure-dependent surface effect, which can enhance the carrier–carrier scattering, leading to the formation of e–h plasma at the surface. The laser spot size on rod- and belt-like structures exhibits the greater surface e–h plasma effect in the belt-like structure due to the fact that its surface acts as a reflector due to the Zn deficiency or Te inclusion on its surface. The polarized PL emission of both Mn- and Ni-doped samples provides suitable information on their structure, suggesting that the zinc blende structure is obtained during high temperature growth *via* the VLS mechanism.²⁸⁵ Normally, the high temperature growth of ZnTe leads to the formation of a cavity, which exhibits an NIR emission, and the literature suggests that the cavity formation is normally observed in the wurtzite structure. In the case of the zinc blende cubic structure, the possibility to observe the cavity-related NIR emission is related to the twinning or polymorphs in the structure during high temperature growth, as reported in ZnTe.^{52,291–293}

Fig. 9a and b show the first-principles calculations study of the magnetic and opto-electronic properties of Fe(II)-doped ZnTe based on the RKKY-exchange. Fe(II) occupies the tetrahedral crystal field, showing a spin-allowed d–d transition in the IR region. The optical bandgap reveals a blue shift due to



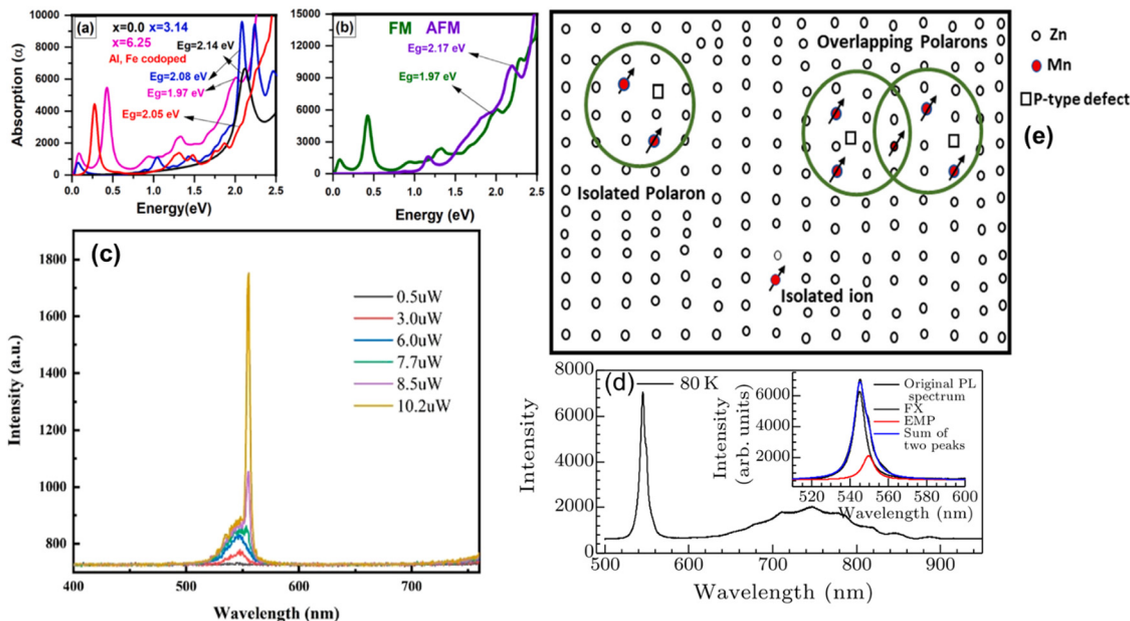


Fig. 9 Absorption spectra of (a) single Fe doped, double Fe doped and (Al, Fe) co-doped ZnTe and (b) FM and AFM coupled ion systems. Copyright © 2023, Elsevier.² (c) PL emission spectra of ZnTe microbelt under femtosecond pulse excitation with a variation in pump power. Copyright © 2023, the American Chemical Society.²⁹⁰ (d) PL spectra of Co-doped ZnTe at 80 K with Lorentz fitting. Copyright © 2018, Chinese Physical Society.²⁸⁹ (e) Schematic diagram of bound magnetic polaron model for Mn(II)-doped ZnTe, Copyright © 2023 Elsevier.¹

AFM coupling, whereas the FM coupling results in a redshift in the emission spectrum. The Fe ion in the host lattice has no gain in energy due to the double exchange of the spin states (e_{2g} and t_{2g}) that are fully occupied, supporting the FM interaction. Similarly, the two spin ions of the same spin gain energy due to the super-exchange of half-filled e_{2g} support the AFM interaction. The exchange splitting of the d-state in the FM state is smaller than that in the AFM interaction and the energy gained due to the super-exchange interaction is larger than that due to the double exchange interaction. In the case of the incorporation of an n-type dopant ion in the host lattice, it occupies the spin down partially filled e_{2g} state, which supports the FM configuration. Therefore, the large exchange splitting of the d-state supports that the FM interaction with double exchange is larger than the super-exchange, which favors the FM state in n-type Fe(II)-doped ZnTe. The incorporation of Fe ions in ZnTe and co-doped with Al ions is responsible for the intermediate d-states between the conduction and valence band. The absorption peaks of the Fe ion associated with the intra-band transition appear at 0.2, 0.92 and 1.03 eV, corresponding to the electronic transition of the E_2 to T_2^3 and E_2 to T_2^3 states, respectively. With an increase in the number of Fe ions, the number of d-states also increases, with the appearance of additional d-d transition peaks due to the Fe-Fe pair system, corresponding to the transition states of $E_1^2-E_2^2$, E_2^2 , $E_1^2-E_3^2$, $E_1^2-T_1^3$ and $E_1^2-T_2^3$ having energies at 0.25, 0.45, 0.88, and 1.31 eV, respectively. The d-d transition intensity increases with an increase in Fe concentration but Al co-doping reduces the d-d transition together with the band-edge transition peaks. Moreover, an increase in the concentration of Fe(II) results in a redshift in the band-edge transition and d-d transition

related with the Fe-Fe ion through sp-d exchange coupling. When Al is co-doped in ZnTe, the Fe(II)-Fe(II) ion pair shows a blue shift in its bandgap due to the Burstein-Moss effect. Additionally, in the Fe(II)-Fe(II) ion pair system, the intensity of the absorption coefficients increases with an increase in the Fe concentration, which decreases after co-doping. According to the density of states peaks, the FM and AFM configurations show the occupied and unoccupied d-orbital states for both Fe(II) and Fe(II) co-doped in ZnTe. The d-d intra-band transition exhibits a red and blue shift for the FM and AFM coupled systems, respectively, with different energies showing the influence of the different types of magnetic coupling in the absorption spectra, as confirm by both theoretical and experimental data. The improvement in the opto-electronic and magnetic properties of Fe(II)-doped ZnTe upon n-type (Al) co-doping provides insight into the potential applications of these materials in spin-related photonic and spintronic devices such as photovoltaics and remote sensing.²

Dong *et al.* showed that the terahertz generation by optical rectification in ZnTe is usually degraded by the following factors: (i) harmonic generation, (ii) two-phonon absorption and (iii) free carrier absorption.²⁹⁴ The inclusion of Te in ZnTe for THz generation was observed by IR transmission spectroscopy. The Te inclusion has a hundreds of picoseconds region and wider width, while that for the ZnTe matrix is several picoseconds. Alternatively, the central frequency and bandwidth of the Te inclusion are 0.1 and 0.6 THz, whereas they are approximately 0.2 and 1.8 THz for ZnTe, respectively. Additionally, the amplitude of the THz waves from the Te inclusion at the center is higher than that from the ZnTe matrix. The Te inclusion in the ZnTe matrix is responsible for



the unique emission, which may enhance the multiple reflection of THz pulse. This multiple reflection enhances the phonon polaritons within the host matrix, which is related with the second harmonic generation.^{294,295} The energy transfer mechanism based on the anti-Stokes emission is beneficial for various technological applications including biological imaging, up-conversion lasers, solar cells and many others.²⁹⁶ Excited state absorption (ESA) showed that the electron transition involves the use of two absorbed photons in intermediate electronic states to create highly excited carriers. Energy transfer induced up-conversion (ETU) is similar to the Auger effect, in which energy transfer occurs from an excited e-h pair to another carrier. Two-photon absorption (TPA) is the simultaneous absorption of two photons in the intermediate state to achieve a high electron transition. The TPA-related emission shows a large anti-Stokes shift without acquiring a real intermediate state at a high excitation density on the nanoscale timescale. Thus, a high density and shorter timescale are key factors in TPA-related linear upconversion absorption. Phonon-assisted anti-Stokes PL (PA-ASPL) only appears where strong electron-phonon interaction occurs at a very low excitation power. The phonon-related anti-Stokes emission involves an indirect optical transition due to the hot exciton effect in polar semiconductors, as discussed above for both ZnTe and CdS nanobelts. Excitons are strongly coupled with optical phonons through deformation potential interaction or Frohlich interaction, which is responsible for the higher order Raman peaks due to resonance phenomena.

Fig. 9e presents a schematic of the BMP model responsible for the ferromagnetism in p-type Mn(II)-doped ZnTe. Different mechanisms, such as double exchange mechanisms, bound magnetic polarons, super-exchange mechanisms, and band-coupling models, have been used to explain the ferromagnetism in DMSs. The most widely considered defect-mediated mechanism for understanding ferromagnetism in dilute magnetic semiconductors is the bound magnetic polaron model. Zn vacancies play a crucial role in understanding the magnetism in Mn(II) in ZnTe, where the 3d states of the Mn ions around the defects couple ferromagnetically to each other, where the hole carriers are bound to form BMPs. The polaron-polaron interaction is ferromagnetically coupled with the carrier-ion-carrier interaction. The paramagnetic (PM) component is attributed to the isolated polarons and Mn(II) ions. In addition, the co-dopant and Zn vacancy boost the room-temperature ferromagnetism. Furthermore, we inferred that the BMP model (defect-mediated model) fit can help understand the origin of ferromagnetism. Our optical study revealed that Mn(II) doping at the Zn site widens the bandgap of ZnTe and produces spin-forbidden d-d transition peaks on the low energy side of the bandgap. The p-type defects in Mn(II)-doped films produce absorption peaks in the infrared region and improve the absorption efficiency. In addition, the optical bandgap and spin-forbidden d-d transition of Mn(II) to different modes of spin-spin interactions were correlated, and we observed a red shift in the d-d intra-band transition peaks and an optical bandgap in the FM-coupled Mn(II) ion system and a blue shift in the AFM coupled ion system, supporting the experimental observations. The theoretical calculation of Mn(II)-doped ZnTe showed that the occupied spin

level of Mn leads to a super-exchange mechanism in p-type semiconductors. The incorporation of Mn leads to ferromagnetism through additional hole carrier formation with the dopant ions. Moreover, the Zn vacancies and co-doped coupled the hole carriers with the spin-up of the Mn(II) ion with stable ferromagnetism behavior. The hybridization of the Te-p level and Mn(II)-d level around the Fermi level supports the stabilized AFM of the d-level due to its full occupation by the spin electron after gaining energy to support the super-exchange mechanism. The double-exchange mechanism appears with no energy gain, which is located just below the conduction band or conduction state minimum. Meanwhile, the overlapping spin-up and spin-down show no magnetism in the AFM coupled ion system. The interaction between two-spin ions with opposite shifts at the d-orbit show the exchange spin splitting of the large AFM state than the FM states. Therefore, the origin of AFM coupling in the ZnTe slab-like structure is due to the super-exchange of electron hopping occurring between the Mn ions *via* the intermediate p-state of the anion. The experimental results reveal that the hole carriers play a vital role in the ferromagnetism in DMS together with the defect states and co-doping enhances the p-type behavior in semiconductors. In general, the FM state is highly stabilized with additional hole carriers, which are localized at the topmost valence band in ZnTe with broadening due to the symmetry breaking relaxation. Secondly, the Zn vacancy acts as a double acceptor, which generates two holes in the spin-up states by charge transfer in the coupled ion systems. The states split into occupied and empty states in the majority spin channel due to the transfer of charge from the Mn-d state to the defect level of the Zn vacancy. The energy difference is about 135 meV greater than the thermal energy, indicating that the FM state is more stable than AFM above room temperature. The magnetic moment is contributed by the following: (i) Mn(II) ion doped in ZnTe or Mn-Mn interaction, (ii) hybridization between Mn-d and Te-p orbitals with minor magnetic moment and (iii) spin distribution of Mn(II) with Zn vacancy. A large density of spin exists around Mn(II) and the interaction of the Mn ion with the nearest Te atoms is antiferromagnetic. Moreover, the co-doping also enhances the ferromagnetism behavior given that it acts as double acceptor with an energy difference of about 138 meV lower than the AFM state.¹

Fig. 9d shows that Co-doped ZnTe exhibits a free exciton emission together with a shoulder peak with the energy difference of 19.4 meV less than the LO phonon at 80 K assigned to the EMP emission. The aggregation of the Co ions results in ferromagnetic spin, which binds with the free excitons to form EMP. Moreover, the broad emission in the range of 600–900 nm corresponds to the d-d transition associated with ${}^4T_1(P) \rightarrow {}^4A_2(F)$ of the ferromagnetic-coupled Co ion in the host lattice from the high d-level of the intra-band transition. Moreover, the power-dependent emission reveals a redshift in the shallow-trapped states (564 nm) due to the generation of a high carrier concentration, which normally appears in low excitonic binding energy semiconductors such as ZnTe (12.6 meV). The high excitation power generates a large number of photoinduced carriers, which are responsible for electron-hole plasma formation on the surface. Secondly, the large radius of the photons and



excitons increases the possibility of electron–phonon coupling an increase in electronic scattering, which is responsible for the large red shift at 563 nm. Previous studies deduced the formation and emission of EMP in TM ion-doped II–VI semiconductors, specifying the emission at the long-wave side of FX. A double-peak structure is observed in the range of 540–580 nm and the emission of EMP appears due to the coupling between the free excitons and ferromagnetic spins from the transition metal ions. The emission peak at 572 nm is related with EMP, which corresponds to the phonon and spin interaction. The redshift in the EMP emission indicates that the carrier interaction is dominant and the increase in intensity shows that the strong spin binding effect at a high power is due to the large polaron radius. Therefore, the EMP becomes more dominant even at a high concentration, providing stability to the spin–spin interaction even at room temperature and high excitation power.²⁹⁷

Ni(II) ion doping in ZnTe results in the generation of magnetic polarons together with the hot carrier effect, which can be understood based on power and temperature-dependent PL. The spin–exciton, spin–spin coupling and carrier–phonon interaction emission account for its ferromagnetic properties. The power-dependent luminescence behavior responsible for the hot carrier effect suppresses the charge trapping effect at a low exciton binding energy (8.12 meV). The large polaron radius exhibits the strong interaction between the carrier and magnetic polarons, resulting in an anharmonic effect in which the side-band energy overtones to the LO phonons. The photon-like polariton shows the polarized spin interaction with LO, exhibiting strong spin–phonon polaritons. Multiple emissions show the non-uniform distribution of the Ni ions in the ZnTe nanostructure with a strong exciton emission at the near band edge and lower energy side due to the ferromagnetic coupled Ni pair or cluster interaction with the traps/defect states. The localized spin-induced excitons interact with hole exchange (heavy hole greater than electron) at the topmost valence band in ZnTe to form LEMP. The non-radiative decay of localized spin/excitons is responsible for multiple emissions at room temperature related with photon-like polaritons. The optically polarized anisotropy explains the dielectric contrast due to the degeneracy of the exciton states at the topmost valence band, representing the spin–spin coupling and exciton–spin–phonon and electron–phonon interaction of magnetic polarons of low excitonic binding energy semiconductors for future spin-based optoelectronic devices. The free excitons (FX) interact with the 2LO phonon–spin interaction, corresponding to the $^3T_1(^3F) \rightarrow ^1T_1(^1G)$ and EMP peaks with ferromagnetically coupled Ni ions at $^3T_1(^3F) \rightarrow ^1E(^1G)$. In addition, the other d–d transition of single Ni ions (600–900 nm) appears at the low energy side with an energy shift of 14–38 meV due to the localized states. These results demonstrate the spin–spin magnetic coupling and spin–phonon interaction at room temperature, making the new horizon of optically controlled dilute magnetic semiconductor applications more realistic.²⁹⁸

3.2.4 ZnO. Zinc oxide semiconductors with a hexagonal wurtzite structure, direct bandgap (3.37 eV) and large exciton binding energy of 60 meV have been widely used as transducers, transparent conduction electrodes, solar cells, and wide ultraviolet

(UV) optoelectronic devices. 1D ZnO nanostructures exhibit high charge transport, mobility, electron affinity, and thermal stability, making them attractive for solar cells and other electronic devices, which have been synthesized by both physical and chemical methods.^{299–303} ZnO NW layers used as the transport layer in hybrid solar cells based on poly(3-hexylthiophene) (P3HT)/ZnO composite are benchmark systems that have attained high power conversion efficiencies.^{25,304} ZnO also exhibits a strong electron–phonon interaction, which may lead to the formation of multiple phonon emissions or hot exciton-related emissions that can be observed in absorption, PL and scattering spectra. Zou *et al.* discussed the exciton coupled with magnetic ions in micrometer-sized DMS in ZnO and CdS. They found that the ferromagnetic and optical responses strongly depended on the size, composition in the local structures, and nature of the TM ions. In the case of the TM-doped II–VI semiconductors, QDs show confined excitons to form EMP in QDs. Moreover, the size-dependent quantum confinement revealed giant Zeeman splitting, photo-induced magnetism and anomalous temperature-dependent EMP emission. An increase in the TM composition gradient in the QDs resulted in an energy shift and enhanced the spin–orbital coupling and spin–spin coupling, which enhanced the EMP formation without LO phonons. In the case of 1D nanostructures or other effects than the quantum confinement effect, the exciton interaction with spin changes due to the long-range coherence of EMPs. The luminescence behavior showed that the condensation of EMP formation occurs in higher dimensions, corresponding to the collective spin influencing the electrical and optical response. In this case, II–VI semiconductor NWs and NBs prepared *via* the CVD mechanism for DMS highlighted their long-range exciton–spin interactions and optical behaviors even at a low concentration. The spontaneous migration of the TM ions resulted in cluster formation during high temperature growth with no new phase in the host matrix.⁵ Dietl *et al.* predicted that the transition temperature (T_c) in p-type ZnO DMS will be greater than 300 K, which can possibly be used for the fabrication of various room-temperature spintronic devices.^{6,7,105,305}

3.2.4.1 Transition metal ion-related luminescence behavior.

Fig. 10a shows the power-dependent PL spectra of ZnO:Mn NW prepared *via* CVD upon excitation by a 355 nm fs laser. At a low pump power, the free exciton peak appears at 3.305 eV with another peak at 3.168 eV ahead of the free exciton peak. The energy difference between the two peaks is about 137 meV, corresponding to the overtone of the A1 2LO mode originating from the polaronic exciton of a bound exciton, which involves spin–phonon coupling. A further increase in excitation power resulted in a superlinear increase in peak B, as shown in Fig. 10a. Also, peak A, which is related with free excitons, slightly increased and decreased swiftly when the power increased from 0.3 to 3 μ J. Fig. 10b and c show the intensity and FWHM as a function of excitation power, respectively. In the high power range, peak B became sharp with a reduction in FWHM. The lasing line of peak B is blue shifted by about 50 cm^{-1} and analogous to the phase space filling effects of excitons at high excitation, which represents exciton repulsion, not a carrier effect. Above 4.0 μ J, the lasing line narrowed by around 3 meV, which reflects the



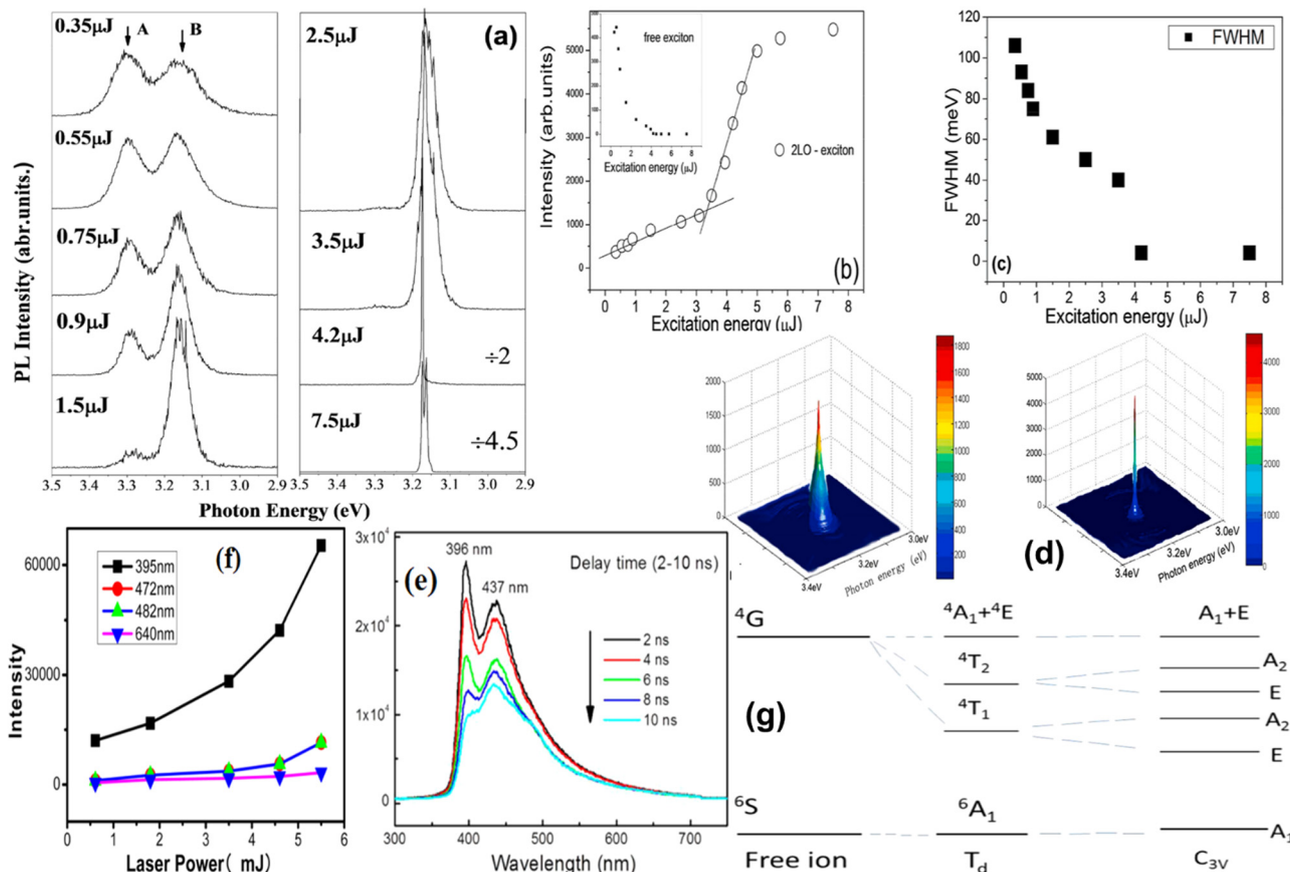


Fig. 10 (a) Photoluminescence spectra at various pumping energies of ZnO nanowires. (b) Luminescence intensity vs. laser excitation energy. (c) Luminescence B band FWHM vs. laser excitation energy. (d) Energy profile of the luminescence B line at 3.5 μJ and 4 μJ . Hot luminescence spectra of Mn-doped ZnO NWs. Copyright © 2011, IOP Publishing.³⁰⁷ (e) Time-delayed photoluminescence of ZnO:Mn NW under a laser pulse at 355 nm (duration 2 ns) from the third harmonic of an Nd:YAG laser with the pumping energy of 3.5 mJ. Observed spectral variation after the NW was excited within 2–10 ns delay time range. (f) Dependence of the emission intensity of various wavelengths on the excitation powers. (g) Diagram of Mn(II) levels in ZnO matrix. ${}^6\text{A}_1$ is the ground state and ${}^4\text{T}_1$, ${}^4\text{T}_2$, ${}^4\text{E}$, and ${}^4\text{A}_1$ are the excited states in ZnO, respectively. Copyright © 2014, the American Chemical Society.⁴

complete coherent emission with an increase in intensity, as show in Fig. 10b. A further increase in power led to the formation of an additional line in the spectra, corresponding to the dipole-dipole interaction of excitons from the coherent single mode. Fig. 10c shows the single mode lasing of ZnO:Mn NW with the narrow FWHM related with BEC excitons. The low doping profile of ZnO NW showed an emission at a low excitation power due to exciton coupling and electron-phonon coupling at room temperature for the large exciton binding energy (60 meV) semiconductor. Alternatively, a high concentration of Mn ions in the host lattice may weaken the UV emission at low power due to the trapped states within the bandgap. The emission shows the non-linear phonon-phonon coupling in 1D NS with an enhancement in A_1 phonons along the axial direction. This strong non-linear coupling may lead to the formation of a 2LO phonon interaction with excitons, exhibiting the boson nature of bipolaron excitons in the 1D system at about $4 \mu\text{J cm}^{-2}$. The increase in intensity and narrow FWHM reflect the coherent two-phonon-exciton formation and condensation, which lead to BEC states at room temperature for ZnO:Mn. The stable BEC in the 1D system is related with the following: (i) coherent and high-density exciton

states produced by fs laser, (ii) side crystal face reflection may enhance the high-density exciton, and (iii) the aggregation of free excitons with different orientations assisted by coherent 2LO A_1 phonons in the longitudinal direction. Therefore, the high-density bipolaronic exciton emits coherently, reflecting the characteristic of the dynamic BEC statistics. Fig. 10d shows the 3D contour plot of the coherent exciton emission for different laser powers. The emission spectrum of Mn-doped ZnO clearly reveals that the Stokes signal interacts with the LO phonons because electrons interact more strongly with LO phonons rather than other phonons due to the deformation potential interaction.³⁰⁶ In the case of ZnTe, the LO phonon undergoes relative displacement due to its oppositely charged ions, which induce polarization, strongly interacting with electrons through Frohlich interaction.²⁹⁶ Mn-doped ZnO NWs exhibit strong electron-phonon interaction, as observed for the eleven order of LO phonon modes, in which the deformation potential of the dopant ion influences the atom displacement from the equilibrium position. Therefore, the displacement of the atom enhances the lattice polarity and shows multiple phonons due to the strong electron-phonon interaction in Mn-doped ZnO NWs. Secondly, the slight shift towards a higher wavelength is due



to the sp-d hybridization between the localized d-electron of the dopant atom and VI-group anion, as observed for other II-VI semiconductors. Thus, magnetic ion doping quenches the band edge emission owing to the radiative absorption by the color center, which highlights the high-order phonon peaks in the hot luminescence process under 325 nm and 355 nm.³⁰⁶

TM ion doping in ZnO is important for DMS and their structure and magnetic origin need to be understood. Liu and co-workers explored the magnetic exciton relaxation and spin-spin interaction of Mn-doped ZnO NW prepared *via* the CVD method.^{4,308} Fig. 10e shows the time-delayed PL of a ZnO:Mn single NW under a laser pulse at 355 nm (duration of 2 ns) from the third harmonic of an Nd:YAG laser with the pumping energy 3.5 mJ. The spectral variation was observed after the NW was excited within a 2–10 ns delay time range. The emissions at 396 nm and 437 nm were observed at a low excitation power, and as the delay time was extended, a shoulder peak appeared at 480 nm together with a shift in the peak at 396 nm to 402 nm. At a high delay time of about 28 ns, the emission peak 472 split into multiple peaks and became sharp under a further delay with the emission of another peak at 636 nm, which is the Mn-related emission. Fig. 10f shows the dependence of the emission intensity of various wavelengths on the excitation powers. According to the results, we conclude that each excitation emission in the luminescence spectrum is time dependent and related to the electronic configuration, carrier-carrier interaction, carrier coupling with spin ion interaction, carrier-phonon interaction and exciton-phonon interaction in the respective host material crystal symmetry.^{309,310} All the emission peaks observed under different time domains show an increase in the superlinear intensity with laser excitation power, as shown in Fig. 10f. In the case of 1D nanostructures, these phenomena become more interesting, which are specially related to the exciton and exciton-spin relaxation characteristics. Initially, the exciton-related emission in the spectrum exhibits that further recombination with phonons, impurities or magnetic ions occurs, as observed by a fs laser producing free excitons and bipolaron excitons in Mn-doped ZnO NW.³⁰⁷ Moreover, the excitation in ZnO is orientation dependent, as discussed above for vertically aligned and horizontally aligned NW, revealing different quantum efficiencies. Thus, the intermediate states with a lower energy are populated after the relaxation of photoinduced carriers.^{5,310} The emission at 394 nm for a 2 ns pulse is assigned as the bi-polaron exciton (BPE) or EMP.^{4,307} According to the schematic diagram (Fig. 10g), the d-d transitions from the 6A_1 ground state to the different intermediate 4T_1 , 4T_2 , 4E , and 4A_1 excited states in ZnO were calculated to be 2.55, 2.85, 2.97, and 2.99 eV, respectively, through CI energy band calculation and photoemission spectra.^{4,311} Therefore, both the theoretical and experimental correlations provide suitable information about the emission spectra of the ZnO crystal symmetry matrix with and without doping at room temperature. Also, 1D-Mn doped ZnO NWs show coherent emission localized exciton magnetic polarons, which are the source of ferromagnetism in the host matrix to produce coherent EMP emission-related different time domains. This type of 1D single structure is significant for both spintronic- and spin-related

photonic devices. The epitaxial growth of a ZnO NW array on sapphire *via* the chemical vapor transport method and determination of the external quantum efficiency related to the PL of ZnO NW at room temperature have been reported.^{301,312} Interestingly, the bandgap-related external quantum efficiency (EQE) is strongly dependent on the excitation power density and spot size. In the case of EQE, it is necessary to understand the absorption first, where the band edge emission improves or becomes more intense with an increase in the incident laser power rather than the exciton-related radiative recombination becoming dominant. Alternatively, the defect-related emission becomes more intense than the near band edge, indicating that the non-radiative recombination related with the defect state is dominant in the single structure.

3.2.5 Ternary alloys of zinc chalcogenides (ZnX; X = S, Se, Te, O). Ternary alloys of Zn chalcogenides with a tunable composition for bandgap engineering have been prepared *via* the VLS mechanism and solution process method. A variation in the composition of II-VI semiconductors causes optical inhomogeneity, which may lead to the formation of more scattering and screening of charges but excitons still play a vital role in lasing when excited by two-photon pump lasing for $ZnS_{1-x}Se_x$ nanoribbons.³¹³ Quantum confinement has a predominant effect on optical gain media including tunable emission in a broad spectrum, low threshold lasing and temperature insensitive lasing performance on the nanoscale. In the case of low-dimensional systems, stimulated emissions and lasing are quite challenging to achieve, but the synthesis of high-quality colloidal QDs can overcome this problem. In the case of one-photon pumping in colloidal QDs, it has poor spatial resolution especially in the biomedical field. Therefore, the two-photon or multiple photon absorption method has been employed to investigate biophotonics without strong scattering from tissue or biological samples. ZnS/CdS/CdSe colloidal QDs were excited with one-, two- and three-PA (photon absorption) with fs laser pulses at 480 nm, 800 nm and 1300 nm, respectively. The stimulated emission of CdSe/CdS/ZnS core-shell QDs excited with 1300 nm was reported.²⁷¹ Fig. 11a shows the photoelectron transfer diagram of $Zn_xCd_{1-x}Se$ and $Zn_{0.9}Cd_{0.1}Se/GQD$ nanocomposites prepared *via* the hydrothermal process. The Zn/Cd ratio can alter the crystal structure from wurtzite (CdSe) to cubic (ZnSe) with a high current photocurrent for a ratio of 0.9/0.1.³¹⁴ With composition gradient changes within the single structure, the emission line was also tuned depending on the Se or S composition modulation. As the fs laser excitation increased, the lasing phenomena were also studied above the threshold value of 1.2 to 5.13 mJ cm⁻² for ZnSe to $Zn_{0.43}Se_{0.57}$ under different excitation powers. In the core-shell structure, ZnTe was used as the type II material with CdSe or ZnSe to cover the visible spectrum range. The ZnSeTe QDs emitted blue color to green emissivity by controlling the Se/Te ratio. However, the realization of red-emission becomes highly exciting mainly due to the strong band gap bowing between the bulk ZnSe and ZnTe. An identical triple shell with ZnSe inner and ZnS outer layers showed PL emission at 608 nm with a quantum yield of 55%. An enhancement in the PL performances was realized with an interlayer of ZnSeTe with a



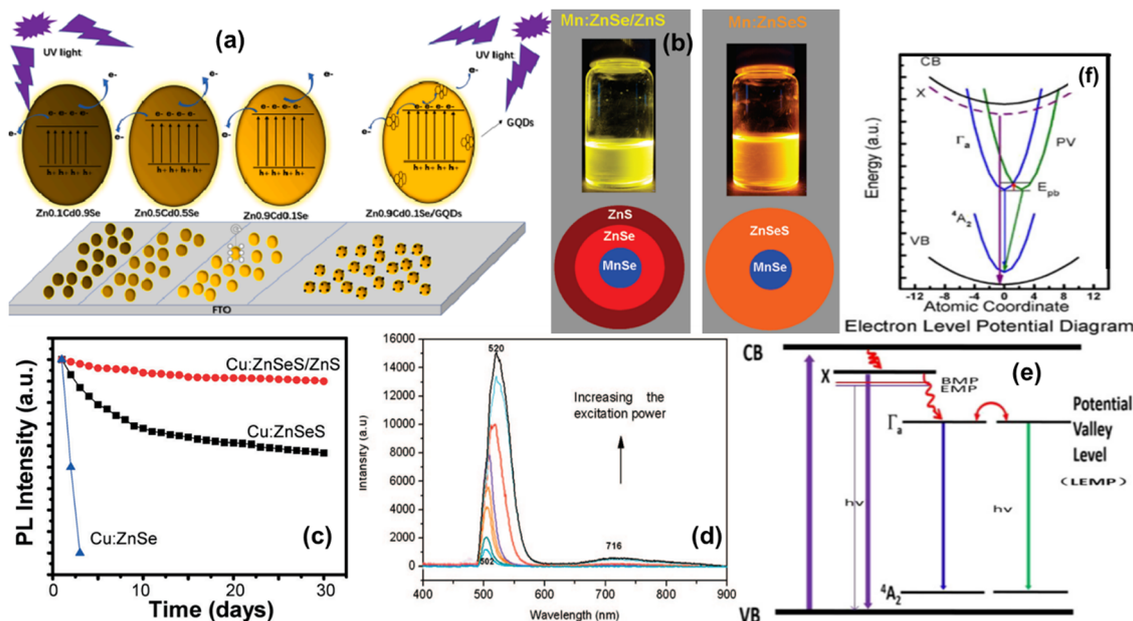


Fig. 11 (a) Photoelectron transfer diagram for $Zn_xCd_{1-x}Se$ and $Zn_{0.9}Cd_{0.1}Se/GQD$ nanocomposites, Copyright © 2022, Elsevier.³¹⁴ (b) Digital image of $ZnSe/ZnS:Mn$ (left) and $ZnSeS:Mn$ (right) upon exposure to a UV lamp, Copyright © 2011, ACS publishing.³¹⁷ (c) The photostability curve plot of $Cu:ZnSe$, $Cu:ZnSeS$ and $Cu:ZnSeS/ZnS$ d-dots in air and light, Copyright © 2011 The Royal Society of Chemistry.³¹⁸ (d) PL of 6-fold symmetry branched nanostructures using Ar^+ 488 nm as the excitation light source at room temperature, Copyright © 2008, ACS.³¹⁹ (e) Schematic of recombination dynamics of excitons (X), EMP or BMP near band edge at high excited state to ground state of d-d transition of Co^{2+} ion (blue color) and STE (green color) within the bandgap of $Co(II)$ -doped $ZnSe$ NR. (f) Electron level potential diagram with atomic coordinates in Co -doped $ZnSe$ NR, Copyright © 2021, IOP publishing.²¹⁶

lower Te/Se ratio as the interlayer, which not only improved the PL emission but also improved the quantum yield up to 63% with a slight redshift. Employing Te as the interlayer has potential application for red emission in display devices.³¹⁵ $ZnTe/CdSe/(Zn, Mg)Te$ core/double-shell NWs grown by MBE show an emission corresponding to the spatially indirect exciton recombination at the $ZnTe/CdSe$ type-II interface. Thus, it is significant to study the blue shift in the emission energy with an increase in excitation fluence, which induces electron-hole separation at the interface. The optical emission in the NIR region originates from NWs and not from the 2D residual deposits between them. Both the $CdSe$ shell thickness and Mg concentration induced an NIR emission related with strain or defect-related layer within the heterostructure NWs, as measured by cathodoluminescence.^{203,316}

Tang *et al.* reported the gradient alloy of $ZnS/Cd_xZn_{1-x}Se_yS_{1-y}$ core-shell QDs for liquid crystal displays based on reversible thermal quenching based on the multiphonon process, temporal trap states and re-existing trap states. $CdSe$ provides an effective confinement effect to prevent electron and hole recombination in large band gap materials such as ZnS . Secondly, the core-shell structure confines the wave function of electrons and holes for exciton recombination, which may reduce the generation of interfacial trap defects. Under illumination, the Auger recombination determines the non-radiative recombination at high power. Therefore, the luminescence quenching in QDs is related to the static charge distribution of trap states and surface trap states. Both resonance tunneling and thermally active physical processes

followed by radiative recombination with and without tunneling and non-radiative recombination are activated through tunneling.^{146,171,243,320} $ZnCdS$ and $ZnSSe$ QD alloys show blue emission with a quantum efficiency of 25–50%. In the case of core-shell nanostructures, ZnS can be used as both type-1 and type-II semiconductors with a core of different II-VI semiconductors.²¹ $ZnSSe$ NRs show spontaneous emission as excited by two-photon pumping with an 800 nm fs pulse laser. An increase in the Se ratio in a single crystal NR resulted in a red shift in its emission from 422 nm to 464 nm. The lasing emission line was achieved at 5.13 $mJ\ cm^{-2}$ with a sharp emission intensity (Fig. 12b). A further increase in excitation power resulted in a super-linear emission intensity in a single crystal, which further narrowed the FWHM due to the occurrence of a stimulated emission. A slight blue shift in the emission location corresponds to the phase space-filling effect of excitons at high excitation for low threshold materials such as $ZnSSe$ NR. A two-photon pump with fs laser is a beneficial tool to understand the lasing generated by free excitons and their scattering process in $ZnSSe$ single crystals. The lifetime provides suitable information on the lasing behavior originating from the free excitons or bound excitons in ternary alloys of $ZnSe$. The amplitude of the decay time of free excitons increases and the bound excitons amplitude decreases above a threshold value due to cavity effects, which saturate the binding states at a high carrier density and reduce the bound exciton amplitudes. The decay time also decreases with an increase in pump energy, which suggests a transition from spontaneous emission to stimulated emission with a narrow bandwidth and super-linear emissive intensity.



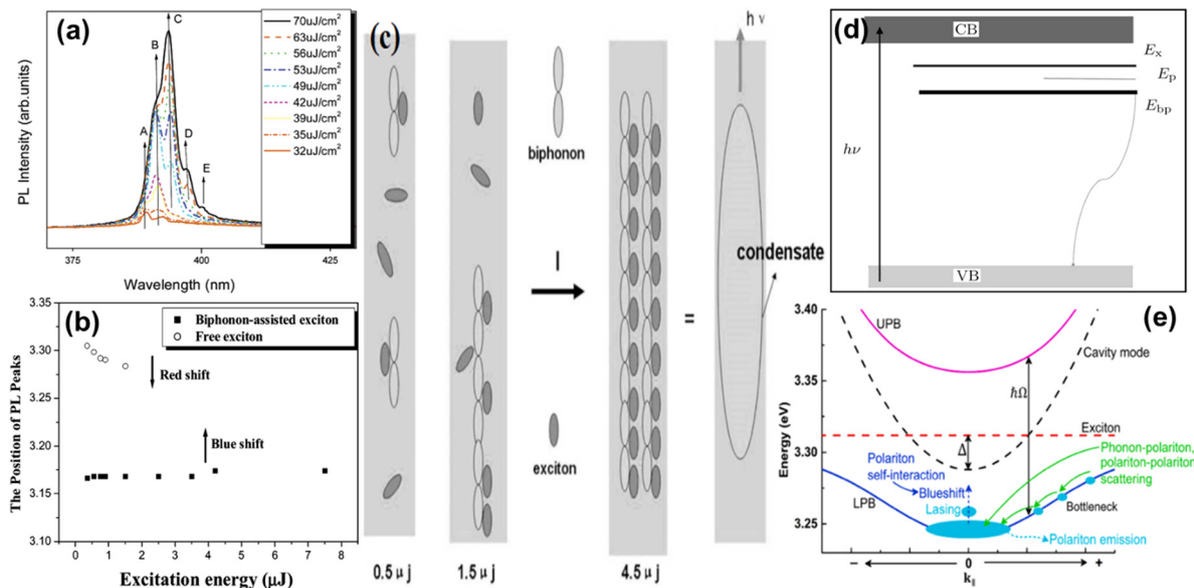


Fig. 12 (a) Detailed variation in lasing multi-modes with an increase in the pump fluence in ZnO nanowires. The direction of the arrows shows the increase in pump fluence; the mode marked by arrowhead B appears after A with an increase in pump fluence, and further, the modes marked by arrowheads C, D, and E appear. (b) Photoluminescence peak position from free excitons and 2LO phonon-assisted complex, respectively, showing their dependence on the excitation power, whose magnitude corresponds to the exciton density in the nanowires. Photoluminescence spectra at various pumping energies of ZnO nanowires, Copyright © 2006, ACS Publishing.³²² (c) Electronic states and optical processes of ZnO:Mn nanowires. (d) Dependence of peak positions of the photoluminescence from free excitons and 2LO phonon-assisted complex on the excitation energy corresponding to the carrier density in the nanowire. Copyright © 2011, IOP.³⁰⁷ (e) Schematic illustration of the polariton dispersion, polariton relaxation, emission and lasing process. The horizontal dashed line (in red) shows the uncoupled exciton energy and the curve dashed line (in black) shows the uncoupled cavity photon mode. The lines represent the upper polariton branch (UPB, in magenta) and lower polariton branch (LPB, in blue), respectively. $\hbar\Omega$ is the Rabi splitting energy. Δ is the cavity detuning. Copyright © 2020, Elsevier.³²³

The different modes illustrate the gain competition and distinct shifts as a function of excitation density.^{313,321}

Fig. 11d shows the power-dependent PL of ZnS and CdS as 6-fold symmetrical branched NS PL at room temperature. The strong luminescence behavior was observed with a red shift effect ranging from 502 nm to 520 nm, which was assigned to the band-edge dominant recombination of excitons or shallow trapped electron pairs of CdS. Moreover, the bandgap of ZnS and CdZnS is larger than that of CdS; thereby, the CdS-related emission is more prominent in the heterostructure prepared by CVD. The red shift in the spectral behavior is attributed to two factors, as follows: (i) excess carrier formation and (ii) trap state formation due to the presence of nonstoichiometric sites or defects of CdS at the interface. Therefore, a high excitation power is responsible for excess carrier generation due to the nonstoichiometric interface and enhances the carrier-carrier scattering, which may lead to electron-hole plasma formation at a threshold power, resulting in a redshift in the emission. At a low excitation power, the strong emission at 503 nm corresponds to the ZnS/CdS interface because the CdS:Zn layer at the interface may possess a slightly larger bandgap and larger radiative recombination efficiency than individual CdS. As the laser power increases, the amount of photoinduced excitons will increase, which move a longer distance from the interface to CdS ends with a redshift effect and broadening of the emission line. In the case of high power, another peak appeared far from the band-edge at around 673 nm to 800 nm, which was centered at 716 nm.³¹⁹

3.2.5.1 Transition metal ion-related luminescence behavior.

Fig. 11b and c show the luminescence behavior of a ZnSe/ZnS core/shell alloy NC doped with Mn and Cu. The Mn-doped ZnSeS alloyed NC phosphine-free approach resulted in the successful formation of MnSe nanoclusters together with ZnSe/ZnS core/shell (Fig. 11b; left). The formation of small-size nuclei of MnSe required a selenium source at 280 °C, resulting in the formation of MnSe clusters. At the interface of the MnSe core, the ZnSe shell is formed depending on growth temperature of about 260 °C. At 220 °C, the third layer of ZnSe was coated on the doped NC. The PL emission at 584 nm corresponds to the ${}^4T_1 \rightarrow {}^6A_1$ transition of the Mn^{2+} ion in ZnSe, as discussed earlier.^{317,324} The slight redshift in absorption indicates that the ZnSe layer grew after each injection of the Zn precursor. The high PL quantum yield was achieved because the Mn ion emission centers were isolated from the surface defects. Further, the ZnS shell grew to form a core/shell nanocrystal, showing a noticeable redshift without alloying between the core/shell with PL emission at 595 nm. The absorption spectra showed a small peak at 400 nm, which is attributed to the excitonic excitation of the host ZnSe/ZnS core/shell NC. Similarly, the alloy formation of ZnSeS showed a blue shift in its spectrum at around 350 nm (Fig. 11b; right). The Mn:ZnSeS alloy showed the Mn-related peak at 605 nm with an increase in the PL quantum yield, indicating a reduction in the interfacial scattering in the alloy shell.³¹⁷ Fig. 11c shows the Cu-doped ZnSe/ZnS core/shell NC and ZnSeS alloy PL quantum yield after 30 days, exhibiting the high stability of the green fluorescent material of Cu-doped NCs.



Therefore, the high quantum yield and stability of these NCs make them suitable for various optoelectronic applications such as bioimaging and LEDs. The emission of Cu:ZnSe d-dots was quenched after two days, while that of Cu:ZnSeS NCs decreased slightly but they still exhibited a higher quantum yield than Cu:ZnSe. In contrast, Cu:ZnSeS/ZnS core/shell d-dots exhibited better stability under air and light, maintaining ~90% of their initial intensity after 30 days. This is because ZnS as the shell protected them from the outer environment and stabilized the Cu emission of the NCs.³¹⁸ The D states belong to (i) surface defects, (ii) point defects and (iii) impurity defects, while the X state is related to the Co doped in the host matrix (Fig. 11e and f).²¹⁶ The 1D structure possessed a twin structure (wurtzite and zinc blende), which was formed during VLS growth, as confirmed by the structural analysis, and may be responsible for the D state or the above-mentioned defects.^{209,313} In the case of the doped nanostructure, the dopant ion is incorporated in a potential valley (green) near the intrinsic atomic potential.^{224,242,325,326} At room temperature or above room temperature, the potential barrier overlap disappears due to the mixing of the two potential wells, as shown in Fig. 11f. Another reason is the formation of a twinning structure of this state in a potential valley, which removes the energy barrier between the two states.

Similarly, the formation of an Mn-doped ZnSe/ZnS type-I core shell NW alloy through the solution process showed different emissions due to bandgap engineering.^{21,190,320} Mn doping in ZnSe is another strategy to tune the bandgap in 1D nanostructures and improve the overall luminescence of materials. ZnSe heterostructures formed with small bandgap materials such as CdSe cover a broad emission spectrum and can be employed in various optoelectronics applications. CdZnSe NWs have been prepared *via* the solution process and physical deposition method by numerous research groups.^{327–330} A change in the Zn content in the heterostructure causes a blue shift in the energy of the near band-edge and exhibits donor–acceptor pair recombination, corresponding to the presence of Zn vacancies near the valence band or deep acceptor states and shallow states near the conduction band. Moreover, the incorporation of Cd in heterostructures suppresses the zinc vacancies or Zn-defect related emission. Moreover, the adsorption of oxygen or water molecules at the surface of NWs will enhance their photosensitivity due to more surface electrons activating surface resistance. Therefore, the depletion layer on the surface of an n-type semiconductor will extract more electrons than low surface resistance. This capability of heterostructure devices improves the photo-conducting behavior together with switching application. Zn chalcogenide alloys with TM ion doping prepared by the CVD method and solution process can be useful for bandgap tuning by controlling their composition ratio, making Zn chalcogenides very efficient for various optoelectronic applications such as photodetectors, solar cells and LEDs.

4. Application of II–VI nanostructures

Binary and ternary semiconductors of Zn- and Cd-chalcogenides possess remarkable physical and chemical

properties and are popular due to their wide range of electronic and optical properties. Most importantly, materials belonging to II–VI semiconductors possess direct band gaps together with high optical absorption and emission coefficients. In addition, binary and ternary II–VI semiconductor alloys are readily miscible, providing a continuous range of properties. Therefore, they can serve as efficient light emitters, such as light diodes, lasers, solar cells, and radiation detectors operating in a wide range from IR to UV and X-ray region. Besides general photovoltaic applications, II–VI semiconductor binary and ternary alloys are also potential candidates for a variety of electronic, electro-optical, sensing, and piezoelectric devices; while few II–VI semiconductors cover the terahertz range. In particular, nanoparticles of II–VI semiconductors, such as QDs, 1D structures, and core–shell structures, can be used for the development of gas sensors, electrochemical sensors, and biosensors. As the device scales down to the few nanometer range, their tunable bandgap, high extinction coefficient, possible multiple exciton generation, and unique electronic and transport properties make them highly attractive in the research community. Therefore, the preparation methodology, doping concentration and crystalline quality of this class of II–VI semiconductors make them more important than III–V and other semiconductors. Currently, the blending of II–VI semiconductors with perovskites has been performed to form a new class of hybrid devices, achieving high sensing, opto-electronics and bio-medical device applications. In addition to the device-centric research, herein we also discuss the potential applications of II–VI semiconductor nanostructures in fields such as sensing, imaging, and quantum information processing. Thus, these applications can be further explored in future studies, investigating new sensing modalities, enhancing the sensitivity, or integrating these materials into advanced imaging systems with improved resolution and sensitivity. Moreover, our research underscores the significance of understanding the fundamental properties and phenomena in materials, such as carrier dynamics, spin-related effects, and optical properties in II–VI semiconductor nanostructures. Future studies can focus on unraveling these intricate mechanisms to pave the way for tailored material design and enhanced device functionalities.^{331–334}

4.1 Photonic and optoelectronic applications

1D nanostructures have diverse potential applications, especially in photonics and optoelectronics due to their excellent optical properties. 1D nanostructures of II–VI semiconductors have been employed in a wide range of photonic applications such as tunable lasers, leading to white light generation, photonic integrated circuits and many sensing applications. This section highlights some of the interesting instances of the application of binary and ternary alloys and TM ion-doped II–VI semiconductor matrix in detail.

4.1.1 Lasers. The first 1D NW laser was demonstrated by Yang and co-workers in 2001 based on the ZnO semiconductor.³³⁵ Afterward, several research groups demonstrated lasing phenomena in 1D ZnO NWs and NBs. Moreover, other II–VI 1D systems have also been extensively used for lasing application such as CdS, CdSe, ZnS and CdSSe.^{166,173,336,337} This is because 1D structures



not only exhibit spontaneous emission but also their reported stimulated emission results in optical amplification based on their cavity wall acting as a reflector. Binary and ternary II–VI materials and TM ion incorporation in 1D nanostructures are promising materials for use in next-generation laser application with the capability of tunable emission wavelength in integrated devices. Lasing in the ZnO nanostructure has been extensively studied due to its hexagonal wurtzite structure and formation of a cavity during high temperature growth, as reported by various groups.^{307,323,338–340} Gargas *et al.* demonstrated that a disk-shape architecture performed low threshold voltage whispering gallery mode (WGM) lasing with a small loss in comparison with the FP cavity in hexagonal wurtzite ZnO for on-chip nanophotonic integration. Both experimental and theoretical correlation showed that UV WGM lasing can be achieved using ZnO nanostructures with subwavelength diameters.^{300,301} The room temperature emission of single-mode lasing was observed from nano-disks having a diameter of 842 nm upon excitation by an optically pump pulsed laser. At a low laser pump power, the emission spectrum was broad and wide, but as the pump power increased above the threshold value, the emission spectrum became sharp with the maximum FMHW value of about 0.8 nm. The integrated peak intensity at 391 nm was dependent on the pump power or in other words threshold value dependent. According to the experimental result, nanodisks with different diameters exhibited a lasing emission at around 385 nm to 391 nm, which was firmly due to the overlap of WGM with the gain media of ZnO.^{335,340–342} Moreover, the lasing threshold was tuned as a function of diameter, where a large diameter exhibited a low lasing threshold owing to the overlap of the WGM mode to shift the spectral window of optimal gain. Similarly, a smaller diameter compromised the spatial overlap between the WG mode and the ZnO gain medium. The optical mode diminished confinement caused a reduction in the quality factor owing to the increase in the leakage power. The simulation study revealed the 7.5% reduction in radiative power lost through the substrate can be improved by 20.7% with a taper base increase in the *Q*-factor. Both experimental and simulation based on FDTD show promising outcome for nanoscale laser and on-chip device for nanophotonics.^{300,312}

Fig. 12a shows that heavily Mn-doped ZnO shows different modes in its luminescence spectrum under different excitation powers. With an increase in the pump power, the carrier interaction became stronger with a redshift due to the amplified spontaneous emission. A high excitation power may lead to electron–hole plasma formation, which increases the carrier interaction in a specific direction to support the lasing mode simultaneously. In comparison with the low doping profile (Fig. 10), the free exciton emission is suppressed by the strong scattering of the impurity ions and phonons. At a low pump excitation, the stimulated emission is due to the exciton–exciton interaction. A further increase in power resulted in an increase in the emission intensity, reflecting the giant oscillator strength and cooperative radiation of localized excitons. At a high power excitation, electron–hole plasma formation becomes dominant, which results in a redshift in emission with multimode lasing

for a high Mn concentration. Fig. 12b shows the low Mn doping profile in ZnO at different fs laser excitation at room temperature, as discussed in Fig. 10 (peaks A and B). The blue shift in the emission is due to the phase space filling effect without carrier scattering of excitons at high power. At 4.2 μJ , the coherent transition of the bipolaronic state enhances the intensity and narrows the FWHM of the emission spectrum. With a further increase in pump power (7.5 μJ), the peak energy width became 10.5 meV (second peak appears in the emission spectrum), which is assigned to the dipole–dipole interaction of different coherent states. According to the experimental results, the emission behavior is similar for the low and high doping profiles at a high excitation with a change in the threshold power, which is dependent on the doping profile. Secondly, the redshift in peak A is attributed to the aggregation of neighboring excitons or screening to enhance the Coulomb interaction between free carriers through bandgap renormalization.

Liu *et al.* revealed that the lasing behavior in Mn-doped ZnO is due to the condensation of magnetic polaronic excitons in 1D NWs prepared by the VLS mechanism.³⁰⁷ To understand the condensation of polaronic excitons, it is necessary to understand all the possible transition states in the host material. At a low excitation power, the exciton and exciton–phonon related peak becomes prominent in the spectrum, whereas at high excitation power, the exciton–phonon interaction is dominant and generates polaronic excitons (bound exciton). The lasing phenomenon in Mn-doped ZnO is owing to the bound excitons, similar to the emission of Bose–Einstein condensation (BEC) excitons, as shown in Fig. 12c. Fig. 12d shows the change in the electronic properties of ZnO with the incorporation of dopant ions. A low doping profile results in strong electron–phonon interactions due to the 2LO of the A1 mode coupled with excitons. FM coupling with coherent excitons result in magnetic polaronic exciton interaction, which condensates by a fs laser to realize the BEC in real confined space. Another key factor related to lasing is Mn doping in the host crystal. A high doping concentration has a higher lasing threshold value than the low doping samples. Therefore, free excitons, coupled spins and bi-phonons lead to the formation of exciton magnetic polarons and condensation-related emissions, which are a function of the doping concentration and pump power for ZnO:Mn lasing; thereby, the intra- and inter-band transitions need to be understood to determine the shift in the emission spectrum under different conditions. The phonon–exciton complex has a higher binding energy than free excitons and two-phonon mode, and therefore both the free exciton- and bi-phonon-related emissions appear at a low excitation pump power owing to their high thermal stability at room temperature. BEC exists in ZnO:Mn-doped nanostructures due to the following reasons: (i) free excitons transform efficiently into bi-polaronic excitons, (ii) NW side reflection may confine the high excitons, which reduces the diffusion or expansion effect and (iii) the aggregation of free excitons with the coherent 2LO A1 mode at a high density. In the case of 1D nanostructures, their crystal orientation is also an important factor that can lead to a reduction in the exciton energy in their interaction and limit the bi-phonon-related emission. In the parallel orientation, the



dipole repulsion increases the exciton energy, which exhibits a redshift at a low excitation pump power. Meanwhile, coupled excitons aggregate and the related repulsion increases in the parallel orientation, which will enhance the bipolaronic excitons at high density and characteristic of dynamic BEC emitting light coherently without other available states.⁵

Optical characterization tools such as PL provide suitable exciton information under different powers. In the case of a low Mn concentration, the free excitons are depleted by the coupling of LO phonons and ion spins at high power, leading to the formation of EMP with a large exciton binding energy and strong electron–phonon coupling semiconductors such as ZnO. Strong exciton–phonon coupling materials normally exhibit an increase in LO mode phonons with the addition of impurity ions. Thus, the LO phonon mode coupled with bound excitons results in the polariton–exciton emission. Mn:ZnO exhibits the EMP corresponding to the 2LO–exciton state at a low excitation energy due to multiphonon exciton but the 3LO and LO coupled exciton emission band is not coupled with free excitons. EMP attain higher thermal stability due to their higher exciton binding energy than thermal energy (26 meV). High exciton binding energy semiconductors possess higher stability of exciton–phonon coupling and exciton–spin interaction at room temperature. A high doping concentration enhances the exciton–phonon coupling and increases the trapped states together with the bandgap. Under these conditions, the emission spectra reflect the forbidden selection rule for one-phonon transition and allow a two-phonon transition or facilitate the bi-polaron process, leading to non-linear interaction in 1D structures. Therefore, the bosonic phonon and exciton dominant the coherent excitonic state of the bound exciton in polar semiconductors, dominating the EMP emission. The EMP peak shows a rapid enhancement, narrowing FWHM and slight blue shift depending on the power due to the exciton–exciton interaction and coherent two-phonon–exciton coupled with spin. Bi-polaron exciton transformation in 1D nanostructures is produced by temporal, coherent and high-density excitons upon excitation by fs laser. High-density excitons in 1D nanostructures are dependent on their crystal face reflection or aggregation of free-excitons, exhibiting coherency in a specific orientation or crystal direction. In the case of the wurtzite structure, the polarization of light–matter interaction shows the orientation-dependent exciton emission due to a change in the dipole force along the parallel or perpendicular direction. In spin-dependent semiconductors, the FM spin exchange coupling in a magnetic field changes the dipole force and aggregation of excitons or spin ions responsible for EMP formation in 1D nanostructures. Therefore, bi-polaronic excitons aggregated in a high density emit coherent light, which resembles a characteristic of the dynamic exciton BEC state (similar to Cooper pairs in a superconductor) in a semiconductor 1D NW structure. Recently, Zou *et al.* explained the identification of coupled spins in 1D NWs and its origin. According to the magnetic measurement, the origin of the FM coupled with spin inside the host lattice was doubtful. However, the ns time-delay PL spectra show PL relaxation due to the local d–d transition. The strong electron–phonon coupling and spin ion in the host lattice indicate FM

coupling in the polariton–exciton emission with the energy position, indicating the origin of the ferromagnetism in DMS. Moreover, the NW cavity demonstrates exciton decay into a very narrow mode of coupling, which is strong enough to produce polariton lasing and polariton condensation in 1D nanostructures. This long-range elementary excitation has spin-polarized character and can give collective EMP condensate at a high density. In QDs, EMP can be easily formed, but without long-range LO phonon coupling, as observed by the difference in PL behaviors. PL can be employed to identify the spin ion FM coupled aggregate d–d transition in 1D structures with a slight redshift with its FM spin cluster. This type of cluster in 1D nano/micrometer sized structures favors EMP formation and enables strong single-mode lasing from the BEC-like state of collective EMPs in the DMS structure.⁵ Zhao *et al.* achieved low threshold lasing in a thin 1D pure ZnO NW WGM at room temperature through surface plasmon promoted by single-mode polaritons.³²³ Fig. 12e shows a schematic diagram of polariton dispersion for lasing, corresponding to the relaxation and emission of coherent plasmon–polariton in cavity-mode ZnO. The horizontal dashed line (in red) shows the uncoupled energy (exciton energy) and the curve dashed line (in black) shows the uncoupled cavity photon mode. The upper polariton branch (UPB, in magenta) and lower polariton branch (LPB, in blue) show the Rabi splitting energy ($\hbar\Omega$) and cavity detuning (Δ), respectively. A nano-laser of ZnO NW on graphene–insulator–metal platform exhibited gate-modulated lasing based on the sensitivity of surface plasmon polaritons to the cavity effect. The electric field distribution on integrated circuits on the nanoscale has potential application as a function of gate voltage. Under an applied voltage, the electric field is shifted away from the metal surface, which results in the internal loss of lasing. Therefore, a high-applied voltage modulates the low lasing threshold and vice versa. The PL-related peak intensities were recorded in a repeatability test over five periods, where VG was set as 6V and 0 V in the on-state and off-state, respectively, and the switching period was 1 s. In each positive cycle of 6 V, the on-state accumulated surface electrons on the metal surface, which may increase the plasma frequency at the metal side, resulting in the internal loss of the surface plasmon polariton mode at a fixed pumping power. Therefore, the lasing signal could be dynamically modulated by gate switching.³⁴³

4.1.2 Solar cells. Solar cells convert light into electricity through the excitation of e–h pairs in semiconductors. Currently, silicon-based solar cells are the main type, having an efficiency of around 25% have expensive manufacturing cost. Therefore, other materials such as II–VI semiconductors have been introduced due to their high efficiency, cost effectiveness and low manufacturing cost. Specifically, 1D II–VI nanostructures are highly crystalline, possess a large surface to volume ratio and photo-active in a wide spectrum range (from UV to IR regions), which are considered next-generation photovoltaic devices.^{16,31,39,344} Law and co-workers reported the fabrication of organic, organic and inorganic hybrid and dye-sensitized cells for solar energy conversion based on ZnO NWs, ZnO particles and TiO₂ particles.³⁴⁵ Fig. 13 shows the scanning electron microscopy cross-section image of a vertically aligned



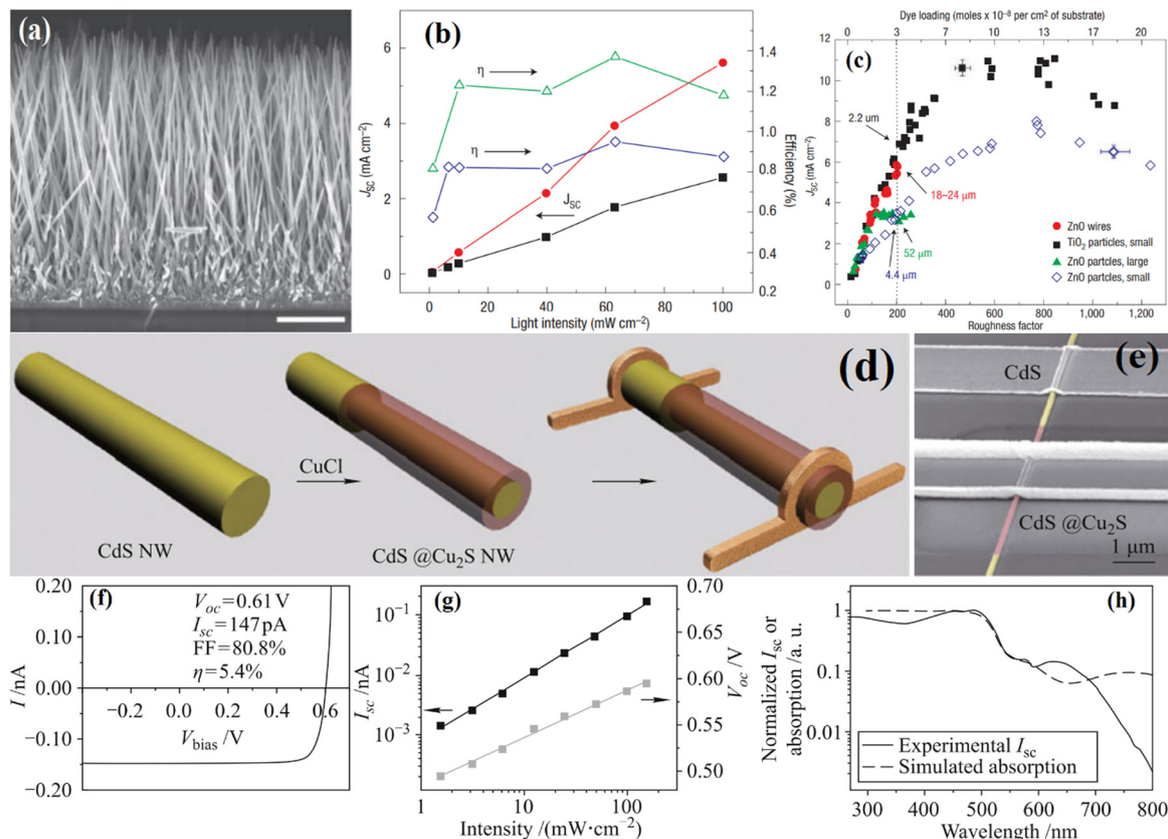


Fig. 13 (a) Scanning electron microscopy cross-section image of ZnO nanowire array on FTO for dye-sensitized cell. The wires are in direct contact with the substrate, with no intervening particle layer. (b) Short-circuit current density and efficiency as a function of light intensity for cells with roughness factors ranging from 75 to 200. Each of the four parameters is represented by data from two different devices to provide an estimate of the range of their variability. Active cell size: 0.8 cm^2 . (c) Short-circuit current density versus roughness factor for cells based on ZnO wires, small TiO_2 particles, and large and small ZnO particles, Copyright © 2005, Nature.³⁴⁵ (d) Schematic diagram of core-shell CdS/ Cu_2S single NW. (e) SEM image of a single NW. (f) I - V curve of a single-NW under 1 sun (AM 1.5 G) irradiation. (g) Variation in J_{sc} and V_{oc} with light intensity. (h) Wavelength dependence of photocurrent compared with simulated absorption, Copyright © 2011, Nature.⁵⁵

ZnO NW array on FTO, which was used for the fabrication of a dye-sensitized cell. Moreover, the NWs were in direct contact with the substrate and no other intervening particle layer was present. The performance of the ZnO NW array is compared with that of NP in Fig. 13b, where the plot shows the light intensity as a function of short-circuit current density and efficiency of the two solar cells. The array-like structure provides direct charge extraction but more traps directly affect the transport of carriers, as discussed above. Also, the array-like geometry allows the formation of a full depletion layer at the semiconductor-electrode interface, leading to electric field-assisted charge separation, which provides better charge transport in comparison with NPs. Another factor that directly affects the solar efficiency is the surface roughness. In this case, the NW array of ZnO has a lower surface roughness in comparison with the NP film. Therefore, the lower surface roughness, higher trap occupation density and lower light absorption of the vertically aligned NWs than the film-like structure result in a greater efficiency, as shown in Fig. 13c.^{302,345}

Tang *et al.* reported that a single CdS NW showed a better response than planar devices.⁵⁵ The high-crystalline single CdS

radial NW p-n junction exhibited an open circuit voltage of 0.6 V and fill factor of 80%, as shown in Fig. 13f. The CdS/ Cu_2S core-shell NWs showed higher values than the planar devices at a low open circuit voltage with a reduced light intensity (Fig. 13d and e). The interfacial junction between core-shell materials should be high quality and the VLS mechanism should provide a high-crystalline interface between CdS/ Cu_2S , with the calculated overall efficiency of 5.6% with a lower current density (Fig. 13g and h). Moreover, the solar cell efficiency is also dependent on the thickness of the shell materials, where an increase in the shell thickness leads to absorption within the interface, affecting the efficiencies of radial solar cells. Additionally, horizontal devices can facilitate the study of the uniformity of charge collection as a function of orientation and single NW. For this purpose, the scanning photocurrent mapping (SPCM) of horizontal NW demonstrated that the core-shell region is more active for charge separation, while the edge exhibits a sharp drop-off. The radial p-n junction has benefits of charge collection from CdS, which suffers from a low minority diffusion length. Moreover, multiple devices with parallel and series configurations based on core-shell single NW



provide a flexible absorption range at low cost and better photonic device performance than planar devices. CdS/CdO core shell and other ternary II–VI nanostructures are potential materials for future photonic applications.^{75,174}

4.1.3 Photodetectors. 1D nanostructures have attracted significant attention due to their unique optical and electrical behavior on the nanoscale. Also, their high surface to volume ratio and improved crystal quality provide additional benefits to enhance the charge carrier transportation in photoelectric devices. Thus, 1D nanostructures based on CdS, ZnO, CdSe, ZnCdSe and other II–VI semiconductors have been employed for the fabrication of tunable photodetectors.^{201,346–348} $Zn_xCd_{1-x}Se$ alloyed NWs were synthesized *via* the CVD method for variable wavelength detection. Yoon and co-workers successfully showed their tunable emission by adjusting their composition gradient, as discussed above, but they first claimed the successful representation of wavelength-controllable optoelectronic devices of II–VI ternary alloyed NWs.³³⁰ The slow response shown by CdSe is owing to the adsorption of ambient gas molecules on its surface, similar to other n-type semiconductors (such as ZnO). Thus, the capture of oxygen ions at the surface generates a depletion layer on the surface or increases the resistance, which is responsible for its slower response time than ZnSe and $Zn_{0.72}Cd_{0.28}Se$, respectively. The two other photodetectors possessed a less active surface in the ambient environment and practically applicable as variable wavelength photodetectors covering a wide range of the visible spectrum from blue to red.³³⁰ Fig. 14 shows the CdSSe chip sensor that can respond to moisture, and its humidity dependent *I*–*V* curves in the dark and light under bias condition are shown in Fig. 14a and b, respectively. At 5 V (dark), the current increased from 3.36×10^{-7} to 1.15×10^{-6} A with an increase in the relative humidity from 25% to 80%. Under illumination, the photocurrent changed from 7.48×10^{-6} to 1.38×10^{-5} A. The inset shows the *I**V* characteristic response under different moisture conditions. The polyimide (PI) on the surface tightens the contact between the CdSSe NW layers and under the appropriate bias voltage gives a good response in the light and dark (Fig. 14c and e). Fig. 14d shows the resistance of the sensor for different humidity levels, showing a linear dependence for both dark and light, which suggests the feasibility and convenience to predict the humidity of moisture. The relative humidity of moisture-dependent resistance in the dark field can be well fitted with a linear function ($R_2 = 0.9779$), and also with illumination ($R_2 = 0.9948$).^{188,189}

Recently, many groups have devoted their efforts to fabricating flexible photodetectors based on 1D and 2D structures. In the case of II–VI 1D nanostructures, there are many reports on both ZnO and CdS related to their photo-detection properties, which depend on the following: (i) composition gradient, (ii) incorporation of dopant and (iii) synthetic mechanism. 1D CdS NW and WSe_2 2D flakes were used as the substrate for the fabrication of a photodetector by Lin and co-workers.³⁴⁷ Both Lin and Du highlighted the importance of van der Waals heterostructure growth.^{347,349} The Xiong group reported the growth of various II–VI semiconductor nanostructures,

reporting van der Waals epitaxial growth by the VLS mechanism extensively.¹⁰ Their photo-response as a function of time and illumination intensity were investigated. With an increase in the illumination intensity, the photo-response of the device decreased due to the interface trap states. The *I*–*t* characteristic curve showed the stability of the photodetector under different illumination intensities at a fixed bias condition of $V_D = +2$ V. Also, it was shown that its response time is less than 1 s and its switching behavior was retained after multiple cycle tests under strain. Similarly, the *I*–*V* characteristic curves under light and without light for the flexible photodetector under different strains were recorded together with the effect of different strain on the output characteristic curve of the photodetector. According to the results, the current of the CdS NW increased under illumination but strain had no impact on the performance of the device. Lou *et al.* also fabricated a 1D ZnO–SnO₂ photodetector for UV photodetection on a PI substrate.³⁵⁰ The ZnO–SnO₂ devices showed excellent UV photodetection under ambient condition. The conductivity of both materials improved due to the oxygen on their surface enhancing the carrier response, which may improve the high I_{ON} – I_{OFF} ratio, fast response time and good reversibility.³⁵⁰ Moreover, Yang *et al.* reported that a ZnO NW covers a broad range of the spectrum for detection under different laser excitation and exhibits ultra-high sensitivity, fast response speed and good flexibility for future optoelectronic circuits, switches, *etc.*^{60,301,351} Guo *et al.* synthesized highly crystalline tin-doped CdS NW for tunable optical loss and multi-band photodetection using the CVD method.^{96,173,188} CdS NWs showed pump power-dependent propagation loss of the near band edge emission and defect-related emission. According to the low temperature PL measurement, the activation energy was calculated to be 34.29 meV, illustrating carrier decay owing to exciton recombination. The high photo-response (51.2 A W^{-1}), high gain (313.3) and stable optical switching at $V_{DS} = 3$ V revealed the importance of 1D II–VI semiconductors in the design of integrated optical interconnection chips.⁹⁶ Lieber *et al.* employed CdS NW on silicon to produce a nanoscale injection laser to cover a wide range of the spectrum, and also integrated it as a single or multiple color laser in silicon microelectronics and single wire on-chip devices.^{40,352–354} CdS single NWs have a high injection rate due to cavity formation and some issues can also be addressed by axial composition, core–shell heterostructure, composition gradient and TM ion doping or metal doping on the surface of 1D nanostructures. Moreover, the radial p–n junction of n-CdS and p-Si NW for an avalanche photodiode demonstrated ultrahigh sensitivity with detection limits of less than 100 photons for ultradense integrated systems, sensing and imaging applications.³⁵³

4.1.4 Waveguides. II–VI 1D nanostructures can be functionalized to confine and propagate electromagnetic waves to specific directions. II–VI semiconductors cover a wide range of the spectrum and possess larger dielectric constants and refractive index than their surroundings, allowing the transmission of light with low loss through total internal reflection. These inherent properties make II–VI 1D nanostructures suitable for various optical applications on the nanoscale. Moreover, ternary



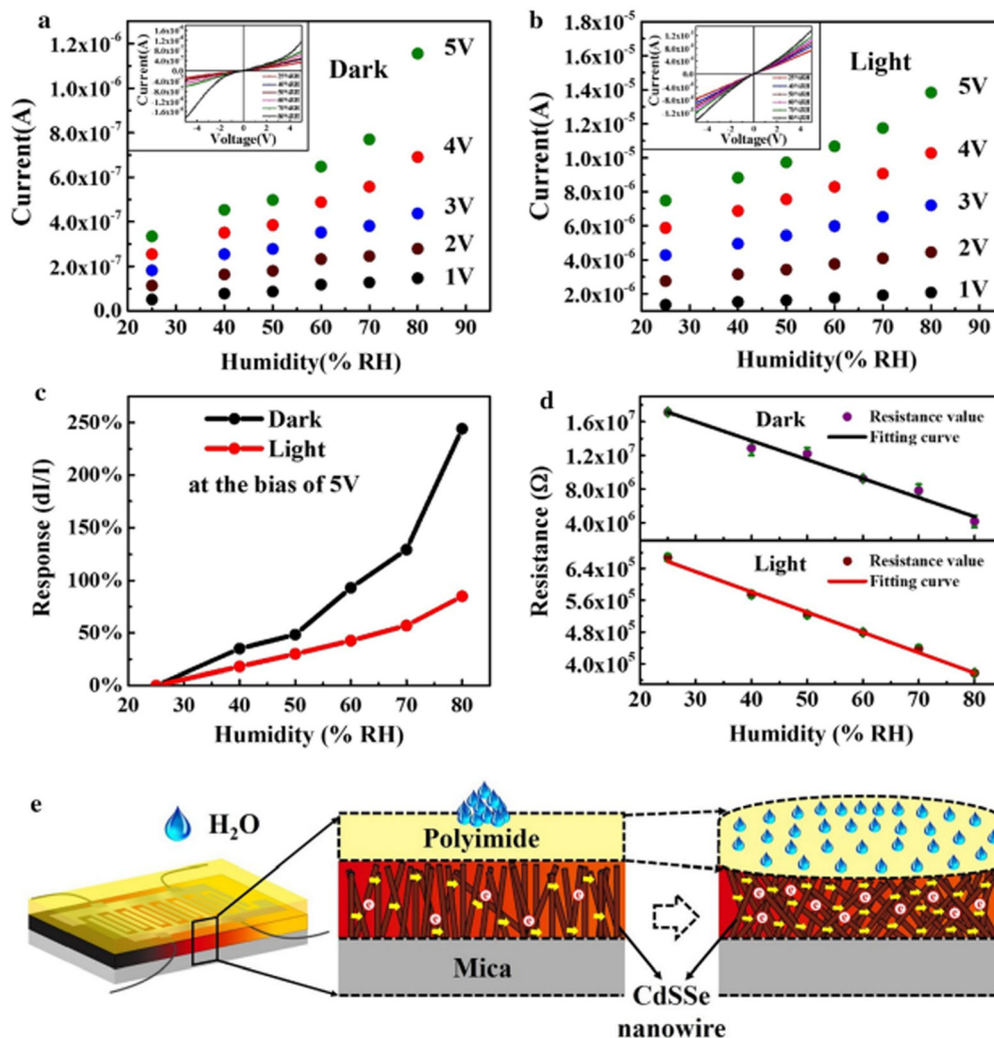


Fig. 14 (a) Dark current and (b) photo-current at various humidity levels under different bias and the inset shows the IV characteristics in the dark (a) and light (b) at different relative humidity. (c) Relative humidity-dependent moisture responsivity of the sensor in dark and bright fields, respectively. (d) Resistance at various relative humidity of the sensor in dark and bright fields, respectively. (e) Schematic theoretical diagram of humidity sensing by the sensor.¹⁸⁸

alloys of 1D II–VI semiconductors enable bandgap engineering and metal ion doping with large dielectric constants and higher refractive index facilitate the propagation of light in one direction. Several groups demonstrated the waveguide behavior of II–VI 1D binary, ternary and metal dopants in II–VI nanostructures in the last decade. Pan *et al.* showed that single crystalline NB of CdS prepared by the CVD method exhibits waveguide behavior.^{77,175,355} Johnson *et al.* and others described the significance of ZnO NWs in wave guiding behavior, while simultaneously behaving as optical cavities (such as FP cavity), and presence of flat and parallel end faces act as reflectors.^{335,341,356} Lieber and co-workers presented the future prospective of CdS NWs, where a single wire can act as an active photonic circuit element for guiding light at sub-wavelength scales.^{352,354,357} Active wave guiding in 1D NWs operated near the band edge was demonstrated by the Zou group for CdS nanoribbons and CdSe NWs.^{89,175,355,358} TM ion doping or metal doping (such as tin) enhanced the cavity emission near the IR region depending

on the metal ion.^{89,96,359} Fig. 15a shows the waveguide demonstration of CdS_xSe_{1-x} nanoribbons and Fig. 15b shows the emission from a branched NW. Fig. 15c shows the PL emission spectra recorded from different marked points in Fig. 15b. The light is emitted in the wide spectrum range in one branched NW; one end is excited with the band edge emission and light propagates along it; while other end of the NW acts as a waveguide. Moreover, the emission is only seen at the extreme end of the branch NW, which indicates the guided nature of the light, as shown in Fig. 15a, where the magnitude of the redshift in the spectra is strongly related to (i) the composition gradient and (ii) degree of crystallization of the nanostructures on the nanoscale. Fig. 15c shows that the different emissions in the branched CdS_xSe_{1-x} nanoribbons are related to the above-mentioned points. The understanding of waveguides in 1D NWs/ribbons offers the opportunity of using them in highly integrated and compact photonic circuits with improved speed limitation and power dissipation in current electronics.



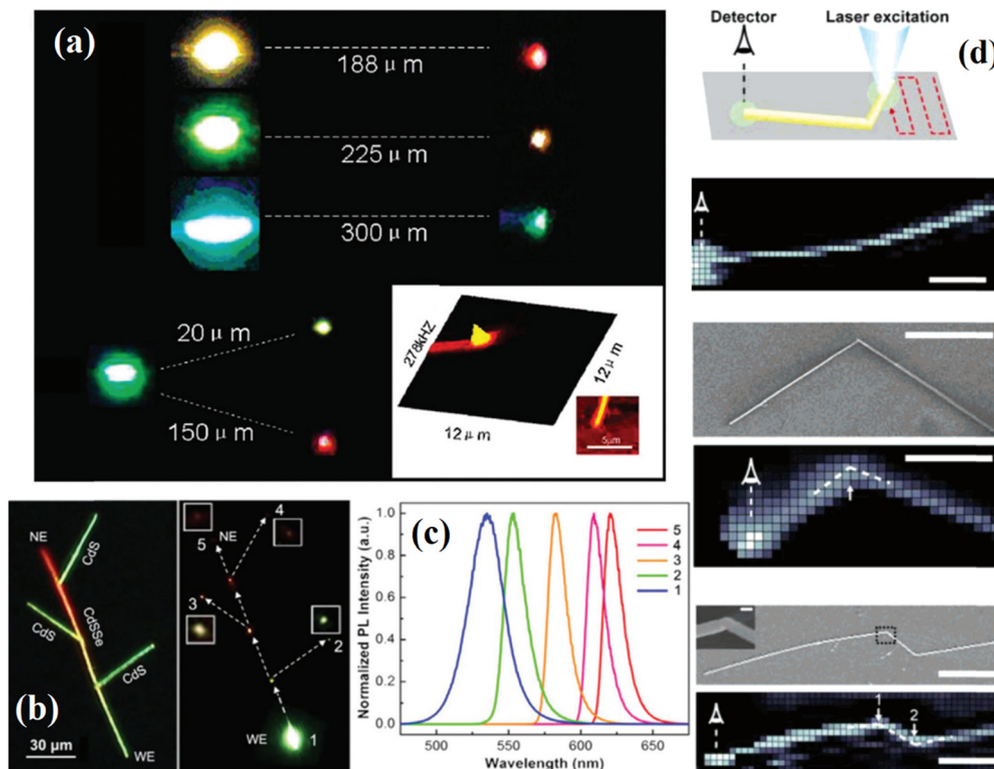


Fig. 15 (a) Waveguide demonstration of $\text{CdS}_x\text{Se}_{1-x}$ nanoribbons. Copyright © 2007, the American Chemical Society. (b) Selective waveguide from the branched architecture of the $\text{CdS}_x\text{Se}_{1-x}$ nanostructures and (c) PL spectra of the marked locations in (b), Copyright © 2012, the American Chemical Society.⁷⁸ (d) Characterization of optical wave guiding in straight and bent nanowires. (top) Schematic diagram. (2nd) Scanning optical microscopy (SOM) image illustrating the focused laser spot scanned over the sample, while monitoring light emission from one end of the nanowire. (3rd) SEM image of a single CdS nanowire and (4th) corresponding SOM image. (5th) SEM image of a single kink CdS nanowire and (last) corresponding SOM image at different positions (marked 1 and 2), Copyright © 2004, the American Chemical Society.³⁵²

Therefore, the manipulation of light from the UV to IR range is beneficial to develop favorable single NW integrated photonic circuits of binary, ternary and metal ion-doped II–VI materials.⁷⁸

Fig. 15d shows the optical waveguide in straight, bent or kink CdS NWs reported by the Lieber group using scanning optical microscopy.^{352,357} The light propagation through a single structure only shows moderate loss through a sharp and acute angle bending due to the (i) surface roughness or (ii) change in refractive index at an interfacing angle.

Due to its wide bandgap with cylinder geometry or hexagonal structure with large dielectric constant, Sn doping in CdS show good waveguide behavior, which was activated as a superlattice 1D nano/micro-wire, as shown in Fig. 16a. The inset shows an optical micrograph of the Sn-doped CdS. Its optical behavior was similar to a 1D photonic crystal (PC). Fig. 16b shows a schematic diagram of the CdS/CdS:SnS₂ junction within the single wire-like structure. The CdS segment in the superlattice wire confines the transport of photons, while due to the large refractive index of SnS₂, it acts as a reflector to trap light within its optical micro-cavities. Periodic bright emission arises from the interference of the coherently scattered light wave within the coupled cavities not related with the Fabry–Perot mode (Fig. 16c–h).¹⁹⁴ The different emission modes with a variation in wavelength can be imaged with this 1D PC, behaving like a local photon bunch. Each mode comes from

the neighboring two cavities and can propagate at a very long distance. The macroscopic mode number can follow the Bloch oscillation character in a broad emission range, indicating a strong nonlinear photonic structure for exciton–photon interactions. Moreover, the incorporation of Sn(IV) in the CdS lattice results in a bandgap edge and trap state emission due to the nonlinear electron–phonon coupling or anharmonicity effect. The exciton trapped by local impurity states is related to paired states or the deep trapped states involve different recombination rates of relaxation, as shown in Fig. 16i.^{194,196,359} Similar behavior of color-tunable period spatial emission was observed in ZnO/FeZn₂O₄ and CdS_{1-x}Se_x/Sn:CdS_{1-x}Se_x superlattice by varying the composition gradient from 0 to 0.4.^{96,196,360–362} The light transports along the axis and is periodically emitted at the adjacent joint segment. A change in the refractive index of CdSSe and Sn forms the neighboring alloy segment, which acts as a microcavity through the reflection interface, where light partially leaks through the joints. The periodic bright emissions arise from the interference of coherently oscillated light within the coupled microcavities, producing multi-peaks at the low energy side. The strong absorption above the bandgap cannot transport far within the wire, and thus it cannot produce waveguide modes. Fig. 16i shows that PL spectrum exhibits a strong band edge emission at ~551 nm and several accompanying shoulders related to the coupled cavity modes.



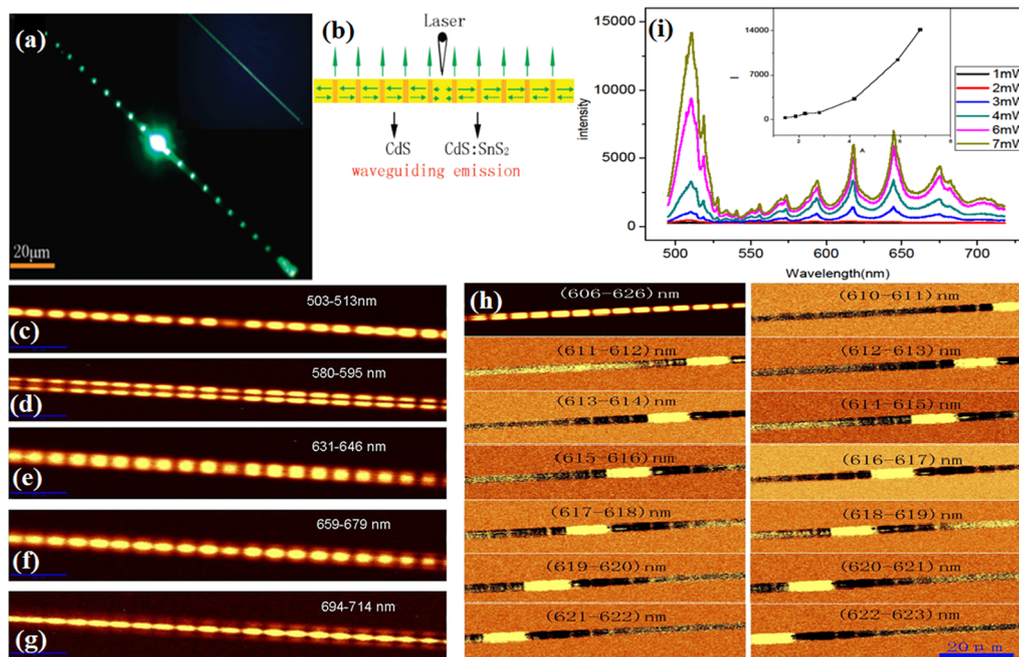


Fig. 16 Far-field PL images of an excited sample with different reaction times: (a) 20 min and (b) 30 min. Insets in (a) and (b) are the corresponding optical images. (c) Schematic representation of the emission process in the 1D superlattice wire, Copyright © 2010, the American Chemical Society.¹⁹⁴ (d) Schematic diagram of the branched nanostructure of 355 nm incident laser (inset is the optical microscope image). (e)–(g) Real-color dark-field photographs under the illumination of a 355 nm laser with an increase in pumping power density, with a scale bar of 20 μm , Copyright © 2012, Wiley-VCH. (h) The photoluminescence mapping of CdS:Sn microwire shows photonic modulation in multi-peak range.¹⁹⁴

The mapping profile in the wavelength range of 606–626 nm is shown in Fig. 16h. The photonic modulation and possible exciton coherent emission are combined in different optical microcavities. Thus, we can conclude that these emission contours were produced by the combined contributions of the coupled optical microcavity modes and local excitons within this 1D photonic crystal wire.³⁶²

Branched-like nanostructures have been extensively used for various applications, as suggested by the Lieber and Yang group. Liu *et al.* also fabricated a nanoscale laser array of branched-like structures of CdS doped with Sn, as shown in Fig. 17a–d. Sn acts as a junction, which splits into a multichannel nanostructure, resulting in effective light confinement and oscillation. Under excitation, a strong band-edge emission was observed and Sn reflected the light scattered from the branch segment. Thus, the oscillation of light within the branch and trunks shows the multi-point lasing high-quality factor of up to 990 with a low threshold. The waveguide performance of the hierarchical trunk-branch CdS NW array excited by UV light showed bright green emission that propagates along the branch and emits from the tips, as shown in Fig. 17b. As the central part is excited, the light propagating from the tips of the branches is stronger than the trunk-branch (Fig. 17b) due to the propagation distance, as shown in Fig. 17c. As can be seen in Fig. 17b and c, the branch-like structure shows a waveguide nature and its propagating distance shows marginal damping in emission. The excited trunk shows efficient light coupled between the trunk and branches in a monolithic structure, as shown in Fig. 17d. This exceptional optical coupling motivates optical integrated device applications

such as active waveguide trunk–branch cavities for 1D NWs to hundreds of branch wires.^{89,96,100,359,363}

4.1.5 LEDs. II–VI semiconductor 1D nanostructures show good electric conductivity and light emission due to carrier injection through radiation recombination. However, the homoepitaxial growth of the same II–VI materials results in a poor performance, as reported in ZnO. Although the lattice mismatching-related defect can be overcome in homostructure growth, the overall performance is compromised. Accordingly, most of the research available on the fabrication of LEDs using II–VI materials focused on their growth mechanism and dimensional growth.³⁶⁴ Among the different types of 1D growth, radial heterojunctions and homojunctions show a much better device performance. The major research on LEDs has been reported for QD-, NC- and NP-based binary, ternary and metal ion-doped heterostructures. QDs based on type-I core-shell ZnSe/ZnS exhibit a high quantum yield in the range of 45% to 65%.^{365,366} A type-I ZnSe/ZnS NR heterojunction showed a 50% quantum yield and emission wavelength in the range of 421–485 nm.³⁶⁷ In the case of a type-II CdSe/Te NW core-shell-like structure, it exhibited a quantum yield of about 20–60% and cover a wide range of the spectrum of 530–760 nm.³⁶⁸ Similarly, the type-II double-shell structure of ZnSe/CdS/ZnS exhibited a quantum efficiency of 43–61% in the broad-spectrum range of 470–760 nm.³⁶⁹ The 1D growth of NW and NB-like structures can form a junction with other nanostructures or they can be grown on a substrate for displays and electroluminescence. ZnO and CdO normally exhibit n-type conductivity, whereas CdTe and ZnTe are inherently p-type conductivity semiconductors. Therefore, the junction for electroluminescence study must be among the II–VI n-type conductivity



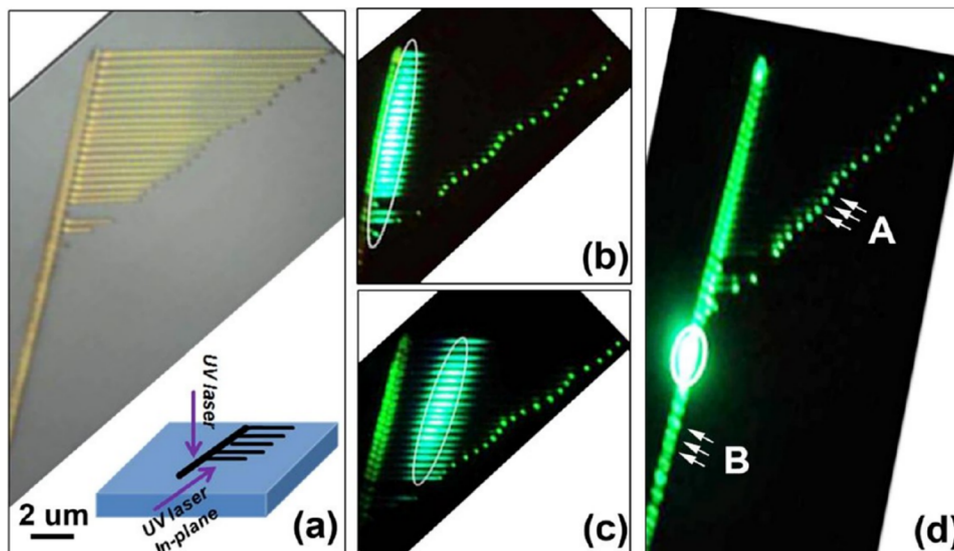


Fig. 17 (a) Optical microscopy image of the yellowish CdS comb structure on an Si wafer. The inset schematically shows the different illumination directions and areas of the UV laser (GaN laser diode at 405 nm wavelength). (b) Green photoluminescence with lateral (in-plane) illumination collimated at the trunk-branch junctions, (c) at the central part of the branches, and (d) vertically collimated as indicated by the white ellipses. Note that the arrows marked by B indicate the periodic emission of light resulting from the Fabry-Perot cavity for lasing; the arrows marked by A indicate the light emission from the end-tip of the branches, Copyright © 2013, ACS Publications.⁸⁹

materials with p-type or p-type III-V semiconductors such as GaN.^{370,371} Thus, electroluminescence in II-VI materials has attracted significant attention, which can be easily integrated with electronic devices for a wide range of LEDs, display panels, *etc.* 1D II-VI nanostructures have advantages over other synthesized growth dimensions, including their direct bandgap, which can be tuned due to their large bandgap coverage, tunable bandgap, large aspect ratio, small size, and high crystalline quality. Their tunable bandgap emission facilitates the coverage of a broad-spectrum range from the UV to IR regions. Also, their large aspect ratio can facilitate their device integration and their small size can generate a smaller emission beam spot, resulting in a higher display resolution. In addition, their high crystalline quality improves the proficiency of radiative recombination. However, despite these advantages, the challenges currently encountered are improving the conversion efficiency and device stability. Lieber and co-workers comprehensively explained that the green emission from n-CdS NW on a p-Si substrate is owing to the FP cavity modes formed inside the NWs during their growth *via* the VLS mechanism. Consequently, the green light lasing by the n-CdS NW heterojunction with p-Si showed a good conversion efficiency.^{336,352}

The electroluminescence (EL) of CdSe/ZnS QD (QLED) and multilayer micro cavity-based QD (MQLED) was used to study the performance of green and blue LEDs (Fig. 18b and c). The CIF diagram of MQLED shows the efficiency achieved for different colors for RGB (Fig. 18a). The significantly improved optoelectronic performances of MQLEDs demonstrate that the optimization of optical properties plays the same important role in determining the EL performance as that of electrical properties. Meanwhile, the microcavity structure can be applied

in low-threshold lasers, cavity quantum electrodynamics, biological detection, and high-performance filters. A further improvement in its *Q* value can lead to an extremely narrow FWHM and a low-loss optical resonator, which indicates that a high *Q* microcavity is a promising and practical structure for achieving electrically pumped QD lasers.³⁷² QDs as a display medium for holographic 3D applications with hydrogel composites exhibit very attractive properties such as high transparency, highly saturated color, fast response, low cost, simple process and scalable and healable characteristics. According to the results, the fabricated PVA hydrogels with different QDs are potential smart display mediums for colorful, real-time 3D displays and have potential in other intelligent optical devices.³⁷³

4.2 Electronic and sensing applications

4.2.1 FETs.

Field effect transistors (FETs) are building blocks in many devices such as integrated circuits, digital data storage devices and flat panel displays. FET devices fabricated from 1D single nanostructures such as NWs/nanoribbons have been established to demonstrate superior performances owing to their high crystal quality, reduced phonon scattering, device material decoupled from the substrate material, large surface-to-volume ratio and quantum confinement.^{14,18,39,374} Jie *et al.* fabricated CdSe NR on SiO₂/p⁺-Si for a single nanostructure FET on a substrate and measured the gate-dependent current and voltage relation, which is also known as the gating effect.^{375,376} The conductance of the nanostructure increased with an increase in the gate voltage for n-CdSe owing to the presence of selenium vacancies. Under illumination, the generation of e-h pairs significantly increases the conductivity, which is related with the increase in carrier concentration under light,



shifting the threshold voltage in the negative direction together with a decrease in the on-off current ratio. Moreover, the 1D system has a high surface to volume ratio, which directly affects the transport properties by the surface absorption and desorption of atmosphere. Surface absorption from the atmosphere increases the resistance, or in other words, a depletion layer is formed near the surface, which decreases the carrier concentration/conductivity under bias condition. Thus, 1D nanostructures can be effectively utilized for gas sensing, chemical sensing and biological detection. Binary and ternary II-VI material single nanoFETs have been extensively explored for potential applications.^{377–382} Nano-network FETs or integrated nanoFETs are promising materials for logic circuit, storage and sensing applications. Fig. 19a shows a schematic diagram of a CdSe nanoribbon as a logic circuit, in which the real intensity measurement system with the fiber tips together with a photomultiplier tube as the detector unit is attached around the single structure. The top inset in Fig. 19a shows the dark field image of a square nanostructure having a size of $25 \times 25 \mu\text{m}$. The fiber tips with a diameter of 250 nm are arranged around the nanostructure. The bottom right inset in Fig. 19a shows a schematic diagram of the top view measurement. The emission intensity at the edges is position-dependent, which changes along the top edge (black square) and right-side edge (red circle) clockwise, respectively, as shown in Fig. 19b, above the threshold intensity level. The first four columns represent different excitation positions of (1000) for region I, (0100) for region II, and so on. The columns with numbers from 0 to 7 represent actual emitting-out states at eight different edge positions corresponding to the different excitation regions. In each table unit, the real emission intensity is given in brackets, as shown in Fig. 19c. Therefore, the directional light propagation and its emission is along the normal and perpendicular axes to form the localized light selection. Based on the light distribution pattern guided rule, an optical logic encoder was successfully developed by Li *et al.* The basic logic element '1' is defined by the real intensity of the illumination part being higher than the threshold, whereas '0' corresponds to the dark noise level.³⁸³ The large high-intensity difference guarantees the validity of a signal for communication. Moreover, its integration in nanodevices was demonstrated by Ma *et al.*, in which nano-inverters with a high voltage gain were fabricated by combining two CdS NW MESFETs for NOR and NAND gates

as integrated devices.³⁷⁹ He *et al.* also described the realization of logic circuits by synchronizing two top-gate CdSe:In NW FETs, which improved the inherent photo-response properties of CdSe NWs.³⁷⁸ The $I_{\text{DS}}-V_{\text{GS}}$ curves of two gated FETs show the hysteresis sweeping rate of 6 V s^{-1} . The series and parallel connection of two gate FETs act as the basic logic circuit of 'AND' and 'OR', respectively. The logic circuits demonstrated a great enhancement in performance, stability, operational speed and high reproducibility. A $\text{CdS}_x\text{Se}_{1-x}$ nanoribbon with lateral heterostructures demonstrated high-performance photo-detection with high $I_{\text{ON}}/I_{\text{OFF}}$ (10^6).⁷³ Li and Junpeng also reported the fabrication of a $\text{CdS}_x\text{Se}_{1-x}$ nanoribbon and $\text{CdS}_x\text{Se}_{1-x}$ NB-based FET, respectively. Under illumination, the overall performance of the ternary and metal-doped binary and ternary alloys improved due to e-h pair generation, which increased the overall device conductivity. Therefore, an integrated circuit with binary and ternary II-VI alloys was experimentally realized but achieving high-density integration without compromising the performance of individual structures is still a challenge that needs to be addressed together with low cost technology device fabrication. This is because high-density integration on the nanoscale for multiple nanostructures involves complicated fabrication steps, which is not only costly but also manipulates the performance of individual nanostructures.^{377,384}

4.2.2 Sensing in bio-medical applications. Lieber and co-workers extensively reported the fabrication of nanoscale transistors that work as nano-bioelectronics, involving the interdisciplinary field of biology and nanoscale electronics to address and overcome the existing challenges in bioelectronics.^{14,18,28} Nanoscale devices exhibit high sensitivity and biocompatibility, opening a new horizon in fundamental biology and healthcare. Devices with the same size as molecules reveal various advantages such as high surface to volume ratio and real-time sensitive detection of proteins and molecules based on high-performance FETs. The surface of a nano-FET can be functionalized with biomolecule receptors that can selectively bind to biomolecule targets. By applying an electric potential at the gate, the nano-FET becomes charged through the biomolecules, and thus its charge carrier density is tuned, leading to a change in electrical conductivity associated with bio-molecular binding events in real time. The same diameter of both the acceptor and receiver provides higher sensitivity detection, which can be further improved with an array of NWs for multiplex detection in the same way. This method is

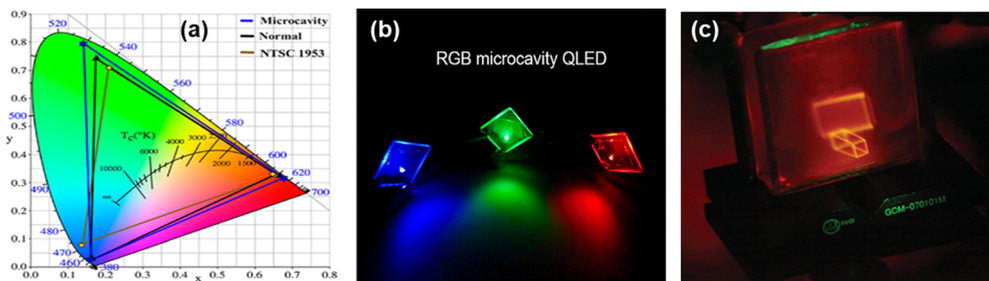


Fig. 18 (a) EL performances of best-performing RGB QLEDs. (a) CIE characteristics. (b) Image of RGB MQLEDs, Copyright © 2020, ACS Publications.³⁷² (c) Digital camera image for holographic reconstruction of cuboid under room light, Copyright © 2014, Wiley-VCH.³⁷³



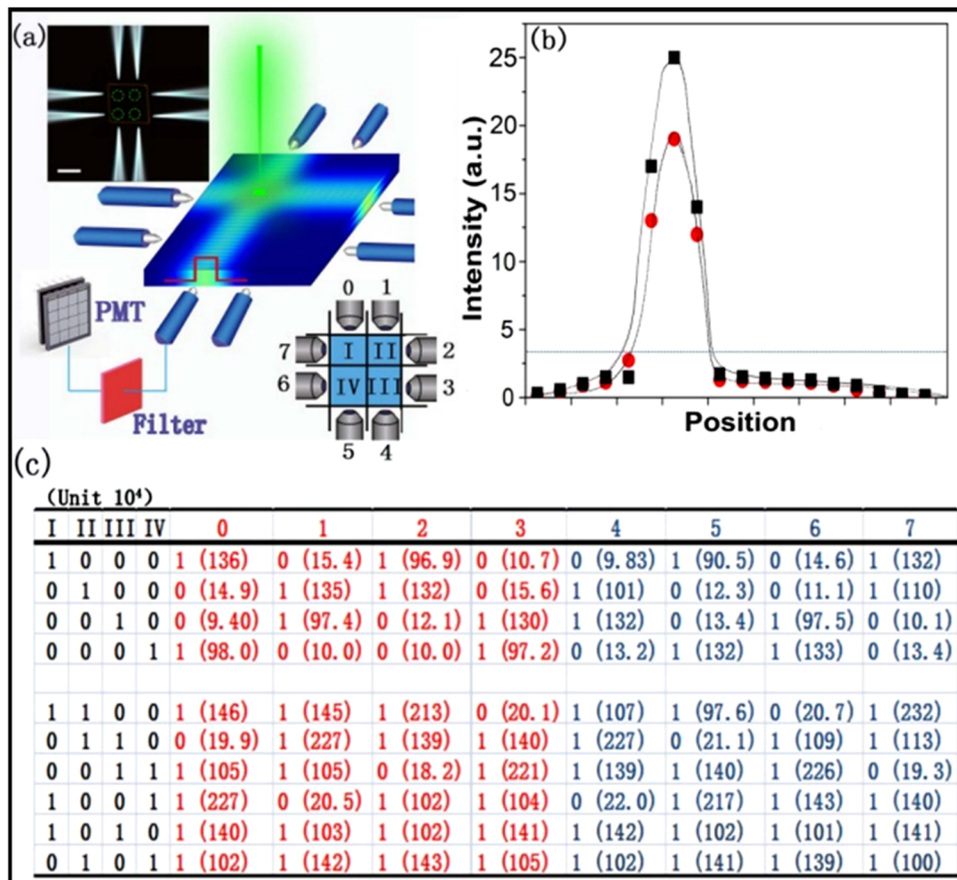


Fig. 19 (a) Schematic diagram of a real-intensity measurement system with the fiber tips together with a photomultiplier tube as the detector unit based on a square nanoplate (dark blue). Top inset: Real dark-field image of a square nanoplate with a size of $25 \times 25 \mu\text{m}$. The fiber tips with a diameter of 250 nm were arranged around the nanoplate. Scale bar is $10 \mu\text{m}$. Bottom right inset: Simple schematic of the top view of the whole microsystem. (b) Lighting position-dependent intensity change along the top edge (black square) and right side edge (red circle) clockwise, respectively. The x-axis represents the scanning position. The line corresponds to the threshold-intensity level. (c) Detailed state of each fiber together with the excitation position. The first four columns represent different excitation positions of (1000) for region I, (0100) for region II, and so on. The columns with numbers from 0 to 7 represent the actual emitting-out states at eight different edge positions corresponding to the different excitation regions. In each table unit, the real emission intensity is given in brackets. Copyright © 2017, IOP Publishing.³⁸³

quite effective for the detection and recognition of various chemical and biological systems.^{40,51,385}

4.2.2.1 FET sensors. Similar to planar FETs, nanofETs can be controlled by applying an electric potential to control the conduction in their channels, making nano-FETs ideal candidates for chemical and biological sensing.³⁷⁴ For example, in p-type Si NW, the receptor (chemical/bio-molecule) binding of molecules with negative charges leads to the formation of an accumulation layer in its channel, resulting in an increase in conductance. However, binding of molecules with positive charges will deplete the holes in the active layer, which will reduce the conductance. Thus, NW FETs produce real-time directional electrical information on biological events related to binding/unbinding, enzymatic reactions, and electron transfer, with superior capabilities for emerging platforms to investigate biological samples. Also, the SiO_2 layer on the surface of Si NW is nontoxic and can sense the hydrogen in lipid bilayers with and without ligand-gating, amino groups and silanol

groups (Si–OH). Silicon NWs have been extensively used owing to the fact that they are mature materials in comparison with other II–VI materials and widely used for bio-molecular sensing. However, ZnO NWs have also been used for the development of FET biosensors.^{351,386}

4.2.2.2 Protein sensors. The sensitive detection of proteins related to disease markers offers substantial potential for the diagnosis and treatment of diseases, where real-time protein sensing was first demonstrated in 2001. Nano FETs have the capability to sense biotin receptors, detect streptavidin (concentration down to 10 pM), biotin-mono-clonal anti-biotin binding and calmodulin in real time. In a recent study, a basic approach was used to demonstrate the successful concentration-dependent detection of cardiac troponin T390, SARS virus nucleocapsid proteins, and bovine serum albumin together with genomics and proteomics for diagnosing complex diseases such as cancers. Moreover, the availability of altered



biomarkers matched with different phases of diseases can help in their initial detection and robust diagnosis.^{18,28,387}

4.2.2.3 Nucleic acid detection. A major difference between nucleic acid and protein detection is the high density of negative charges on the nucleic acid phosphate, which increases the high ionic strength barriers to screen the repulsion in the active channel region and allow binding when DNA or RNA is used as the probe molecule. However, high ionic strength solutions have a short Debye screening length, which can make detection difficult. Therefore, solutions that have a high ionic strength binding/screening require the use of neutral charge peptide nucleic acids (PNAs), which exhibit excellent binding affinity with DNA at lower ionic strengths. Si NWs with PNA probe molecules exhibit time-dependent conductance changes depending on the selective binding of the complementary target DNA at low concentrations (10 fM). In addition to sensing protein binding/unbinding, the real-time detection of nucleic acids (e.g., DNAs and RNAs) has been successfully carried out using Si and GaN NW FET devices.^{387,388} Multi-segment CdTe–Au–CdTe NWs were used in devices in which the Au segments were modified with thiol-terminated DNA probes, where binding to these probes induced a conductance change in the overall device structure.^{389,390}

4.2.2.4 Virus detection. Viruses are the main origin of infectious diseases, which continue to be the main cause of death globally. In this case, the successful treatment of viral diseases often depends on the quick and precise identification of viruses at ultralow concentrations in living organisms. An Si NW-based nanoFET-based was successfully employed to detect the Influenza A virus for the first time. By recording the analogous electrical conductance fluctuations upon binding/unbinding of the virus particles to monoclonal-antibody-modified Si NWs, the selective detection of Influenza A at the single-particle level was successfully demonstrated. Another example of virus detection is the identification of dengue. A specific nucleic acid fragment with 69 base pairs derived from the dengue serotype 2 virus genome sequence was selected as the target DNA and amplified by the reverse transcription polymerase chain reaction (RT-PCR). The hybridization of the target DNA- and PNA-functionalized Si NW FET sensors increased the device resistance, leading to a sensitivity limit down to 10 fM.^{18,29,389,391}

4.3 Spintronics

Spintronic-based devices open a new chapter in electronic devices scaled down on a single chip to follow Moore's law, which are controlled by the spin degree of freedom of charge carriers. The realization of these devices in 1925 was enabled by the continuous optimization of the nature of TM ions in various semiconductors. However, there is still room for improvement to explore their fascinating functionality for various electronic applications especially 1D systems. The unique optical response of low-dimensional and spin-controlled behavior opens new horizons in spintronic devices based on spin-dependent electronic properties of FM behavior.³⁹² The incorporation of TM ions

in II–VI semiconductors results in spin splitting of their conduction and valence bands through the exchange coupling of sp–d between the delocalized carriers and core spin. Wide bandgap semiconductors such as GaN and ZnO also exhibit the spin–carrier interaction above room temperature. Ternary alloy formation or heterojunction devices can be frequently used for spin-dependent transport across the interface between different materials. Mn ions introduce spins and holes in the valence band in GaAs, point defects in SnTe and MnTe, and shallow acceptor impurities in CdMnTe and ZnMnTe, where the polarized electrons permit the injection of spins in normal semiconductors. The FM order of spin ions is mediated by the carriers in Mn-based DMS in II–VI semiconductors. The spin–spin interaction becomes dominant in the presence of the divalent Mn ion through hybridization between the anion and d-states of the TM ion. The super-exchange of the divalent TM ion may lead to the formation of AFM coupling.^{393,394} The spin–carrier interaction in n-type and p-type semiconductors is related to the spin-polarized electrons or holes in DMS. The spin polarization of injected carriers can be detected by circular polarized light, also known as spin LEDs. However, the spin relaxation time for holes is much shorter than that for electrons in QW based spin-LEDs. Moreover, neuromorphic computing uses brain-inspired principles to design circuits that can perform computational tasks with superior power efficiency based on spintronic nano-devices, in which electronic devices create artificial neurons and synapses with efficient energy integrated circuits. Spintronic-based neuromorphic computing can effectively perform pattern recognition in associative memory.³⁹⁵

At a low temperature (5 K), CdTe/ZnTe QDs doped with Cr ions show spin relaxation under a strong resonant field, which can be tuned by the Cr ion due to the Stark effect. The exchange interaction between the spin atom and exciton further splits the energy level of the Cr ion due to the high polarizability of the crystal field. The carrier dynamics can be understood by employing transient absorption spectroscopy to observe the relaxation time and non-equilibrium distribution of the Cr spin population with circularly polarized luminescence spectra. The Cr ion interaction with phonons within the system results in strong spin–phonon coupling at 5 K, which is an important step forward to control the TM ion coherent spin for future spintronic devices. Similarly, the Cr spin is coupled with excitons near the valence band and shows nano-magnetic behavior due to the optical pumping of electron-Cr and hole-Cr. Resonance and non-resonance optical pumping can be directly used as spin memory (write and erase, respectively). Therefore, the spin channel can be studied with the polarized emission that responds to the relaxation channel of the Cr ion in II–VI semiconductors. Also, the optical pumping and power-dependent PL intensity-related temperature effect directly affect the hole-Cr spin near the valence band and electron-Cr local generation of phonons.¹⁵⁸

4.4 Future research direction

In this review, we focused on the optical properties of low-dimensional II–VI semiconductor nanostructures and their potential application over the past two decades, which have



attracted significant attention as nanoscale active and passive devices in optoelectronics, bioelectronics and electrochemical applications. The interdisciplinary nature of the research offered herein signifies the future of scientific discovery, where the boundaries among chemistry, physics, biology, and engineering are no longer relevant. We tried to cover the intensive research on 1D II–VI binary, ternary alloy and TM-doped semiconductors and understand their luminescence properties related to excitons, photons and phonons within the nanostructures and their coupling with quanta such as photons and plasmons. The light–matter interaction provides fascinating information and tunable optical emission based on the molar ratio and ferromagnetic semiconductors, which usually arises from carrier–spin interaction, spin–spin coupling and sp–d hybridization between anions. Therefore, the understanding of the chemistry and physics of nanostructures can reasonably support the control of their growth pattern, crystal structure and compositional complexity for practical applications toward novel technologies. Luminescence behavior shows the dynamic nature of the magnetic ions through spin-splitting, bandgap narrowing and magnetic polarons. Transition metal ion doping in II–VI semiconductors results in phonon–spin, spin–spin and exciton–spin interactions, producing EMP, BMP or LEMP and exciton antiferromagnetic polarons in 1D and 2D systems. A pulse laser produces bosonic lasing owing to the coherent magnetic exciton Bose–Einstein condensation in a low doping profile, which favors ferromagnetic coupling. Moreover, a high doping profile scatters the coherent EMP due to bound excitons, coexisting EMPs and e–h plasma formation at the surface of the nanostructure. The coupled optical microcavity effect produces enhanced photonic Bloch oscillation in 1D nanostructures with multipeak emission or lasing at low threshold pumping depending on the spatial distribution of photons, corresponding to exciton–polariton formation. Nanostructures such as axial, radial/coaxial modulated, branched and kink nanostructures are important as next-generation building blocks for bioelectronics, photonics and electronic circuits. Increasing the number of devices on a single chip without compromising their properties is important to understand semiconductor/metal/insulator interfacial or hybrid structures with organic semiconductors or 2D materials, which can lead to an enhancement in the device performance with improved responsivity, detectivity and quantum efficiency. Therefore, different advanced synthesis techniques aim to control the size, morphology and composition of II–VI semiconductor nanostructures to ensure their reproducibility, yield, high quality and cost effectiveness as suitable materials for large-scale device fabrication. The spin behavior in II–VI semiconductors opens a new horizon for DMS research on flexible electronics, wearable devices, and bio-sensing platforms, leveraging their unique optical and electronic properties. Yang, Leiber and other research groups focused on 1D nanostructures for biomedical applications such as biochemical sensors, extracellular and intracellular electrical sensors, molecule carriers, injectable electronics, biosynthesis and optical neuromodulation.

5. Outlook

In this review, we provided an overview of TM ion doping in II–VI semiconductors and their fundamental properties, especially their exceptional optical properties and promising optoelectronic applications. Moreover, we highlighted the rapid and prominent development in the research activities within last two decades considering the increasing importance of 1D nanostructures in the current technology market. Recent advances in 1D nanostructures such as NWs and NBs open up opportunities for future generation nanodevice applications. The interdisciplinary nature of the research offered herein signifies the future of scientific discovery, where the boundaries among chemistry, physics, biology, and engineering are no longer relevant. In this review, we focused on the luminescence of 1D nanostructures based on binary and ternary alloys of II–VI semiconductors and their composition gradient, together with TM ion doping in 1D II–VI nanostructures. Based on this, we highlight some important issues to promote the practical applications of II–VI nanostructures, as follows:

1 In the past decade, researchers have realized substantial progress in the synthesis and characterization of II–VI nanostructures in the forms of NW/NT/NB, which have been grown by both physical and chemical methods. The properties of nanostructures are strongly determined by their geometries, stoichiometric ratio, structures, and surface states. The chemical synthesis of NPs, QDs, and others results in unique electronic and optical properties due to the quantum size effect owing to their nanoscale, together with low cost and less energy consumption, but compromises their crystal quality and surface effect. Physical methods for the synthesis of 1D nanostructures such as the VLS mechanism offer highly crystalline and controlled nanostructures, but their composition, orientation and reproducibility are still challenging to control at high temperature. 1D nanostructures of II–VI ternary alloys have been inspired by the rapid development of technology in the last decade; research progress regarding material synthesis, fundamentals of optical properties, and promising electronic properties for optoelectronic and photonic applications. Although most ternary alloy nanostructures and their corresponding applications are new and their technological development is in its infancy, research on ternary alloys is highly attractive and rewarding. Recently, many researchers have realized the advantages of quaternary alloys, blended organic–inorganic hybrid devices and TM ion incorporation to fulfil future device application demands.

2 TM ion doping in 1D nanostructures results in unique optical properties and promotes various electronic and photonic applications. The incorporation of TM ions results in spin–carrier injection, spin–spin coupling among TM ions and p–d hybridization (involving p-orbital anions and d-orbital of TM ions), making II–VI semiconductors suitable for DMS. Exciton magnetic polarons and bound magnetic polarons or localized EMPs are outstanding zones for DMS that need an enormous amount of study to explore Zn chalcogenides and Cd chalcogenides in 1D systems.



(a) TM ion doping in 1D II–VI semiconductors makes them suitable candidates for DMS-related applications. The TM ion concentration can tune the luminescence spectra under different laser excitation powers. A low TM concentration may lead to the interaction between spin ions with free excitons and phonons to form independent EMPs. Even a low TM doping concentration can tune the bandgap and related d–d transition in the emission spectra at room temperature. A low TM concentration with high laser excitation can show bosonic lasing owing to the coherent collective EMP transition from magnetic exciton Bose–Einstein condensation at room temperature. LO phonon and spin ion interaction exhibits the bosonic lasing reported in 1D nanostructures that appears for long range coherent coupling of local EMPs. Moreover, a high TM ion doping concentration scatters the coherent EMP due to (i) bound carriers, (ii) coexisting EMPs and (iii) e–h plasma formation at the surface of nanostructures.

(b) In low-dimensional systems, the free excitons or polaronic excitons or bound excitons will be prominent with TM ion doping in II–VI semiconductors at room temperature. Moreover, the doping profile will enhance the electron–phonon interaction especially for 1D NWs. In ZnO:Mn, the polaronic excitons bind with free excitons for the large excitonic binding energy material to show bipolaronic exciton condensation under fs laser excitation. Also, coherent and high power favor multimode lasing emission at room temperature. Moreover, the LO phonons in the *c*-axis contribute to the condensate formation for BEC of excitons at room temperature.

(c) Optical characterization tools have the capability to observe the dynamic nature of magnetic ions, which can provide strong evidence of the spin splitting, bandgap narrowing and bound magnetic polarons in nanostructures. Strong spin–spin interactions demonstrate additional broadening of the bandwidth or strong inhomogeneous broadening, which is the standard electron paramagnetic resonance of spin ions in DMS. Raman scattering provides suitable information about the bound magnetic polarons at low temperature, which shows the strength of the antiferromagnetic interaction between nearest the neighbor d–d interaction. Time-resolved spectroscopy provides the dynamics of local spin ordering with either bound magnetic polarons or self-trapping related to exciton magnetic polarons.

(d) A deeper understanding of material properties and device behavior is required to determine significance of performance reliability and revolutionary applications from the novel and innovative fundamental properties manifested at the nanoscale. In particular, more 1D nanostructure research should be devoted to gaining a deeper understanding of the interaction properties of excitons, plasmons and phonons within II–VI semiconductors. Therefore, we can explore their coupling mechanism with other quanta, which is influenced by their optical properties, and the related applications greatly influenced by the interactions and coupling strongly depend on the optical and emissive properties of II–VI materials.

(e) Additionally, the development of 1D II–VI nanostructures with CVD is important for various photonic applications. Doping in 1D structures can afford active photonic waveguides to tune their emission and propagation light wavelength with the

propagation distance and endowing the wire end with a tunable wavelength. The tunability comes from the strong exciton–phonon coupling during the light propagation. For example, Sn:CdS, CdSSe NW array and branched-like structures for 1D periodic photonic applications related to exciton–polariton. The coupled optical microcavity effect produces enhanced photonic Bloch oscillation in these wires and multiplex emission or lasing at low threshold pumping. The photonic propagation within the wires show a clear spatial-dependent photon distribution and correlation.

3 1D nanostructures of II–VI ternary alloys have been inspired by the rapid development of technology during the last decade; research progress regarding material synthesis, fundamentals of optical properties, and promising electronic properties for optoelectronic and photonic applications. Although most of the ternary alloy nanostructures and their corresponding applications are still new and infancy of technological development, the research on ternary alloy remains highly attractive and rewarding. In this case, an important foundation is developing efficient and low-cost technology for the synthesis of ternary II–VI 1D nanostructures with high quality and large quantities. Therefore, it is necessary to adopt effective preparation techniques that are more controllable with the simultaneous precision of composition, orientation, location, alignment and dimensions. Therefore, the physical properties of nanostructures are more suitable for massive-scale device integration for a single structure. To achieve the required task, ternary and quaternary or poly-compounds have been designed and synthesized to enable a full-spectrum response. Recently, many researchers have realized the advantages of quaternary alloys. Additionally, the development of ternary II–VI 1D nanostructures is also important due to their tunable bandgap, which is applicable in electronic and optoelectronic circuits. The high surface-sensitivity of 1D systems is owing to their large surface to volume ratio make them appropriate as high-sensitivity gas, chemical and biological detectors. 1D hybrid nanostructures are highly appropriately in various optoelectronic applications such as type-I and type-II junctions; type-I junctions exhibit high fluorescence and light emitter yield (LEDs and lasers), while type-II junctions facilitate the formation of e–h pairs with improved energy conversion efficiency (photodetectors and solar cells). Different prototypes have the capability to assemble and integrate individual structures into functional devices on a large scale. However, achieving high-density, large-scale, integrated devices at a low cost is still a bottleneck.

4 Another important emerging phenomenon in 1D II–VI material research is the essential engineering of large-scale architectures at a low cost without compromising the material properties on the nanoscale rather than focusing on individual structure. For the commercial adaptation of 1D systems, the development of device design must emphasize low cost, environment-friendly features and generation of minimum waste with the utilization of more expensive elements together with the necessary spatial and temporal resolution to implement quality control throughout the fabrication process. This control will facilitate the development of the next era of nanotechnology in the environmental and biomedical fields with economic benefits to society and realization on a global



scale. The Lieber group synthesized IV, III–V and II–VI 1D nanostructures analogous to biological systems and reported their biomedical application, together with a detailed understanding of individual nanostructures and their interfacial interactions for biochemical sensors, extracellular and intracellular electrical sensors, molecule delivery, injectable electronics, biosynthesis and optical neuromodulation for brain science.

6. Conclusion & future prospective

In this review, we discussed magnetic polarons and their interaction with excitons, electrons and spin ions for TM ions doped in various II–VI semiconductors for 1D DMS nanostructures. Moreover, we explored the formation of EMP, BMP and LEMP and their related dynamic spectral response together with their collective response such as bipolaronic BEC excitons at room temperature. Strong electron–phonon interaction materials coupled with spin ions show stable EMP formation. High excitation with a low concentration of magnetic ions may lead to the formation of bosonic lasing due to collective EMPs, which has interesting application in future spin-related optical devices at room temperature. The active photonic waveguide property comes from the strong electron–phonon coupling during light propagation. The coupled optical microcavity effect produces enhanced photonic Bloch oscillation in wires and multiplex emission or lasing at low threshold pumping. TM ion doping in II–VI ternary alloy nanostructures is still in its infancy of technological development. In particular, we believe that more comprehensive and intense research must be conducted to gain a deeper and better understanding of the luminescence properties related to the excitons and phonons within II–VI 1D nanostructures and their coupling with quanta such as photons and plasmons. Also, a deeper understanding of the luminescence properties of binary and ternary alloys, heterojunctions and architectures with metal ion doping in II–VI 1D nanostructures is needed to optimize their overall optical and emissive properties for the emergence of revolutionary applications from the novel fundamental properties. Each of these applications of 1D II–VI nanostructures has been developed based on years of research and fundamental understanding of entire material-related chemistry, physics, and engineering for their practical implementation.

Data availability

The data that supports the findings of this study are available within the review article.

Conflicts of interest

The authors have no conflict of interests to disclose.

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