Colloidal nanocrystals for photoelectrochemical and photocatalytic water splitting

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Abstract
Colloidal nanocrystals (NCs) are among the most modular and versatile nanomaterial platform to study emerging phenomena in different field thanks to their superb compositional and morphological tunability. A promising, yet challenging, application involves the use of colloidal NCs as light absorber and electrocatalyst for water splitting. In this review we discuss how the tunability of these materials is ideal to understand the complex phenomena behind storing energy in chemical bonds and to optimize performance through structural and compositional modification. First, we describe colloidal synthesis approaches that offer a high degree of control over single material NCs and NC heterostructures, including examples on the role of the ligands to modulate size and shape. Next, we focus on the use of NCs as light absorbers and catalysts to drive both the hydrogen evolution reaction (HER) and the oxygen evolution reaction (OER) along with some of the challenges related to the use of colloidal NCs as model systems and/or technological solutions in water splitting. We conclude with a broader prospective on the use of colloidal chemistry for new material discovery.

1. Introduction
Much progress has been made since Honda and Fujishima discovered that TiO\textsubscript{2} is capable of splitting water in 1972 [1]. However, the storage of sunlight into chemical bonds, namely artificial photosynthesis, cannot be efficiently completed yet with robust and Earth-abundant materials. The combination of modeling and experiments have shown that higher efficiencies are achieved with dual tandem absorbers design, instead of one single semiconductor material [2,3]. Dual tandem absorber configurations include two properly matched light absorbers, that generate the photovoltage needed to split water, and catalysts specifically optimized for water oxidation or reduction, that are interfaced with the semiconductor light absorbers [2,3]. Yet, the coupling of photons and electrons, protons and electrons and even the catalytic mechanism behind water splitting have not been fully elucidated. The knowledge gap becomes even larger when looking at the carbon dioxide reduction reaction. Improving the performance of sustainable solar-to-fuels conversion requires probing and understanding these nanoscale phenomena with advanced scientific techniques. This understanding must then be translated into control of the functionality and performance of the materials and chemistry that govern the water splitting and the CO\textsubscript{2} reduction reactions. Here, the ability to produce materials with tailored structure and composition is key to derive meaningful structure/function relations. Conventional synthetic techniques (i.e. solid state reactions, sol-gel processing, electrochemical deposition) often provide

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samples that are heterogeneous in nature. Because the materials can undergo unknown structural changes during the photoelectrochemical and electrochemical testing, significant effort are expected to understand what makes the light absorber, the catalyst and the combination of both efficient. Precise engineering and manipulation of the active materials is instead needed to elucidate the relevant structural parameters governing their catalytic or photocatalytic behaviour.

Colloidal nanocrystals (NCs) are among the most modular and versatile nanomaterial platform to study emerging phenomena in different field due to their unprecedented compositional and morphological tunability [4–25]. A collection of transmission electron microscopy (TEM) images of various NCs is shown in figure 1. They are constituted by an inorganic core, with tunable composition, size and shape, and a shell of amphiphilic surfactants or ligands. The latter are key in controlling the NC morphology. Furthermore, they guarantee the dispersability of colloidal NCs in different solvent, thus allowing their deposition onto various substrates or their integration into various matrices (i.e. inorganic or polymeric). These processes are not possible with powder-form nanoparticulate or NCs synthesized by chemical vapor deposition (CVD) and molecular beam epitaxy (MBE).
The use of colloidal NCs in various catalytic reactions, such as CO oxidation and oxygen reduction, has demonstrated their huge potential to function as model systems to propose new catalytic schemes or to elucidate the role of defects, catalyst/support interface, crystalline phase [26–30]. As for the water splitting reactions, to date most of the work with colloidal NCs has focused on heterostructured NCs comprising a metal catalyst coupled with a semiconductor light absorber to drive the sacrificial hydrogen production.
While long term stability still remains to be solved, compelling efficiencies have been achieved. More recently, colloidal NCs, mostly belonging to the phosphide family, have been used as hydrogen evolution catalyst. Far less studies have focused on the oxygen evolution reaction, yet their number is increasing. Motivated by the growing interest in the field by the nanocrystal community, we have decided to give an extensive overview of the state-of-art work relative to the application of colloidal NCs for photocatalytic (no bias applied) and photoelectrochemical (bias applied) water splitting. The aim is to help to identify the challenges and the research areas which deserve more attention so to direct future efforts towards those. We omit to review studies on nanocrystalline materials prepared by other routes and used in suspensions, instead of uniform colloidal solutions. Therefore, the work from the Domen and the Osterloh groups are not discussed in the present contribution [31–35]. First, we briefly overview single material NCs, NC heterostructures and some of the approaches to synthesize them. Next, we focus on the use of NCs as light absorbers and catalysts to drive both the hydrogen evolution reaction (HER) and the oxygen evolution reaction (OER). We also describe some of the challenges related to the use of colloidal NCs as model systems and/or technological solutions in water splitting. Finally, future opportunities of using colloidal chemistry for new material discovery are succinctly highlighted.

2. General concepts in colloidal nanocrystal synthesis
2.1 Single-material nanocrystals
Colloidal synthesis is a surfactant-assisted approach which has proven to afford a superior tuning of the size, the shape and the composition of nanocrystalline materials compared to other techniques [4–25,36,37]. In colloidal synthesis, the growth medium is a liquid mixture of ligands and surfactants and the whole synthesis process is usually carried out under inert atmosphere, as sketched in figure 2A. Some of the surfactants commonly used include alkylthiols, amines, carboxylic and phosphonic acids, phosphines, phosphine oxides and various coordinating solvents, such as ethers or alkenes [4–25,36,37]. The general scheme of a colloidal synthesis involves thermally activated reaction of organometallic precursors (that carry the atomic species necessary to build the desired material) in the presence of selected surfactants (figure 2B). Once the synthesis is activated, the highly reactive species that are generated, commonly referred to as the “monomers”, induce the nucleation of NCs and sustain their subsequent enlargement. The surfactants play several key roles during the synthesis. Indeed, they form complexes with the monomers, dictating their actual chemical potential in solution [9–13]. Simultaneously, they participate in an adsorption/desorption dynamics on the surface of the growing clusters, which prevents aggregation and uncontrolled growth. Mechanistic studies have shown that monodisperse NCs are produced when a burst of nucleation that enables separation of the nucleation and growth processes is combined with diffusion-controlled growth. Satisfactory tailoring of the NC size and size distribution can be achieved by balancing the relative depletion of monomers between the nucleation and the growth stages either with the aid of suitable techniques (i.e. the “hot injection”), or profiting from the particular reactivity of the system (i.e. “delayed nucleation”), or exploiting digestive-ripening [5,7,38]. It is also worth mentioning that surfactants can affect the specific surface energy of the growing NCs, which has important implications in the tuning of their shape [5,7,8]. Indeed, facet-preferential li
Figure 2: Schematic of (A) the setup used for colloidal synthesis consisting of a three-necked flask, a heating mantle and a condenser; (B) the synthesis of single material NC; (C) the synthesis of hybrid NCs by seeded growth approach.

Gand adhesion can modify the relative growth rates along the various crystallographic directions and/or can favour selective elimination of unstable surfaces by triggering oriented attachment of particles along defined crystallographic directions. In the absence of additional circumstances that can interrupt growth symmetry (e.g., formation of soft micelle templates, the presence of foreign particle catalysts, or the application of external electric or magnetic fields), surfactants remain mostly responsible for the formation of NCs in a variety of anisotropic shapes, such as cubes, polyhedrons, rods, wires, polypods, and rings, as shown in figure 2.

2.2 Nanocrystalline heterostructures or hybrid nanocrystals
NC heterostructures or hybrid NCs (HNCs) are multi-material nano-objects, consisting of two or more different material domains, often interconnected through epitaxial interfaces [14–19,39–43]. One of the most common synthetic approaches to these multicomponent nanomaterials is “seeded growth”. “Seeded growth” relies on performing preferential heterogeneous nucleation and growth of a second material on preformed NC seeds, as shown in figure 2C. Other synthetic strategies include cation exchange, phase segregation, coalescence of an initially amorphous shell. Figure 3 gives examples of HNCs with different topologies and highlights once again the high level of control achievable by colloidal chemistry. These different topologies are dictated primarily by the lattice mismatch between the two domains; however, different synthetic strategies can impact topology as well. As one example, by tuning the reactivity and/or concentration of the monomer of the second material, it is possible to decide whether, starting from one rod-like semiconductor seed, either both ends of the rod, only one end or the longitudinal side is decorated with the second domain [15,39]. It is helpful to consider that the creation of HNCs by seeded growth in solution phase shares some similarities with the more traditional heteroepitaxial deposition processes accomplished by MBE and CVD techniques, in terms of growth mode [44,45,46]. However, what clearly distinguishes colloidal seeded growth from MBE/CVD approaches is the presence of surfactants that indeed alter the surface energy term considerably. This is one of the reasons why strain-
related hetero-epitaxial growth constraints can be greatly overcome by colloidal chemistry. As an example, epitaxial interfaces have been observed in TiO$_2$-Fe$_2$O$_3$ even with a 13% lattice mismatch [40].

**Figure 3.** TEM examples of hybrid NCs with different topological distribution of their component material sections. (A) Pt-decorated CdSe@CdS octapod NCs: (a) TEM image of an ensemble of CdSe@CdS octapod nanocrystals, the scale bar is 200 nm. Representative images of (b) a single Pt-free octapod nanocrystal, (c) a single Pt-tipped octapod nanocrystal and (d) a Pt-covered octapod nanocrystal. Scale bars in (b), (c) and (d) are 20 nm [14]; (B) γ-Fe$_2$O$_3$/TiO$_2$ NC hybrids [15]; (C) BiVO$_4$/TiO$_2$ heterostructures [16]; (D) Pd@Cu core@shell nanocubes [47]; (E) Ag-Pt-Fe$_3$O$_4$ trimers [17]; (F) RuRh-Cu$_2$S nanocages [18]. Reproduced from [14,18] with permission of The Royal Society of...
3. Colloidal nanocrystals as light absorbers

In recent years, the reports on colloidal NCs used as light absorbers to drive the water splitting have notably increased, with the studies on HER greatly outnumbering those on OER. While in this review we will limit our discussion to fully inorganic colloidal HNCs, we notice that light absorber NCs have been successfully coupled with enzymes, specifically iron hydrogenases, and molecular species as co-catalyst for hydrogen generation [48–50]. Studies relative to dye reduction or oxidation are also not included in this contribution.

3.1 Colloidal nanocrystals as light absorbers for HER

The general approach to drive HER with colloidal NCs has been to decorate II–VI semiconductor NCs (or quantum dots, QDs) with metallic nanoparticles. The latter promote charge separation of the photogenerated carriers and also act as the site for hydrogen generation. Most of these semiconductor/metal HNCs have been tested as photocatalysts dispersed in solution, with the exception of few examples [51,52]. To perform the testing in aqueous solutions, the native hydrophobic ligands functionalizing the NC surface are exchanged with shorter hydrophilic ligands, the most common ones being mercaptoundecanoic acid (MUA) with some examples using cysteine and mercaptopropionic acid (MPA). Several reviews on semiconductor-metal colloidal NC heterostructures for HER have been published in the past years [53–55]. We refer to them for more details, especially with regards to studies of charge carrier dynamics which are not included here. For sake of clarity, literature has been grouped by classes of the light absorbing semiconductor materials. It should be noted that most of the papers reviewed in this section reports performance as external or apparent quantum yield (EQE or AQE) of hydrogen evolution and uses a visible laser as the light source. The AQE is defined as two times the rate of collection of H$_2$ to the incident photon flux, given by equation (1). We have reported the AQE values at the irradiation wavelength, when data were available, so to have a measureable comparison between different photocatalysts under similar experimental conditions. Other ways to report the efficiency for photocatalytic hydrogen generation include quantum yield (QE) and solar-to-hydrogen (STH) energy-conversion efficiency [31,56]. QE is the ratio of the number of “reacted” electrons per number of absorbed photons, as shown in equation (2). Instead, the STH indicates the conversion of the energy in the incident solar irradiation to chemical bonds as H$_2$ through oxidation of water to O$_2$, given by equation (3) [31,56]. In equation (3), 237000 $J/mol$ is the standard Gibb’s free energy of water splitting reaction.

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AQE = \frac{\text{Number of reacted electrons}}{\text{Number of incident photons}} = \frac{2 \times \text{mol} \ H_2/\text{s}}{\text{Number of incident photons}} \tag{1}
\]

\[
QE = \frac{\text{Number of reacted electrons}}{\text{Number of absorbed photons}} \tag{2}
\]
\[
S \text{TH} = \frac{(\text{mol } H_2/s) \times (237000 \text{ J/mol})}{P_{\text{inc}} (W/cm^2) \times \text{Area}(cm^2)}
\] (3)

**CdS and CdSe NCs.**

CdS/Pt and CdSe/Pt HNCs are the first and among the most studied systems so far (figures 4A,B,D). CdS and CdSe nanorods (NRs) have been largely preferred to spherical NCs as they have a lower exciton binding energy and offer the possibility to control the distance between reaction sites. Tailored variation of the NR length, diameter and shape as well as Pt domain size have enabled correlations between the structural parameters and the efficiencies [57–59]. As one example, Berr et al. have showed that the hydrogen evolution rate does not benefit from larger Pt domains (4.8 nm) on CdS NRs when compared to subnanometer sized Pt clusters [58]. Similar results were found for Pt-decorated CdSe tetrapods reported by Sung et al. [59]. The higher surface-to-volume ratio of the small clusters and probably different charge separation kinetics have been identify to play an important role in the higher photocatalytic efficiencies compared to the particles. These findings suggest that efficient hydrogen production utilizing II-VI nanostructures with reduced amounts of Pt is possible. In addition to Pt as metallic co-catalyst, a few studies on CdSe/Au have been reported [60,61]. For example, using a well-defined model system of CdS/Au NRs, Ben-Shahar et al. have studied the effect of the gold size tip on the photocatalytic function, including the charge transfer dynamics and the \( H_2 \) production efficiency [62]. They found a non-monotonic behavior with size of the gold tip, leading to an optimal metal domain size. The optimal value is explained in terms of competing processes where for small tips the hydrogen evolution QY is mainly determined by the rate of electron injection to the metal tip, whereas for large tips it is determined mainly by the water reduction kinetics on the metal surface. These two limits show opposite dependence on the metal domain size, leading to an optimal value as explained by a minimalist and general kinetic model [62].

A few examples including a non-noble metal catalyst have been reported [63–65]. Cao et al. have revealed the potential of CdS/Ni\(_2\)P and CdS/Co\(_2\)P by measuring their activity in an aqueous lactic acid solution [63]. In particular they have achieved a AQE of hydrogen generation of 6.8% under LED light (\( \lambda > 420 \text{ nm} \)) by physically mixing CdS NRs and Co\(_2\)P nanoparticles [63]. As for the Ni\(_2\)P they report only rate of hydrogen generation [64]. Zhukovskyi et al have decorated CdS nanosheets with Ni and have been able to achieve AQE up to 25% under laser irradiation at 405nm for the optimal catalyst loading [65].

Iron-based alternatives are especially attractive because Fe is the most abundant transition metal. Cheng et al. have successfully demonstrated a low-cost yet highly efficient and stable semiconductor CdS/FePt photocatalytic system [66]. An AQE of over 35% at 520 nm was measured. This value was 3-fold higher than that of the control CdS/Pt system.

Because photo-oxidation is an issue for stability of semiconductor NCs, the photo-holes need to be quickly extracted from the surface. Thus, a hole scavenger (or electron donor) is used in all the reports. While alcohols are the most common, it is clear that the redox level of the scavenger with respect to the valence band of the semiconductor NCs will
impact the charge transfer kinetics and therefore the quantum efficiency for hydrogen generation. A nice study by Berr et al. illustrates the positive effect of the increased redox potential of the hole scavenger (MeOH < EDTA< TEA < SO3−), which facilitates a faster hole scavenging, on both the activity and the stability of the CdS/Pt hybrids (figures 4A,D,E) [67]. In particular the authors compare the hydrogen evolution in function of the time during exposure under illumination. They found a correlation between the oxidation potential of the hole scavenger and the stability of the photocatalyst. With TEA and EDTA− the amount of hydrogen product dropped from a maximum of 2.5 µmol and 0.4 µmol to 0.6 µmol and 0.07 µmol after 4h of illumination, respectively. The hydrogen generated in the presence of MeOH was below the detection limit. The best stability was measured in the presence of SO3− as hole scavenger. The conclusion of this study was that the higher the oxidation potential is, the more efficient the hydrogen production and the more stable the nanorods are.

High pH has been demonstrated to increase efficiency and stability of CdS/Pt and of CdS/Ni hybrid NCs [68,69]. The authors have demonstrated that although the yield decreases to approximately 20% after 80 h, it then remains stable for up to 220 h. This means that at high pH the system is protected against photo-oxidation. This happens because of the redox couple •OH/OH operating as a shuttle between the surface of the NCs and the ethanol scavenger. As illustrated in figure 4F, the authors suggest that instead of a direct oxidation of ethanol by the photoexcited hole, the small sized hydroxyl anion, present in large quantities in highly alkaline conditions, easily diffuse to the NR surface where it gets oxidized. The AQE of H2 generation overcomes 50% under 446 nm laser illumination of a dispersion of Ni-decorated, cysteine stabilized CdS NRs in the presence of ethanol as a sacrificial agent. This work establishes the record efficiencies for CdS NCs used as photocatalysts to drive hydrogen evolution.

CdSe@CdS HNCs.

In addition to scavenging them with a sacrificial agent, the holes can be confined to a part of the nanostructure that is acting like a trap. The use of heterostructured CdSe/CdS nanointerfaces has been demonstrated to further increase the QY of hydrogen production through the spatial separation of charges in nonadjacent domains. In CdSe@CdS/metal HNCs, the photogenerated charge carriers are separated with hole in the core (CdSe) and electrons in the shell (CdS), that are eventually transferred to the metal domain to generate hydrogen.

Amirav et al. have assembled heterostructures including CdSe@CdS asymetric core@shell NRs functionalized with Pt nanoparticles (figure 4B)[70]. What is interesting about the asymmetric dot@rod core@shell is that the extent of charge separation can be modulated by the length NR shell, and so it is the efficiency of hydrogen production. As a matter of fact, the hydrogen production of the CdSe@CdS/Pt was almost 30 times higher than CdS/Pt for the same length of CdS with a maximum AQE of 20% at 450 nm reached for 60 nm CdS rod with a CdSe core of 2.3 nm. Wu et al. have evaluated the performance of MUA-capped CdS/Pt and CdSe@CdS/Pt NRs in aqueous solutions using methanol or sulfite as hole scavenger [71]. The relative behaviour of these two NR-based systems varied with the nature of the hole scavenger: higher QE (2% versus 0.8%) was observed for CdSe@CdS/Pt when methanol was used; instead, CdS/Pt was more efficient than CdSe@CdS/Pt (10% vs 3%) in the presence of sulfite. Utilizing ultrafast transient
absorption spectroscopy the authors have measured the transfer rate of the electrons to the Pt domain and of the holes to the surface. They found that the electron transfer efficiency to the Pt tip is near unity for both systems. Instead, the hole transfer rate was slower in CdSe@CdS due to the hole confinement within the CdSe core that is harder to access by the electron donor. The hole transfer rates likely depend also on the interaction of the electron donor with the surface ligands and the accessibility of surface sites. The authors found a positive correlation of the observed hole transfer rates to the scavenger with the steady-state H2 generation QE, indicating that hole transfer was a key efficiency-limiting step [71]. Despite the importance of hole transfer in determining both efficiency and stability, electron transfer studies outnumber hole studies. Different groups are investigating this aspect into more details for the CdSe@CdS system [72,73]. Examples are given below in section 5.1.

As an alternative to Pt as HER catalyst, recent works have studied the effect of different metallic and bimetallic co-catalysts on the CdSe@CdS rods tip to improve the efficiency for the photocatalytic water splitting reduction half reaction [60,74,75]. Nakibli et al. have compared the effect of Au, Pt, Au@Pt and Au-Pt deposited simultaneously [74]. Rods decorated with Au cocatalyst showed very low activity, compared with the Pt cocatalyst. When the Au tip was covered with Pt in a core@shell configuration, the performance were improved compared to pure Pt, with the best performance obtained when the Au-Pt were deposited simultaneously. The suggested reason relies on the fact that the injection of the photoinduced electrons from CdSe into the Au tip is faster than their injection into the Pt tip. At the same time, only Pt can completely extract the excited electrons from the semiconductor NRs [76]. Thus, building a golden gate through which electrons can migrate from the semiconductor photocatalyst to the Pt cocatalyst islands results in increased activity for the water reduction half reaction. The same authors demonstrated that the synergistic effects of Au and Pd are beneficial for both the overall stability and the efficiency of the photocatalysts [77]. High-resolution TEM techniques were used to unravel the effects of structural changes at the atomic scale on the overall function and durability of heterostructured photocatalysts. The authors showed that the mobility of Pd atoms has a detrimental effect on the stability of the nanostructures due to atomic scale corrugation processes and that the Au serves as a physical barrier against Pd migration [77]. More recently, the Amirav group has published a study where the Pt catalyst is substituted with Ni domains. Similar AQE (23%) were achieved respect to that for Pt (27%) [74]. Ehrat et al. have also decorated core@shell CdSe@CdS with Ni [78]. They have shown that the hydrogen generation rate increases with the shell thickness, but decreases with the increasing core size. Interestingly, they have found that the photoluminescence quantum yield and lifetime follow the same trend. The correlation between the photocatalytic performance and photoluminescence lifetime is to be expected because they both rely on efficient charge separation [78].

Very recently, the record AQE of 100% for CdSe@CdS core@shell NRs has been achieved by Kalisman et al. when the system was tested at very basic pH illuminated with a LED at 455nm [79]. This result was attributed to the presence the hydroxyl anion-radical redox couple operating as a shuttle to relay the holes, in a similar manner to the work by Simon et al. cited above [69]. The authors also demonstrated 44h stability under non-continuous illumination. This work highlights once more the importance of efficiently extracting the holes to boost efficiencies.
Figure 4. Semiconductor/metal hybrid NCs for HER. TEM images of (A) metal tipped CdSe nanorods [81], (B) Pt tipped CdSe@CdS nanorods [79], (C) Pt tipped ZnSe@CdS nanorods [80], respectively. (D) Schematic representation of the relative band position for the corresponding HNCs in figure A-C illustrating the hole and electron path after illumination toward the scavenger (A) or catalyst (Pt), respectively. (E) Schematic plot of the energy levels of the electrons of \( \text{SO}_2^{-3}, \text{TEA}, \text{EDTA}^{4-}, \) and MeOH that are involved in the reduction of the photohole vs. vacuum and on normal hydrogen electrode (NHE) scale [81]. (F) Scheme of the photocatalytic water reduction by CdS-Ni nanorods (cysteine-stabilized) at alkaline conditions. The photo-excited holes oxidize hydroxyl anions that, as a radical, carry away the positive charges and in turn oxidize ethanol to acetaldehyde. The blue arrows denote movement of the species; the red arrows denote a redox reaction [69]. Reproduced from [79,80], Copyright 2016,2011 American Chemical Society. Reprinted from [81], with the permission of AIP Publishing. Reprinted by
Other Cd-based semiconductor NCs.
To tune the localization of photoinduced holes Acharya et al have synthesized asymmetric core@shell HNCs including ZnSe@CdS/Pt and ZnTe@CdS/Pt (figures 4C,D) [80]. In this study the authors evidenced the role of the NC ligands in mediating the hole extraction from the NC surface. The comparison of photocatalytic activity in ZnSe@CdS/Pt and ZnTe@CdS/Pt revealed that the hydrogen production on the Pt site was efficient only when electron donating molecules were used as surface ligands. Based on band gap alignment, the hydrophilic MPA ligands are expected to scavenge holes from ZnSe but not ZnTe core NCs. This hypothesis was confirmed by the authors from fluorescence lifetime measurements. Furthermore, the addition of excess ligand molecules to the NC solution resulted in a recovered H$_2$ generation rates, in agreement with the hypothesized role for the MPA and with the fact that the photo-holes need to be extracted to the NC surface for the H$_2$ to be produced.

Cu$_{1.94}$S–Zn$_x$Cd$_{1-x}$S heteronanorods (0 ≤ x ≤ 1) have recently been proposed to achieve a tunable composition-dependent band gap (between 2.57–3.88 eV) while also favoring charge separation [82]. Even without any cocatalysts, Cu$_{1.94}$S–Zn$_{0.23}$Cd$_{0.77}$S heteronanorods exhibited an AQE of 8.5% under visible-light irradiation (λ > 420 nm), representing a 59-fold enhancement compared with the pristine CdS catalyst. Meanwhile, deposition of a Pt cocatalyst on the Cu$_{1.94}$S–Zn$_x$Cd$_{1-x}$S surface substantially enhanced the hydrogen production performance by achieving an AQE of 26.4% at 420 nm [82].

As one example of photocathodes assembled from QDs, Korala et al. have recently reported a new method to fabricate high quality CdSe and CdSe@ZnS QDs photoelectrodes by applying gelation strategies to spin-coated thioglycolic acid-capped QD films (see section 5.1 and figure 9A) [52]. This method enables improved charge transport properties in QD solids by enhancing interparticle coupling via formation of oxidation-induced dichalcogenide linkages. Overcoating CdSe QDs with a thin ZnS shell reduces unpassivated surface states, and thus charge trapping, and results in a more than 2-fold increase of photocurrent (200µA/cm$^2$ versus 75 µA/cm$^2$ under white light irradiation and at -0.57V vs Ag/AgCl).

Cd-free semiconductor NCs.
In the search of alternative light absorber to Cd-based materials, Cu$_2$ZnSnS$_4$ (CZTS) combines ideal band gap (direct and 1.5 eV) and a composition based on Earth abundant elements. Yu et al. have constructed HNCs including highly monodisperse quasi-spherical CZTS as light absorbers and Au and Pt nanoparticles as co-catalysts using the seeded growth approach, with Pt being more active [83]. The best Pt loading in order to maximize the H$_2$ production rate without compromising the light absorption of the CZTS NCs was determined. Furthermore, the compositional tunability allowed the authors to report the H$_2$ production trend versus the composition of the CZTS and observed higher activity in the Cu-poor and Zn-rich samples, in agreement with the behavior observed in photovoltaics [83]. An increased activity was observed when the NCs were functionalized with MPA instead of (NH$_4$)$_2$S. While this behavior is not commented, it highlights once more the importance of controlling interfaces and band alignments in the photocatalytic systems. The same group has also investigated CuIn$_{1-x}$Ga$_x$S$_2$ (CIGS) NCs
to identify the best composition to compromise between visible light absorption and efficient charge transfer [84]. In both CIGS and CIGSe, the incorporation of Ga has a major impact on the position of the conduction band minimum, which is shifted to higher energy values; instead, the valence band maximum remains almost unchanged. The additional energy of the electrons at the conduction band minimum when increasing the Ga content, provide extra driving force for charge injection, and thus a faster charge transfer. At the same time though, the band gap increases and so less visible light is absorbed. The optimum composition was found to be CuIn$_{0.3}$Ga$_{0.7}$S$_2$, which was characterized by a band gap of 1.75eV. A two-fold increase in the hydrogen production rate was observed when functionalizing the CIGS NCs with Pt.

Among the Cd-free alternative photoanodes for HER, considerable attention has been paid to CuIn$_2$S$_2$ (CIS) and its alloys; among them an often-studied CIS-based alloy is ZnS-CuInS$_2$ (ZCIS) semiconductor [85]. CIS and ZnS are both direct-gap semiconductors with bandgap energies of 1.5 and 3.7 eV, respectively. By varying the ratio of CIS to ZnS, the bandgap of ZCIS can be easily tuned to absorb light in the visible region, which makes up a large part of the solar radiation spectrum. Only recently Ye et al. have employed colloidal synthesis to obtain ZCIS nanorods alloy successively decorated with Pt and Pd$_4$S [86]. The authors demonstrated the potential utility of these ZCIS NRs in visible light-driven photocatalytic hydrogen generation from water in the presence of S$^2$- and SO$_3$$^2$- hole scavengers. The ZCIS-Pt and ZCIS-Pd$_4$S displayed higher photocatalytic performance than the bare ZCIS [86].

Recently Patra et al. have reported the synthesis of 0D and 2D twin dot-disc Au-CIS HNCs (figure 5) [51]. As these structures have plasmon and exciton coupling, they were explored as photocathode in a PEC cell for water splitting and recorded significant enhancement of the photocurrent in comparison to only CIS nanostructures. The half-cell STH evolution efficiency of Au-CIS photocathode was found to be higher (4.29% at 0.438 V vs. RHE) than other reported comparable nanostructures [51].
3.2 Colloidal nanocrystals as light absorbers for OER

The studies on HER using colloidal NCs largely outnumber the ones on OER. Only a couple of attempts have been made to synthesize colloidal II-VI semiconductor/metal HNCs for OER. However, no oxygen evolution has been demonstrated so far [87]. As discussed in the previous session, the main challenge is the instability of these materials due to photo-oxidation. The latter becomes even more problematic under OER conditions, as the reaction is slower (four holes need to be transferred to liberate one oxygen molecule) and higher overpotentials are usually required. Because of their higher stability in oxidative environment, metal oxide NCs are ideal candidate as light absorbers for OER. A few studies have been reported where photoanode electrodes are assembled from metal oxide NCs. The problem here is that there are not too many stable systems available yet that efficiently absorb visible light. As the previous one, this section is organized by material classes.

**II-VI semiconductor NCs.**

Kalisman et al. have functionalized CdSe@CdS asymmetric dot@rod with IrO$_2$ nanoparticles [87]. While the authors highlight photochemical stability under prolonged illumination, no demonstration of possible utilization of the photogenerated holes is
reported. More recently a study from the same group details the synthesis of complex hybrid NCs comprising both a water oxidation and water reduction catalyst, namely CdSe@CdS/Pt/Ru@RuOx [88]. This is a great example of the impressive control gained by the nanoscience community over the assembly of different building blocks in a single object. However, also this work supplies no evidence of actual activity towards water oxidation and/or reduction.

**Metal Oxide NCs.**

In most of the studies involving colloidal metal oxide NCs, photoanode electrodes are prepared by necking the NCs at a certain temperature. Such a thermal treatment might or might not induce phase transformation and/or grain growth depending on materials and treatment conditions. As illustrated in section 2.1, the facets exposed on the NC surface can be finely tailored by tuning the reaction conditions. One of the advantage of using colloidal NCs as precursors for thin film electrodes is the possibility of directing grain size and orientation, which are known to impact transport properties and catalytic reactivity, through the size and shape of the NCs themselves. Such a tailoring of the microstructure has been demonstrated in materials for photovoltaics and has often lead to enhanced performance [89].

The popular binary metal oxide photoanodes researched upon, like TiO$_2$, WO$_3$, and α-Fe$_2$O$_3$, are readily available by colloidal synthesis. Recently, our group has been able to position nitrogen dopants selectively in interstitial or substitutional sites in TiO$_2$ branched NCs [90]. Correlating the change of band gap and the photoelectrochemical behavior of the doped TiO$_2$ nanostructures to the nature of the dopants was thus possible. When assembled as photoanodes, the substitutionally doped samples outperformed the interstitially doped ones. While N-doped TiO$_2$ is clearly not a technologically relevant photoanode material, this work demonstrates the potential of colloidal NCs as atomically tunable materials to build unambiguous structure/properties relations, often prevented by a poor angstrom-scale material control.

WO$_3$ photoanode electrodes assembled from nanowires by Goncalves et al or from nanoplates by Hilaire et al have showed preferential texturing along the {200} and the {002} planes, respectively [91,92]. These photoanodes exhibited a photocurrent of 1.96 mA/cm$^2$ at 1.23V$_{RHE}$ and 2.7 mA/cm$^2$ at 1V$_{RHE}$, respectively, both with an onset potential around 0.5 V$_{RHE}$. Because of different electrolyte concentration, these results are not directly comparable, yet they serve the purpose of illustrating the impact of texturing on the photocurrents by suggesting a better charge transport along the [001] direction. Goncalves et al have prepared hematite α-Fe$_2$O$_3$ photoanodes by annealing at 820°C thin films of highly monodisperse magnetite nanocrystals with 2.5 nm in diameter [93]. Among the crystalline phases of iron oxide, hematite is the active phase to drive water oxidation. The growth of columnar hematite grains along the [001] direction with grain size around 50 nm was observed. Two main factors were identified to contribute to the lower overpotential and high photocurrent (for 930nm thick films 1.1 mA/cm$^2$ at 1.23V vs RHE and overpotential smaller than 0.8V) recorded for these photoanodes: 1) favorable surface properties of the hematite grains obtained by using a high temperature and an oxygen atmosphere during the sintering process; 2) favorable morphology, where columnar grains are oriented in a favorable crystallographic orientation for electron collection and the mesoporous structure. The same authors achieved better performance (photocurrent up to 1.4 mA/cm$^2$ and 2.7 mA/cm$^2$ at 1.23 V$_{RHE}$ for undoped and Sn-doped
hematite films, respectively) when assembling the colloidal magnetite NCs under an external magnetic field during dip coating deposition [94]. The process is illustrated in figure 6A. Such a process promoted the growth of hematite columnar grains of 180 nm thickness and with a preferred [110] orientation. The study by Wang et al on Sn-doped Fe₂O₃ corroborates an increased electron transport properties along the same crystallographic direction. The authors reported Sn-doped Fe₂O₃ films with a preferred [110] texture, obtained by sintering in Argon Sn-doped magnetite NCs at 550°C [95]. A photocurrent up to 3.32 mA/cm² at 1.23 V vs RHE and onset potential around 0.9V vs RHE was measured for film thickness of 385 nm. Samples annealed in inert atmospheres performed much better than the ones annealed in air and doped samples better than undoped ones. Sintering in reducing atmosphere promotes the creation of oxygen vacancies which can play a key role in increasing conductivity (which was verified by the authors of this work) as well as modifying the photoactivity towards water oxidation.

In the previous examples, hematite was obtained upon annealing of magnetite NCs. Wang et al. succeeded in synthesizing α-Fe₂O₃ NCs by microwave-assisted colloidal synthesis [96]. Because of the good solubility, the photocatalytic activity was tested in aqueous solution for oxygen evolution. The same particles were deposited by electrophoresis, annealed and tested as thin films electrodes as well. Comparison with other studies is challenging as only oxygen generation rate was reported [96]. However, this work shows the versatility afforded by colloidal chemistry: light absorbing nanocrystals can be tested both as photocatalysts in solution or assembled in photoanode for electrochemical investigation.

In the last years, monoclinic BiVO₄ has attracted much attention as a photoanode for solar fuel applications [97–99]. The best performance for this materials have been obtained by nanostructuring and by interfacing with wider band gap oxide semiconductors [99]. Our group has recently developed a BiVO₄ NC ink that facilitates the assembly of BiVO₄/metal oxide (MO) (MO = TiO₂, WO₃, and Al₂O₃) nanocomposites in which the morphology of the metal oxide building blocks is finely tailored [16]. The BiVO₄ NC ink was constituted by Bi₂O₂.₇/VOₓ heterodimers (figure 6B). Tunable nanostructured BiVO₄/MO were obtained by annealing thin films deposited from mixed solution of the NC heterodimers with TiO₂, Al₂O₃ and WO₃ NC of different size and shape. We used a combination of transient absorption spectroscopy – spanning from picoseconds to second timescales – and photoelectrochemical measurements to reveal that the achieved structural tunability is key to understanding and directing charge separation, transport, and efficiency in these complex oxide heterostructured films. For example, in the BiVO₄/WO₃ nanocomposite, better photoelectrochemical performance was found for smaller WO₃ NRs (figure 6B). This behavior resulted from an improved efficiency of electron injection from BiVO₄ to WO₃ deriving from a more extended interfacial area between BiVO₄ and WO₃, compared to nanocomposites comprising bigger WO₃ nanorods[16]. Our results highlight the significant potential afforded by the assembly of such oxide nanocomposites and unveil relationships between MO properties and PEC performance.

We have also investigated a NC-seed mediated growth of thin films vanadate light absorbers. The seeding process has been traditionally employed to reduce the crystallization temperature in ceramic ferroelectric materials, such a lead zirconate
titanate, to facilitate their integration in flexible electronics as a result of the decreased crystallization temperature [100,101]. Recently, our group has utilized colloidal nanocrystals with variable composition as seeds for the nucleation and growth of vanadate thin films [102]. Specifically, we employed Bi$_x$Sb$_{1-x}$ colloidal NCs with variable composition $x$ to discover a new light absorber, BiV$_{1-x}$Sb$_x$O$_4$. The Bi$_{1-x}$Sb$_x$ NCs were converted into Sb-BiVO$_4$ films by reacting them with vanadyl acetylacetonate upon annealing (figure 6C). This novel two-steps synthetic approach allowed for a precise compositional tuning while preserving the same morphology and crystalline structure up to a 20% antimony content. It is important to note that attempts to synthesize Sb-BiVO$_4$ by more conventional approaches such as sol-gel, solid solution, and spray pyrolysis were unsuccessful and resulted in phase separation, even for low percentage Sb alloys. Thanks to the broad compositional range covered by our approach, a combination of theoretical predication and experimental validation has showed that this photoanode possesses a band gap that linearly decreases with increasing Sb content up to 20% when the band gap is 0.2eV smaller than the one of the undoped counterpart (figure 6C). While this material is still far from being optimal, the results of this work, derived from close coupling of theoretical understanding and advanced synthesis approaches, provide a path to the design of other metal oxides alloys with desirable band gaps and band edge positions for solar water splitting. To demonstrate the general applicability of the NC-seeded growth approach, other high surface area ternary oxide light absorbers were targeted. Bi$_2$WO$_6$, CuWO$_4$, SnWO$_6$, and InVO$_4$ were synthetized by using Bi, Cu, Sn and In$_2$O$_3$ NC seeds, respectively.
Figure 6. Photoanodes from assembly of metal oxide NCs. (A) Proposed pathway to produce hematite thin-film starting from colloidal magnetic NCs and scanning electron microscopy (SEM) cross section image of the obtained hematite thin film [93]. (B) Low-resolution TEM images of as-synthesized Bi$_2$O$_{2.7}$/VO$_x$ NC heterodimers (left) and SEM images on the surface and in cross-section (inset) of the BiVO$_4$/TiO$_2$ nanocomposite.
4. Colloidal nanocrystals as electrocatalysts.

Studies on colloidal NCs as electrocatalysts for HER and OER have proliferated in the last few years and have highlighted the great potential of colloidal NCs both as a platform to deepen the fundamental understanding of structure/properties relations in catalysis and as an actual technological opportunity. The fine tailoring of size, shape and composition serves well the purpose of fundamental studies. The easy processability of colloidal solutions (colloidal NCs can be deposited on flat or porous or temperature sensitive substrates and loading can be easily adjusted) is well suited for integration in different device configurations. Much of the recent work on HER and OER, which is reviewed in this section, focuses on non-noble metal catalysts that are stable in extreme pH conditions, which are relevant for electrolysis and for integrated photoelectrochemical devices [103,104].

4.1 Colloidal nanocrystals as electrocatalyst for HER

Platinum is so far the best catalyst for HER, with the lowest overpotential of -18 mV at -10 mA/cm² and -26 mV at -20 mA/cm² [103]. Examples of colloidal semiconductor/Pt HNCs for HER have been discussed in section 3.1. Recently the Schaak group has been leading the field by investigating the electrochemical activity of metal phosphide colloidal NCs towards hydrogen evolution and the stability in acidic conditions, under which proton exchange membrane-based electrolysis is operational [105]. An overview of this work is given in figure 7. Before performing any electrocatalytic measurement on these NCs, ligand removal was performed by annealing 30 minutes at 450°C in 5% H₂/N₂ without observing any apparent change in the morphology /composition of the particles used in their studies.

Nickel phosphide (Ni₃P) NCs were synthesized as hollow multifaceted particles with an average size around 17nm and high density (001) facets exposed, which has previously been predicted to be the most active surfaces [106]. The overpotential for these particles to produce current densities of -20 and -100 mA/cm² were -130 mV and -180 mV, respectively. The size and shape-dependence of electrocatalytic activity of the Ni₃P NCs has also been investigated [107]. Electrochemical measurements have revealed that Ni₃P nanospheres (Ni₃P NSs) with predominant (001) surfaces exhibit higher HER activity than Ni₃P nanorods (Ni₃P NRs) with the (210) surface. Ni₃P NSs generated a current density of −10 mA cm⁻² at an overpotential of 135 mV in 0.5 M H₂SO₄, whereas Ni₃P NRs produced the same current density at a larger overpotential (270 mV). Furthermore, the turnover frequency of Ni₃P NSs was about thirteen times higher than that of Ni₃P
NRs. The results suggest that the crystallographic facets of Ni$_2$P NCs play a critical role in dictating HER activities [107]. Zhou et al. have also investigated the effect of the size of Ni$_2$P NCs for HER to establish the relationship between size and catalytic activity [108]. Highly monodisperse, small-sized Ni$_2$P NCs with sizes ranging from 2 nm to 10 nm were synthesized. The resulting small-sized product possesses a high accessible surface area and a high density of exposed (001) facets. Experimental results showed that the small-sized Ni$_2$P NCs not only reduced the overpotential but also decreased the Tafel slope, thus significantly improving HER catalytic performance. The best performance were achieved with 5.4nm Ni$_2$P NCs which showed an overpotential of 93 mV at -20 mA/cm$^2$ and for a mass loading of approximately 1 mg/cm$^2$ onto a 0.20 cm$^2$ Ti foil substrate [108].

Multi-faceted cobalt phosphide (CoP) NCs were synthesized by the Schaak group by reacting Co NCs with trioctylphosphine [109]. Electrodes comprised of CoP NCs on a Ti support (mass loading equal to 2 mg/cm$^2$) produced a cathodic current density of 20 mA/cm$^2$ at an overpotential of -85 mV. The CoP/Ti electrodes were stable over 24 h of sustained hydrogen production in 0.50M H$_2$SO$_4$ [109]. The same authors used the tunability afforded by colloidal chemistry to directly compare the activity of CoP and Co$_2$P NC catalysts [110]. Co$_2$P has not been studied as extensively as CoP because of its higher overpotentials. However, unambiguous conclusions from comparing catalysts synthesized by different approaches and tested under different conditions cannot be made. Nevertheless, such comparisons are needed to identify and to understand the key intrinsic factors that underpin high catalytic activity. Morphologically equivalent CoP and Co$_2$P multifaceted and hollow particles were accessed using the same colloidal procedure. This allows similarities and differences in HER activity to be correlated most significantly with composition and structure, while minimizing morphological contributions, such as size, size distribution, dispersability, and faceting. When testing the electrocatalytic activity of CoP and Co$_2$P NCs, the following activity trend was found Co$<$Co$_2$P$<$CoP. While this result is in agreement with previous literature, much lower overpotential (-95 mV and -109 mV to produce cathodic current densities of -10 and -20 mA/cm$^2$, respectively) was found for the Co$_2$P NCs compared to the ones with different morphology reported in the literature (that falls in a range between -160 mV and -200 mV at -20 mA/cm$^2$), once again it reveals the importance of morphology in catalysis. Furthermore, direct comparisons by the same group of multifaceted CoP nanoparticles with highly branched CoP nanostructures that exposed a high density of (111) facets suggested that the high HER activity of CoP is intrinsic to the system and that, for this system, shape may not play a significant role in defining the magnitude of the overpotentials required to produce operationally relevant cathodic current densities [111]. Ha et al. have investigated the active site and stability of the same colloidal synthesized CoP NCs for HER through experiments and DFT calculations [112]. The HER activity of CoP is found to be reduced with potentiostatic holding at potentials above 0.4V vs. RHE. This activity loss was accompanied by leaching of P and lowered Co/P ratio suggested by EDX results. The formation of (oxy)phosphate(s) emerges concomitantly with activity reduction as indicated by signal changes of Co and P in X-Ray absorption spectroscopies. Furthermore, DFT calculations indicate surface P could act as the active site for HER on CoP, which constrasts with the traditional HER mechanisms on the metal site [112]. Iron phosphide (FeP) NCs were synthesized and tested by the Schaak’s group [113].
These particles possess a similar morphology (hollow 10nm spheres) compared to the nickel and cobalt phosphide, though they outperformed those. In fact, they showed an overpotential of -50 mV under the same acidic conditions the other transition metal phosphides had been tested, and -102 mV in a neutral electrolyte, at a current density of -10 mA/cm². In addition to serving as an efficient HER electrocatalyst, FeP nanoparticles were immobilized on TiO₂ particles (Degussa P25) to photocatalytically generate hydrogen under UV illumination in acidic and neutral-pH aqueous solutions.

Amorphous tungsten phosphide (WP) nanoparticles have been synthesized and demonstrated to be highly active HER electrocatalysts in acidic solutions, requiring overpotentials as low as -120 mV and -140 mV at current densities of -10 mA/cm² and -20 mA/cm², respectively [114].

Finally, the Schaak group has also demonstrated that colloidal molybdenum phosphide (MoP) amorphous NCs with diameter around 4nm are active and acid-stable electrocatalysts for HER [115]. Overpotential as low as -90 and -105 mV were requested to produce operationally relevant current densities of -10 and -20 mA/cm², respectively, thus revealing that these particles are much more active than bulkier MoP nanostructured electrode ranging between -125 and -150mV at -10 mA/cm².

Figure 7. Metal phosphide NCs as electrocatalysts for HER: (A) High resolution (HR)-TEM image of Ni₃P NCs, highlighting the exposed Ni₃P (001) facet and the 5.2 Å lattice fringes that correspond to (010) planes [106]. (B) Low and HR-TEM image of FeP NCs [113]. (C) Low and HR-TEM image of Co₂P NCs [110]. (D) Plot of current data vs. potential in 0.5M H₂SO₄ for Ni₃P/Ti electrodes [106]. (E) Plot of current data vs. potential for three distinct FeP/Ti electrodes along with Pt and Ti in (acidic) 0.5M H₂SO₄ as well as in (neutral) 1.0 M Phosphate buffered saline (PBS) [113]. (F) Plot of current data vs. potential in 0.5M H₂SO₄ for Co₂P/Ti and CoP/Ti electrodes [110]. Reprinted with permission from [106,110,113], copyright 2013, 2015, 2014 American Chemical Society.

All the previous catalysts have been tested in acidic conditions. However, finding HER catalysts which are active and stable in alkaline solution is highly sought after as well.
Recently, the Schaak group has reported the colloidal synthesis of molybdenum rich Co-Mo alloy NCs with diameters of approx. 3 nm and they have proved their potential to serve as an Earth abundant alternative to Pt for catalyzing the HER in alkaline aqueous solutions [116]. With an observed overpotential of -75 mV at an operationally relevant current density of -10 mA/cm², the performance of the Co-Mo/Ti electrode places it amongst the best non-noble metal HER catalysts in alkaline solutions. When subjected to both short-term galvanostatic stability testing and longer-term simulated stability testing via cyclic voltammetry, the Co–Mo/Ti electrode was found to be stable in 1M KOH [116].

While none of the catalysts reviewed above outperform Pt, these studies demonstrate that phosphide NCs still represent valid non-noble metal alternative electrocatalyst.

4.2 Colloidal nanocrystals as electrocatalyst for OER

As referred to in section 3.2, OER is more challenging than HER because of the slower kinetics of the 4e⁻/4H⁺ coupled transfer needed to liberate one oxygen molecule. The search towards robust Earth abundant electrocatalysts is frenetic but simultaneously extremely challenging. One noteworthy observation is that iridium and ruthenium are the only OER catalysts active and stable in acidic electrolytes. Surprisingly not many studies on ruthenium and iridium oxide colloidal NCs, prepared in organic solvents, to investigate size and shape dependence have been reported. Under alkaline conditions, a few more options exist, with NiFe-based catalysts being the most investigated so far. To the best of our knowledge only a handful of studies have been reported of OER catalysts prepared by a colloidal approach.

Work from the group led by Bein and Fattakhova-Rohlfing focuses on the synthesis and characterization of NiO and Fe-doped NiO NCs (figure 8) [117]. Those were prepared by solvo-thermal synthesis in tert-butanol and characterized by a size below 5nm with a slight decrease in size as the iron content increases up to 20%. It is interesting to note that, unlike the previously reported ultrathin NiO films, NiO NCs do not completely transform into layered Ni(OH)₂ after prolonged cycling. This behavior is not discussed further by the authors but it is a clear indication that that catalytic mechanism might be different in nanoscale catalysts. While there are no big changes in overpotential compared to bulk, the beneficial effect of the reduced crystal size of undoped NiO to only a few nanometers and the surface properties of the nanoparticles on the catalytic performance is shown by the very high turnover frequency value (0.55s⁻¹), indicating an increased amount of electrocatalytically active sites. As for the Fe-doped NiO, while different studies report a solubility limit up to 5-6% before phase separation occurs, the authors did not observe any formation of additional phases up to 20% according to the results of several characterization methods. The unusually high solubility of Fe in the rock salt NiO structure can originate from the kinetic control of the Fe-NiO phase formation in a solvothermal process, as well as the effect of the nanoscale where the metastable and defect phases often show higher stability than in the bulk. The NC dispersion was deposited as thin films on gold electrodes of the piezoelectric quartz crystals for exact determination of their loading. When doping the nanocrystals with iron, the catalytic activity of the NiO was even further increased with turnover frequency up to 1.9 s⁻¹ for 10% Fe nanoparticles in the present study. In addition to the extremely high electrocatalytic activity, the excellent dispersibility of the FeₓNi₁₋ₓO nanoparticles offers
additional advantages, such as a great flexibility towards the fabrication of different electrode architectures.

Bau et al. have synthesized and tested as OER electrocatalysts 8 nm Ni-Fe-O NCs in a metastable rock-salt phase and variable composition ratios of nickel to iron [118,119]. The nickel-rich NCs showed the lowest overpotential of around 300 mV at 10mA/cm² with optimal electrode thickness corresponding to a couple of monolayers. The similar OER performance of the rock salt phase NCs with other Ni-Fe-O based catalysts suggests that the presence of Fe³⁺ centers on the surface plays a bigger role than the crystalline phase itself.

Figure 8. Fe-doped NiO NCs as OER electrocatalysts. (A-C) TEM, HRTEM and electron diffraction pattern of Fe₀.₁Ni₀.₉O NCs; (D) Cyclic voltammograms of gold electrodes coated with thin films of the NCs with different Fe content. The inset shows across-section HR-TEM image of the NP-10% film deposited by spin-coating on the substrate. (E) TOF values at an overpotential of η = 300 mV (filled circles) and η = 350 mV (open circles) for different Fe contents [117]. Reprinted with permission from [117], copyright 2015 American Chemical Society.
In addition to metal oxides, metal phosphides have recently emerged as potential electrocatalyst candidates for OER. Masud et al. have tested colloidal 5nm FeP NCs and measured an overpotential of 290 mV at a current density of 10 mA/cm² [120]. The electrode was reported stable with full activity for 4 hours. In addition, a hybrid electrode assembled from the same NCs and reduced Graphene oxide (rGO) lowered the overpotential to 260 mV at 10 mA/cm², making it the lowest reported at the time for metal phosphides. The enhanced electroactivity in the nanocomposites was attributed to the increase in the catalytically active surface area. Dutta et al have studied 1D needle shaped Co₂P [121]. These catalysts generated 10 mA/cm² at an overpotential of 310 mV and were stable for 10 hours with 15 mV increase in overpotential. The salient feature of this colloidal synthesis is the needle shaped Co₂P, which has higher metallic content compared to CoP and thereby a most likely better charge transfer. Another interesting result discussed by the authors is the analysis of the electrode after catalysis. A combination of EDX and XPS evidences the presence of oxide or oxo-hydroxide layer around the Co₂P core. The authors suggest that the Co₂P/oxide interface enhances the carrier transportation from the core to the oxide layer, thereby enhancing the electroactivity. NiP nanowires were prepared by colloidal synthesis and reported to exhibit current density of 10 mA/cm² at an overpotential of 330 mV for OER [122]. As for cobalt phosphide, the authors report the formation of a NiOₓ shell during operation which enhances the activity of the electrode for OER and is termed as a synergistic effect which needs further study. The insertion of a second metal has been suggested to lower the thermodynamic barrier of a proton-coupled electron transfer pre-equilibrium while facilitating O−O bond formation, leading to enhanced catalytic activity [123]. Hence CoMnP NCs have been synthesized by colloidal synthesis [123]. An overpotential of 330 mV at 10 mA/cm² was reported, which was lower than the overpotential of CoMnO₂ (390 mV) synthesized by the same approach. The higher intrinsic conductivity of phosphides in comparison to oxide has been hypothesized to account for this behaviour. An increase in overpotential to 370 mV on 500 cyclic voltammetry sweeps was observed and attributed to surface oxidation and leaching of P into the electrolyte. From these initial studies on phosphides as OER catalysts, it is clear that transformation and oxidation of the catalysts occur during operation.

Overall, the studies reviewed in this section also highlights that the minimum overpotential reached for OER with any non-noble metal catalyst tested so far is around 300 mV at a current density of 10mA/cm². Carefully in-situ and in-operando studies are mandatory to get true insights into the activity and reaction mechanisms, so to guide the development of new and more efficient OER catalysts.

5. Outlook
5.1 Ligands off or on?
As described in section 2, the organic ligands functionalizing the NC surface are essential during the synthesis to modulate NC size and shape and, afterwards, to grant solubility to the NCs in a wide range of solvents. However, they are not necessarily beneficial for their final application [124,125]. In general ligands are hydrophobic and highly insulating, and constitute a significant barrier for charge transfer and transport. Thus, whether the photocatalytic, photoelectrocatalytic or electrocatalytic activity of the NCs is tested in thin films or in solution, proper surface chemistry is needed to investigate their
behavior with minimal interference of the native coordinating ligands. Fortunately, the understanding and the progress made in NC surface chemistry are far enough that this does not constitute any obstacle for the use of colloidal NCs as model catalytic systems for different reactions [26–30]. The goal is to eliminate/exchange the bulky organic ligands while preserving size, shape and composition of the NCs. Different options are available, with the best one being highly dependent on the system of interest and on whether the NCs will be tested as thin film electrodes or dispersed in solution.

When the NCs are assembled into photoelectrodes or electrodes, thermal treatment to decompose the ligands is one of the most straightforward options. The temperature required for the decomposition of long-chain ligands usually ranges between 400°C and 500°C. Annealing in air is required to avoid carbon contamination from decomposed ligands, which is a problem for light absorption in the case for photoelectrodes. Clearly, this treatment is limited to systems that are resistant to oxidation. We have found such approach to work well the BiVO₄ NC ink and its nanocomposites (figure 6B) [16]. Nickel iron oxide NCs for OER were also annealed in air (figure 8) [117,119]. The Schaak group uses annealing in a tube furnace at a temperature between 350°C and 450°C under H₂(5%)/Ar(95%) to remove surfactant molecules from the surface of their phosphide NCs (figure 6) [106,111,114–116]. For certain NC compositions, at these relatively high temperatures, sintering might occur, thus invalidating the synthetic efforts to control size and shape. One possibility to overcome this issue is to exchange of the native ligands with shorter chain ligands. This approach brings two advantages: i) decomposition temperature is reduced to <200°C; ii) annealing under inert atmosphere with minimal carbon residue remaining is possible. Formic acid has worked well for assembling conductive electrodes from tin-doped indium oxide NCs [126]. Patra et al have used a similar approach when building photoelectrodes from CIS-Au hybrid NCs (figure 4) [48]. Substitution of oleylamine with 1,2-ethane dithiol decreased the annealing temperature to 200°C and allowed the ligand removal to be conducted in vacuum instead of air [51]. Alternative exchange approaches have been developed during the last years and include a variety of agents that react with the native NC ligands and leave behind a “bare” NCs surface (“reactive stripping”). The nitrosyl tetrafluoroborate and the exceptionally mild Meerwein’s salt are two example among these [127,128]. We have found the latter to be suitable for N-TiO₂ as it allowed annealing under inert atmosphere, which was essential to preserve the nitrogen doping [90]. To build photoelectrodes from CdSe and CdSe@ZnS core@shell, Korala et al created an interconnected NC network by interparticle linking via dichalcogenide bonding (figure 9A) [52]. Such a process was achieved by exchanging the original trioctylphosphine/stearic acid-capped NCs with thiolated ligands (thioglycolic acid) and inducing gelation afterwards by immersing into an oxidizing solution. This last step was followed by annealing at 250°C under argon to form the inorganic sulfur bond between the NCs and an electrically conductive films where size, shape and optical properties of the original NCs were preserved. Fully inorganic ligand, such as molecular metal chalcogenides and halide ligands, have been used to exchange the native ligands and imprint electrical conductivity to NC films [129,130]. Yet, these approaches have not been employed to build photoelectrodes for artificial photosynthesis. Very recently, rapid thermal annealing at 750°C for 30 seconds has been suggested as method to efficiently remove the ligands from Pd NCs supported on alumina or on ceria without causing any change in the size, shape and composition of
these NCs [27]. However, its applicability to other systems has to be proven yet. Oxygen plasma or UV treatments are other possibilities [119,131]. Bau et al have investigated these two different treatments for ligand removal from the surface of Ni-Fe-O NCs for OER: UV irradiation and air plasma (figure 9B) [119]. Interestingly, only the electrodes treated by UV irradiation were stable and preserved their appearance after electrolysis. The authors claim an increased adherence with the conductive substrate [119].

When the NCs are tested in solution, the need to preserve colloidal stability in water demands for the substitution of the native hydrophobic ligands with hydrophilic ligands. Some of the most common ligands that have been used for this purpose are mercaptoundecanoic acid (MUA), mercaptopropionic acid (MPA), cysteine, citric acid. It has been noted that these ligands themselves can act as holes or electrons scavengers depending on the alignment between the energetic level of the NCs and the ligand. One example is given by Acharya et al. and it is illustrated in figure 4C [80]. In a second example, Wang et al have studied the effect of three different ligands on the photocatalytic activity of H₂ generation for CdSe@CdS core@shell NCs [132]. They functionalized the NCs with three ligands containing thiol groups: poly-acrylic acid (PAA), 3-mercaptopropionic acid (3-MPA) and 2,3-dimercaptosuccinic acid (DMSA). The DMSA–capped QDs exhibited the highest efficiency of H₂ production, while the 3-MPA and PAA coatings resulted in the intermediate and lowest photocatalytic activity, respectively. A combination of photoluminescence intensity, lifetime measurements and electro transfer studies illustrated that such a ligand-dependent photocatalytic activity could be mainly attributed to the variations in the surface traps and corresponding charge separation efficiencies of the QDs induced by surface coating with the different ligands [132]. In one last example, Han et al have reported about a robust and highly active system for HER where the dihydropolic acid (DHLA) capping CdSe NCs plays a central role in transferring electrons to the catalysts, which is a soluble Ni²⁺-DHLA complex [48]. A schematic representation is shown in figure 9C. This system achieved QY of over 36% and 360 hours stability under illumination at 520 nm [48].

As discussed already in section 3.1, strategies to stabilize QDs need to be developed for these systems to be considered in a real-world context. Hole accepting ligands with tunable energetic levels can be utilized to conduct detailed charge transfer studies [72,133]. Strategies to improve the stability of quantum dots could emerge as a result of a more fundamental understanding of hole transfer. In a recent work, hole transfer kinetic studies have been conducted on CdSe@CdS core@shell NCs to hole acceptors covalently linked to the nanocrystals via a thiolate binding group [72,133]. The hole acceptor used is a ferrocene molecule that is positioned at the end of the thiol chain. The large thermodynamic driving force for photoinduced hole transfer allows this process to compete with native recombination. Yet, the hole transfer rate was still extremely sensitive to modulation in the electronic coupling achieved by varying the thickness and composition of the barrier material between the CdSe core and the acceptor. This barrier material includes both the length of aliphatic thiol chain and the thickness of the CdS shell. Through this study the authors demonstrate that one can use the design of QD heterostructures and acceptor ligands to mitigate undesirable trap-states while still being able to extract charges and preserve the photostability.

Recently, De Roo et al have introduced the concept of sustained chemically driven ligand displacement which allows to address the activity/stability tradeoff in nanocatalysis
Herein, weakly binding amines and alcohols chemically convert carboxylic acids functionalizing the surface of HfO$_2$ NCs into non-coordinate amides and esters. Thus, the carboxylic acids act both as stabilizer and as reagents and are supplied back during the catalytic cycles [125]. Eventually, similar concepts might also be applied to photocatalysts and catalysts for water splitting. Overall, these last examples point to the importance of ligand engineering in catalysis, which has been recently gaining increasing attention [134,135]. The ligands could indeed play the role of co-catalysts and synergistically contribute to increase the activity or the stability of the catalysts, instead of being always demonized.

**Figure 9.** Various strategies for ligand exchange or removal. (A) Schematic representation of the mechanism of oxidative ([OX]) gelation of a thioglycolic acid-capped CdSe QDs film (sol film) submerged in a tetranitromethane (TNM) solution [52]. (B) Sketch of the functionalization of electrodes with [Ni,Fe]O NCs and subsequent removal of oleate ligands with UV irradiation or air plasma [119]. (C) Cartoon that illustrates the CdSe nanocrystals capped with DHLA with the relevant energies for H$_2$ production. dHA indicates dehydroascorbic acid. Potentials are shown versus that of an NHE at pH = 4.5 [48]. Reprinted with permission from [52,119], Copyright 2013,2015 American Chemical Society. From [48], reprinted with permission from AAAS.

5.2 Colloidal chemistry for material discovery
As evidenced in the previous sections, colloidal chemistry is a powerful synthetic approach to access homo-materials and hetero-materials with a precise control on morphology and composition. One aspect that could be further explored is the use of colloidal approaches for material discovery. New light absorbers and electrocatalysts are required, with a more urgent need in OER compared to HER. Our group has recently started to explore the potential of using NC-seeded growth to
synthesize a new light absorber, Sb-BiVO$_4$ in a wide compositional range (figure 6C) [102]. This same approach could be applied to other systems. More efforts should also be directed towards the colloidal synthesis of these materials. Compared to the classical II-VI QDs and to multinary chalcogenide NCs, the chemistry of multinary metal oxides has not been fully elucidated yet. Also, the complexity of their phase diagrams makes not trivial to isolate them in one single crystalline phase. Often amorphous NCs are obtained. Our work on BiVO$_4$ NC ink is one example of the challenges related to the synthesis of colloidal visible light absorbing metal oxides [16]. While targeting BiVO$_4$ NCs, we obtained only phase segregated Bi$_2$O$_{2.7}$/VO$_x$ NC heterodimers, indicating that the chemistry of these vanadates is far from being fully understood. At the same time, we have also demonstrated the power of a rational synthetic design to position dopants in specific lattice sites, for the case of the well-studied N-TiO$_2$ NCs, and to study their impact on optoelectronic properties and photoelectrochemical performance [37,90]. In addition to enable fundamental studies, overcoming the synthetic challenges to access multinary metal oxide colloidal NCs might open new opportunities towards their actual implementation into baggie-type reactors, which will be discussed in the next section.

Another advantage of colloidal chemistry, being an out-of-equilibrium synthetic approach, is the wider compositional space which can be covered compared to other thermodynamic-governed method. One example cited above concerns Ni-Fe-O OER catalysts but many more could be mentioned [117]. As for the electrocatalyst, non-noble metal catalysts with low overpotential for OER have yet to be found. The situation is even more dramatic in terms of overpotential and selectivity if one starts to look at the CO$_2$ reduction reaction, which is out of the scope of the present review. At the nanoscale property modification can be tremendous because of the unique effects, such as the variation of the electronic state density, the charge redistribution, surface- and strain-driven lattice distortion, that evolve with size. The design of atomically-defined samples with tunable size and shape is crucial in determining structure/properties correlations which eventually lead to optimized performance, as seen in many example discussed in this review. In addition to offer a superb control on the morphology of nanocrystalline materials, colloidal chemistry often permits the synthesis of materials unaccessible in their bulk form. Colloidal NCs can indeed be trapped in metastable crystalline phases thanks to the size-dependence of phase transformations (thermodynamics) and, above all, to the rather unique possibility of tuning the chemical potential of the reaction mixture by simply changing reactant concentration, temperature, precursors, surfactants (kinetics). As one example, the metastable TiO$_2$ brookite phase was stabilized over a wide dimensional range (30-200nm) contrary to thermodynamic predictions (figure 1G) [9]. In another example, V$_2$O$_3$ in its metastable bixbyte phase was observed for the first time by exploiting the slow kinetics of transformation in the colloidal reaction mixture [136]. As another example already cited above, Bau et al. were able to synthesize Ni-Fe-O NCs in a wide compositional range in the metastable rock salt phase [119]. The access to phases which are not available in bulk could also open new opportunities in catalysis for water splitting.
5.3 Colloidal nanocrystals for water splitting: only model systems or technologically relevant systems?

This review certainly evidences the role of colloidal NCs as excellent model systems. One question arises about whether colloidal NCs could become technologically relevant systems and which type of water-splitting reactor is better suited for such possibility. The implementation of colloidal NCs into various optoelectronic and electrochemical devices has been increasing over the last years. NC-based solar cells, flexible electronic circuits and, more recently, electrochromic windows and batteries have benefited from the NC solution processability and from the readily tunability of size, shape, crystalline phase and stoichiometries [4,21,137–139]. Devices for water electrolysis, based on electrochemical, photoelectrochemical and photocatalytic approaches, are emerging technologies [140]. Different reactor types have been considered, specifically particle suspension (baggie-type) reactors and panel arrays [2,31,141,142]. Single bed (type 1) and dual bed (type 2) baggie reactors have been proposed. In the former, each particle drives both OER and HER. The production of both H₂ and O₂ in the same vessel might constitute an explosive hazard, thus separation is needed. In the latter, one baggie performs OER and one baggie performs HER. A redox shuttle is required and its choice is crucial to reduce losses in the system. Panel arrays, fixed or tracking concentrator, are constituted by photoanode and photocathode thin films. Photovoltaic electrolysis is an alternative to such integrated photoelectrochemical devices [140,143]. The advantages of one versus the other have been discussed elsewhere [140,144,145]. Colloidal NCs can be assembled in electrodes/photoelectrodes or utilized in solution. A schematic representation of implementation of NCs in different reactor types is shown in figure 10.

Progress still need to be made for an actual practical device based on colloidal NCs. For baggie-type reactors, the state-of-the art for the HNCs driving HER is quite advanced and record quantum yield of hydrogen generation up to 100% have been reached [69,79]. Yet, their activity under solar irradiation remains to be tested and their stability to be improved. As discussed in chapter 3, the slow transfer and subsequent reactions of the photoexcited holes and the ensuing high charge recombination rates often limit the efficiency of the photocatalysts. The use of hole scavengers with proper redox potential has been reported to improve the stability [67]. Also high pH has been reported to improve the stability of CdSe/Ni NRs system because of the redox couple ‘OH/OH operating as a shuttle between the surface of the NCs and the ethanol scavenger in highly alkaline conditions [68,69].

No example relative to hybrid NCs for OER has been reported yet. Studies on redox shuttle are also far from being exhausted. As for thin film photoelectrodes assembled from NCs, new device architectures must be sought after to permit to fully benefit from the size-tunable properties of colloidal semiconductor NCs. At the same time, new metal oxide light absorbers need to be discovered, as discussed in the previous section. The easy processability of colloidal NCs should facilitate their integration in electrolyzers, once long term stability under operational conditions is demonstrated.
Figure 10. Schematic representation of colloidal NCs implemented in different reactor types for water splitting.

6. Conclusions
Herein, we have reviewed the utilization of colloidal NCs as light absorbers and electrocatalysts for water splitting. We have shown how the tunability of these materials is ideal to investigate the complex phenomena behind storing energy in chemical bonds and to optimize performance through structural and compositional modification. Nonetheless, the full potential of colloidal chemistry to advance studies of artificial photosynthesis is far from being fully explored. In this review, the emerging challenge of electrochemical CO₂ reduction has not been discussed. Yet, there are already many examples in the literature which illustrate how colloidal chemistry can aid in revealing design guidelines for producing new catalysts with improved activity and selectivity [20,146,147]. The superb control offered by colloidal chemistry is rapidly expanding towards NC-based multifunctional hybrid materials, where colloidal NCs are interfaced with chemically diverse building blocks, such as polymers or glass matrixes [139,148]. These hybrids are expected to satisfy the demand for complexity of energy devices which is hard to meet by single component materials, thus they might open opportunities to uncover novel phenomena in artificial photosynthesis. As the interest increases and more research groups dedicate their effort to this field, basic research will be key to discover new materials and to propose new device architectures. At the same time, more applied studies will aid to demonstrate the effective utility of colloidal NCs in technologically-relevant photoelectrochemical and electrochemical reactors.
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