Recent Advances in Niobium-Based Materials for Photocatalytic Solar Fuel Production

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Abstract: The search for renewable and clean energy sources is a key aspect for sustainable development as energy consumption has continuously increased over the years concomitantly with environmental concerns caused by the use of fossil fuels. Semiconductor materials have great potential for acting as photocatalysts for solar fuel production, a potential energy source able to solve both energy and environmental concerns. Among the studied semiconductor materials, those based on niobium pentacation are still shallowly explored, although the number of publications and patents on Nb(V)-based photocatalysts has increased in the last years. A large variety of Nb(V)-based materials exhibit suitable electronic/morphological properties for light-driving reactions. Not only the extensive group of Nb2O5 polymorphs is explored, but also many types of layered niobates, mixed oxides, and Nb(V)-doped semiconductors. Therefore, the aim of this manuscript is to provide a review of the latest developments of niobium based photocatalysts for energy conversion into fuels, more specifically, CO2 reduction to hydrocarbons or H2 evolution from water. Additionally, the main strategies for improving the photocatalytic performance of niobium-based materials are discussed.

Keywords: niobium; water splitting; artificial photosynthesis; Nb2O5; niobates

1. Introduction

Increasing energy demand concomitantly with global warming is a major world concern. Currently, fuels and electricity supplies are still strongly dependent on nonrenewable sources. For instance, in 2018, the worldwide consumption of fossil energy was almost 36.5 billion barrels of oil, 3850 billion cubic meters of natural gas, and 3800 million tons oil equivalent of coal per year. As a consequence, more than 33,000 million tons of carbon dioxide was emitted, which represents an annual increasing of 2.0%, the fastest growth for seven years [1]. This scenario can trigger serious problems for the current and next generations such as the insufficiency of fossil fuel resources, air pollution, and also climate changes as a result of the greenhouse effect [2].

In the face of this issue, the development of methods to obtain energy in a clean and sustainable way becomes imperative. Therefore, solar light as a primary energy resource stands out for its advantages in terms of spread availability and accessibility in an inexhaustible and inexpensive way around the world. In addition, obtaining energy from the sun may cause less damage to the environment in relation to waste production, polluting gases emission, or impacts to different ecosystems. The possibility to convert light into valuable solar fuels from water or even CO2 is a promising approach to provide clean
energy and, at the same time, contribute to diminishing the global warming effect [3]. Additionally, the total sum of renewable energy from all the reserves is around 1% of the solar energy supplied to the earth surface by the sun, that is, 173,000 TW. This amount of energy is more than 9000 times larger than the current total energy consumption of the world (17.91 TW in 2017). The challenge lies on the development of economically viable technologies for solar energy harvesting, storage, and utilization [4].

Thus, since Fujishima and Honda [5] reported the possibility of promoting the photo-oxidation of water on the surface of TiO$_2$ photoanodes, semiconductor materials have been extensively investigated as photocatalysts to store solar energy in chemical bonds. Among different photocatalysts, it is worth pointing out Nb(V)-based materials that have been investigated owing to their suitable electronic and morphological properties for light-driving reactions. Nb(V) is characterized by its high affinity to oxygen and its oxides exhibit different properties depending on the preparation and desired application [6,7]. Niobium pentoxide is considered the most thermodynamically stable Nb-based compound with more than 15 different structural configurations. Depending on the crystalline phase, its band-gap energy can vary from 3.2 eV to larger values such as 5 eV [8–10].

Besides Nb$_2$O$_5$ polymorphs, Nb(V) can also combine with oxygen to form extended lamellar structures resulting in polynoy anions, the niobates. They are basically constituted of stacked negative charged layers with intercalated cations [11]. This configuration provides a considerable surface area and allows easy modification either by intercalation, superficial modification, or formation of nanosheets and nanoscrolls [12–14]. Furthermore, Nb(V) is also frequently applied as a doping agent. It is commonly combined to TiO$_2$ and considered one of the most promising dopants for modifying the matrix crystalline structure and minimize the electron–hole recombination [15–18].

In this paper, the recent developments of niobium-based materials for photocatalytic solar fuel production, especially those in the last three years, are systematically reviewed. The main strategies for improving the photocatalytic performance for H$_2$ production from water splitting and CO$_2$ reduction to hydrocarbons, focused on niobium species, will be presented. In Section 2, application to H$_2$ eVolution is discussed, followed by Section 3 related to CO$_2$ reduction.

2. H$_2$ eVolution and Water Splitting

2.1. Niobium Pentoxide (Nb$_2$O$_5$)

Nb$_2$O$_5$ is a typical n-type semiconductor that, along with other oxides, has been widely studied as a photocatalyst thanks to its electronic properties such as excellent chemical and thermal stability, large abundance, rich morphologies, and several polymorphs [19–22]. It is a white solid powder, insoluble in water, and can be dissolved just by fusion with strong basic or acid fluxes. In general, its structure is formed by NbO$_6$ octahedral, which can result in an amorphous state or may crystallize in a wide range of polymorphs with different physical properties [6,8]. Some of those polymorphs are pseudohexagonal (TT-Nb$_2$O$_5$), orthorhombic (T-Nb$_2$O$_5$), and monoclinic (B, H, R, and N-Nb$_2$O$_5$) [19]. The most known nomenclature is based on Brauer’s system, in which the crystalline phases are identified by letters corresponding to the temperature at which they are obtained [8].

Despite these properties, bulk Nb$_2$O$_5$ is not able to achieve a considerable photoactivity owing to its low specific surface area and fast recombination rate of the photogenerated charge carriers [21]. In this sense, several approaches have been applied to increase its photocatalytic performance. One of them is to control the shape and morphology of particles in order to tune the Nb$_2$O$_5$ properties. This can be possible, for instance, by selecting a suitable precursor to obtain the desired configuration of Nb$_2$O$_5$. Wen and coauthors firstly synthesized orthorhombic Nb$_2$O$_5$ (O-Nb$_2$O$_5$) by a hydrothermal reaction using K$_4$Nb$_6$O$_{17}$ 4.5H$_2$O as a precursor. This sample exhibited rectangle nanosheets with a dominantly exposed (010) plane (Figure 1a–d) [22]. After acid treatment of K$_4$Nb$_6$O$_{17}$ 4.5H$_2$O to get H$_4$Nb$_6$O$_{17}$ 3H$_2$O, the ion-exchanged material was treated hydrothermally for 24 h at different conditions of temperature (195–215 °C) and pH (1.5–11.5), as shown in Figure 1e. At first, by changing
the temperature at pH 3.5, X-ray diffraction (XRD) results showed that the O-Nb2O5 phase began to be formed at 200 °C, and at 210 °C, the peaks of H4Nb6O17·3H2O disappeared completely and all observed peaks could be indexed as the O-Nb2O5 phase. From the Figure 1e, it can be also observed that the O-Nb2O5 was obtained in all selected pHs; however, the authors reported that, at pH 3.5, the material showed higher crystallinity and more regular morphology. A commercial O-Nb2O5 sample with spherical morphology was used for comparison and exhibited a band gap value of 3.2 eV, in contrast to 3.0 eV for the as synthesized O-Nb2O5 rectangle nanosheets. Then, the photoelectrochemical activity of both samples was evaluated by measuring the photocurrent resulted from H2 evolution under 370 nm UV light irradiation (30 W·m⁻²). Both photoelectrodes show very low dark current, but under illumination, the O-Nb2O5 nanosheet-based photoelectrode exhibited photocurrents 4.3 times higher than the commercial sample. The authors could justify the improved performance to a larger number of charge carriers coming from the exposed (010) facet with lower band gap energy, fewer recombination centers, and also faster charge transfer by the regular rectangle nanosheets with high crystallinity.

![Figure 1](image_url)

Figure 1. O-Nb2O5 microscopies images of (a) field emission scanning electron microscopy (FE-SEM), (b) high-resolution transmission electron microscopy (HRTEM), (c) transmission electron microscopy (TEM), and (d) selected area electron diffraction (SAED) pattern and (e) diagram for the transformation of O-Nb2O5 nanosheets from H4Nb6O17·3H2O (HnbO) nanosheets under hydrothermal conditions. Reproduced with permission from the authors of [22]. Copyright (2019), Elsevier, Amsterdam, Netherlands.

Later, the same strategy was employed by the same group to obtain crystalline hexagonal Nb2O5 (H-Nb2O5) nanobelt clusters with an exposed (−110) plane [23]. In this case, a fiber-like K2Nb2O6·H2O material was used as precursor. In order to improve the photocatalytic performance of H-Nb2O5 under visible light, metallic Ag was deposited by in situ photoreduction. UV/vis diffuse reflectance spectra showed that the absorption edges were 375 nm for the H-Nb2O5 nanobelt clusters and 423 nm for the Ag-modified sample. Then, both samples were evaluated photoelectrochemically under simulated sunlight illumination (100 W m⁻²) at zero applied voltage. The Ag-modified sample achieved a photocurrent of 10.73 μA, 535 times higher than that observed for bare H-Nb2O5, which was attributed to the improved light-harvesting efficiency as well as the better charge transfer performance.

Zhou and coauthors reported monoclinic Nb2O5 nanorod superstructures by a similar route using Sm2Nb2O7 nanoparticles as the precursor [24]. SEM images showed that Nb2O5 nanorods were obtained with lengths around 2 μm and diameters of 200 nm. The light absorption properties
of the Nb$_2$O$_5$ nanorods were compared to those of commercial Nb$_2$O$_5$ powders and a considerable blue shift was observed in the absorption edge, which could be related to the nanosize of the former. Photocatalytic H$_2$ production was also evaluated with 0.5 wt% Pt as co-catalyst in 25% methanol aqueous solution, employing a 500 W high-pressure mercury lamp as light source. The Nb$_2$O$_5$ nanorod superstructures sample showed a H$_2$ evolution rate of 91 µmol h$^{-1}$, whereas the commercial one exhibited a rate of 0.6 µmol h$^{-1}$, ca. 150 times lower than that for the Nb$_2$O$_5$ nanorod. Moreover, Nb$_2$O$_5$ nanorod superstructures maintained an almost constant photocatalytic H$_2$ production rate after 24 h. The higher performances were explained by the authors based on the superstructures of the material (Figure 2), in which photogenerated electrons could be transferred along the 1D nanorods direction and also transported to the neighboring nanorods, leading to a more efficient separation of photogenerated charge carriers.

![Figure 2. Scheme of Nb$_2$O$_5$ nanorod superstructures synthesis and the improved photogenerated charge carriers separation. Reproduced with permission from the authors of [24]. Copyright (2019), Elsevier, Amsterdam, Netherlands.](image-url)

Clearly, such morphological and structural transformations lead to changes in the optical and electronic properties of niobium oxide. In this sense, Zhang et al. reported an enclosed porous structure of pseudo-hexagonal tief-tief (TT) Nb$_2$O$_5$ nanowires, which showed a significant decrease in the band gap from 3.22 to 2.95 eV in relation to the nonporous Nb$_2$O$_5$ nanowires, and then an enhancement of H$_2$ evolution under visible light irradiation ($\lambda > 420$ nm) [25]. The material was prepared by a solvothermal synthesis using ammonium niobate(V) oxalate hydrate and oleic acid mixed with trioctylamine. The sample was calcined at different temperatures. Thermal treatment at 600 °C and 700 °C causes the formation of orthorhombic phase, while at 580 °C, the pristine TT-phase was formed, so this last condition was selected for further analysis. In TEM images, it was observed that the porous nanowires were shorter than 1 µm and exhibited well separated pores inside the structure, whereas nonporous ones were longer than 5 µm and showed ripple-like contrast owing to bending-induced strain. The authors could explain the formation of such porosity to the replacement of oxalate anion by oleic acid and trioctylamine in the Nb center, resulting in a structure with hydrophobic core and surrounding Nb species. Then, the subsequent heating treatment induced the formation of pores from the cores and the Nb$_2$O$_5$ crystalline nanowires from the coordination structure. However, the samples were not characterized by isotherms of N$_2$ adsorption to be properly classified as porous material. The authors suggested that under-coordinated ions in the twisted lattice around the pores could generate mid-gap states, and hence lower band gap energies. As a result, the H$_2$ production rate of the porous sample was 243.8 µmol g$^{-1}$ h$^{-1}$, considerable higher than the nonporous one, which exhibited H$_2$ evolution rate of 113.1 µmol g$^{-1}$ h$^{-1}$ under similar conditions, and even higher than that for P25 TiO$_2$ (129.6 µmol g$^{-1}$ h$^{-1}$).

Besides the morphology, other strategies can be applied to increase the photocatalytic activity of Nb$_2$O$_5$ species. The introduction of defects into the crystal lattice could be used to engineer the electronic transport properties along the oxide structure or even to create new trap states inside the forbidden band gap region [26]. Zhao and coauthors reported improved solar-light absorption
via partial reduction of Nb$_2$O$_5$ nanorods with active exposed (001) surfaces, which result in the black niobium oxide [27]. In summary, the previous prepared Nb$_2$O$_5$ was thermally treated in an aluminum reduction device, where the sample and aluminum powder were positioned independently in a two-zone tube oven, and then the Nb$_2$O$_5$ and aluminum powders were heated to 500 °C and 800 °C, respectively. This procedure led to the formation of Nb$^{4+}$ centers and hence substantial oxygen vacancies that resulted in a black powder, and thus an enhancement in visible and infrared light absorption compared with the pristine Nb$_2$O$_5$. The reduction of Nb$_2$O$_5$ could be confirmed by Raman and X-ray photoelectron spectroscopies. The Nb$^{4+}$ centers were also characterized by electron paramagnetic resonance. High resolution transmission electron microscopy indicated predominant (001) crystal plane exposed on the nanorod surfaces, suggesting good crystallinity of samples before and after reduction. The samples were evaluated as photoanode using dip-coated FTO glasses in a 1 M NaOH electrolyte under 100 mW cm$^{-2}$ illumination with a 150 W Xe lamp. At 1.23 V, the reduced Nb$_2$O$_5$ sample showed a photocurrent density 138 times higher than that for Nb$_2$O$_5$. The carrier density was calculated by Mott–Schottky plots and was $3 \times 10^4$ times higher for the reduced sample, evidenced an enhanced electrical conductivity, charge transport, and separation. Finally, the samples were evaluated to produce H$_2$ with 0.5 wt% Pt and 20% methanol solution under full-sunlight irradiation from a 300 W Xe lamp, which, for the reduced Nb$_2$O$_5$, resulted in 13 times higher production rate of 274.8 $\mu$mol h$^{-1}$ g$^{-1}$ in comparison with Nb$_2$O$_5$ nanorods (21.1 $\mu$mol h$^{-1}$ g$^{-1}$).

An extended visible light activity of orthorhombic Nb$_2$O$_5$ was achieved by Kulkarni et al. as result of nitrogen doping using a solid state reaction of urea and niobium salt [20]. At 500 °C, ammonia is released and works as nitrogen source to be introduced into the Nb$_2$O$_5$ lattice, filling oxygen vacancies. Field emission scanning electron microscopy (FE-SEM) analysis showed that the sample concentration was crucial for the observed morphologic changes. Concerning the optical properties, the modified samples were analyzed by UV/vis diffuse reflectance and compared to the pristine Nb$_2$O$_5$, which had a band gap energy of 3.4 eV. In general, all the N-doped Nb$_2$O$_5$ samples showed significant absorption in the range of 400–600 nm and a higher red shift in the band gap absorption when the urea concentration was increased (2.6–2.4 eV). Density functional theory (DFT) calculations described that N 2p bands falls above the O 2p bands, and so the top of the valence band gives rise to the reduction in the band gap. At the highest Nb/urea ratio investigated (1:15), this trend was not observed and a higher band gap edge than the expected trend (2.5 eV) was obtained. This would have resulted from nitrogen saturation of the niobia matrix. Photoluminescence measurements indicate that the optimal Nb/urea ratio was 1:10. This sample exhibited the lowest luminescence intensity among the other samples, indicating the efficient electron/hole separation. These results were in agreement with the photocatalytic H$_2$ evolution experiments.

Another strategy to achieve higher photocatalytic performances is coupling to different Nb$_2$O$_5$ compounds with special attributes to create an efficient heterostructured photocatalyst. Owing to its flexibility in morphological, electronic, and structural properties, Nb$_2$O$_5$ is a promising candidate to design an effective heterojunction. Huang and coauthors reported a synergic combination of highly ordered mesoporous Nb$_2$O$_5$ (MNB) nanostructure to two-dimensional N-doped graphene (NGR) through a good interfacial connection (Figure 3) [28]. UV/visible diffuse reflectance spectroscopy evidenced gradual shifts to longer wavelengths as the NGR quantity was increased, so the band gap energies were shift from 3.12 eV up to 2.67 eV. Photoluminescence analysis suggested that NGR supported an efficient separation of charge carriers owing to the fast electron transfer from the conduction band (CB) of MNB to NGR sheets.

Huang and coauthors reported an in situ growth of mesoporous Nb$_2$O$_5$ microspheres on the surface of g-C$_3$N$_4$ [21]. In this case, assuming the photocatalytic activity of g-C$_3$N$_4$ under visible light, the enhanced photoactivity was mainly achieved thanks to the effective interfacial charge transfer between g-C$_3$N$_4$ and Nb$_2$O$_5$. Therefore, it is mandatory to ensure sufficient interfacial contact, and this condition was succeeded by the authors via a so-called in situ self-assembly process of synthesis. The introduction of Nb$_2$O$_5$ microspheres on g-C$_3$N$_4$ caused an expressive enhancement
on the specific surface area. UV/visible diffuse reflectance spectroscopy showed that g-C₃N₄ and the composites presented an absorption edge at ca. 450 nm, while pure Nb₂O₅ had slight a band edge of ca. 420 nm. The slightly visible light absorption of Nb₂O₅ was justified by traces of carbonate species from the thermal decomposition of the triblock copolymer used in the synthesis. Photoluminescence spectroscopy evidenced that pure g-C₃N₄ exhibits stronger emission than the composites. Photocatalytic activities for H₂ eVolution over the samples were eValuated and the composite with 38.1 wt% of Nb₂O₅ exhibited the highest H₂ eVolution rate. With higher amounts of the Nb₂O₅ (69.6 wt%), the photocatalytic performance was still higher than pure g-C₃N₄, but slightly decreased in relation to the composite with 38.1 wt% of Nb₂O₅. This fact could be related to a less effective interfacial contact between the two semiconductors and to a considerable reduction of light absorption by the composite. Other successful combinations between Nb₂O₅ and g-C₃N₄ are reported elsewhere [29,30].

![Diagram](image_url)

**Figure 3.** Schematic illustration of the transfer of photoexcited electrons and production of H₂ photocatalyzed by MNb–N-doped graphene (NGR) under visible light. Reproduced with permission from the authors of [28]. Copyright (2019), The Royal Society of Chemistry, London, United Kingdom.

Therefore, it can be observed that Nb₂O₅ pristine or modified has been employed successfully as a photocatalyst for H₂ eVolution. The role of the crystalline phase, morphology/shape, defects/oxygen vacancies, and heterojunction formation were investigated and showed great contributions for H₂ eVolution activity of Nb₂O₅.

2.2. Niobium Layered Compounds

Layered niobates are composed by a repetition of [NbO₅] octahedral units connected through adjacent or opposite sharing edges or corners, resulting in an extended 2D layered arrangement. This configuration is built by the stacking of negative charged layers separated by cations in the interlayer space [11,31]. The first niobate phases were reported in 1955 by Reisman and Frederic [32]. In this work, they could investigate the products K₃NbO₅, KNbO₅, K₄Nb₆O₁₇, KNbO₆, and “K₆Nb₄₄O₁₁₇” from the reaction of K₂CO₃ and Nb₂O₅ by differential thermal analysis [32,33]. Then, in 1969, the K₄Nb₆O₁₇ and KNbO₆ crystals were further investigated by Nassau and coauthors [33]. In 1982, a new series of phases MCa₂Nb₅O₁₀ (M = Li, Na, K, Rb, Cs, NH₄, Tl) was reported by Dion et al. [34]. Similarly, some years later, Jacobson and coauthors were able to prepare and investigate the interlayer reactivity of the series of layered compounds K[Ca₂Naₙ₋₃Nb₅O₃n+1] [35]. Since then, a class of layered perovskite with the general formula Aₓ[Bₘ₋₁NbₓO₃n+1] is known as Dion–Jacobson compounds (where A represents an alkaline monocation; B an alkaline earth ion; m = 1, 2, and 2 ≤ n ≤ 7; n indicates the number of [NbO₅] chains that form each perovskite-like slab) [11]. Moreover, plenty of different niobates are currently known such as alkali (M⁺NbO₃), columbite (M²⁺Nb₂O₆), and rare-earths orthoniobates (YNbO₄) [6].

These materials are known as promising photocatalysts not just because of their different structures and band gap energies, but also their extended 2D layered arrangement, which provides a considerable surface area and allows easy modification either by intercalation, superficial modification, or formation
of nanosheets and nanoscrolls through exfoliation with bulky $n$-alkylammonium salts of mboxiteB11-catalysts-682272,B31-catalysts-682272,B36-catalysts-682272. Recently, our group investigated the influence of the medium, co-catalyst precursor, and heat treatment in the photoactivity of Pt-modified hexaniobate (K$_{4-x}$H$_x$Nb$_6$O$_{17}$) for H$_2$ evolution [31]. The effect of the Pt(0) precursor and the deposition method (absorption or impregnation) was also investigated. These different conditions of the samples’ preparation mainly affected their configuration and morphology. At the end, the H$_2$ evolution performance was dependent on those conditions, where a ‘soft’ photoreduction of platinum precursors was identified as the best method for the preparation of those photocatalysts.

Suspended hexaniobate layers (K$_{4-x}$H$_x$Nb$_6$O$_{17}$) in tetrabutylammonium hydroxide (TBAOH) showed a good performance in designing the photocatalytic film through the layer-by-layer (LbL) deposition technique [36]. Thin films were assembled by alternative immersions of FTO substrate into exfoliated hexaniobate with pre-adsorbed [Pt(NH$_3$)$_4$]$^{2+}$ suspension (pH = 8) and poly(allylamine hydrochloride) solution (pH = 4) up to 25 bilayers. Further thermal treatment at 500 °C removed the organic species, leading to a fuzzy assembly of hexaniobate nanoscrolls with Pt evenly distributed in the surface, as shown in the FE-SEM image (Figure 4). In this case, it could be concluded that the scrolled morphology was favored in relation to the opened sheets. The Nb-based films were photocatalytic to produce H$_2$ from 20% (v/v) methanol/water solutions under UV-irradiation. The bare hexaniobate LbL films were able to photocatalyze the H$_2$ evolution, with the apparent quantum yield being proportional to the number of deposited bilayers. This behavior evidenced that no photocatalytic surface is lost owing to material deposition. Moreover, when the Pt nanoclusters were added to the film composition, the observed H$_2$ evolution rates were approximately two times higher, reaching 4.0% ± 0.5%.

![FE-SEM image of the surface of Pt-modified hexaniobate layer-by-layer (LbL) films with 25 bilayers (the bright small spots are Pt nanoclusters). Reproduced with permission from the authors of [36]. Copyright (2019), Elsevier, Amsterdam, Netherlands.](image)

Oshima and coauthors reported the photocatalytic activity of Pt-deposited KCa$_2$Nb$_3$O$_{10}$ for water splitting [37]. With an adsorption method, cationic precursor of Pt ([Pt(NH$_3$)$_4$]Cl$_2$) was attached on niobate surface (1.3 wt%) by the TBA$^+$/Ca$_2$Nb$_3$O$_{10}^-$ suspension, followed by restacking with potassium hydroxide and reduction with H$_2$ gas in various temperatures. The authors noted that the deposition of Pt occurred in the interlayer spaces of the restacked nanosheets. Additionally, the increase in the annealing temperature up to 973 K decreased the Pt content in the KCa$_2$Nb$_3$O$_{10}$ interlayer space. Diffuse reflectance and X-ray photoelectron spectroscopies suggested that the adsorbed Pt was reduced from an oxidized to the metallic state according to the applied temperature, and that interlayer Pt species were less susceptible to reduction. The samples were evaluated for water splitting in NaI 10 mmol L$^{-1}$ solution and using a 300 W xenon lamp (λ ≥ 300 nm). The maximum rate of both H$_2$ and O$_2$ production was achieved with the sample treated at 473 K, and higher reduction temperatures.
led to a decline in the photoactivity. This fact could be explained by an improved contact between Pt and the niobate and the formation of suitable active sites. At higher temperatures, the Pt species were located mostly at the external surface. The authors also investigated the impact of the Pt valence state on water splitting activity, comparing the performance of samples heated at the 573 K under H2 and air atmosphere. By TEM, both methodologies led to the formation of Pt with the same size. However, more electron-deficient Pt species were identified on the sample treated in air, which performed a better overall water splitting reaction than the sample treated at H2.

Distinct species to form the layered Dion–Jacobson materials AB2Nb3O10 also play an essential role in the physical properties of the material and especially in their photocatalytic performance. From this point of view, Kulischow et al. investigated a family of layered Dion–Jacobson perovskite-type materials AB2Nb3O10 (A = K, Rb, Cs and B = Ca, Sr, Ba) and their photocatalytic activity for hydrogen production [38]. Seven different niobates were prepared by the molten salt method mixing BCO3 or B(NO3)2 (B = Ca, Sr, Ba), Nb2O5, A2CO3 with ACl (A = K, Rb, Cs), and at specific reaction temperatures (750–1200 °C). UV/vis diffuse reflectance revealed that the band gap energies varied only with different B-cations, seeming to be independent of A-cations. In the case of Ca2+ as B-cation, the band gap value was 3.6 eV, whereas it was 3.3 eV for Sr2+ and 3.0 eV for Ba2+. This trend suggested a dependence on the ionic radii of the B-cations that filled the spaces among NbO6 octahedral units. With bigger B-cation, the overlap of the Nb 4d orbitals was greater, stabilizing the conduction band energy and, as a result, the band was shifted negatively. The prepared niobates were eValuated for H2 eVolution in 10% methanol aqueous solution without a co-catalyst, and then Rh was in situ photodeposited (0.05 wt% Rh loading) on the catalyst. Without co-catalyst, the KCa2Nb3O10 sample exhibited the highest H2 eVolution rate; however, after the Rh addition, the highest overall hydrogen production was achieved by CsCa2Nb3O10. In this case, photocatalytic behavior was influenced by the type of A-cation at the interlayer, suggesting a dependence on the ionic radii in the order Cs > Rb > K, with the exception of RbSr2Nb3O10. The B-cation influenced the hydrogen generation rates owing to more cathodic conduction band edges in the order Ca > Sr > Ba.

Zhou and coauthors showed that the performance of KCa2Nb3O10 for H2 eVolution could be enhanced through a nitrogen and tetravalent niobium doping to generate yellow and black [Ca2Nb3O10]− nanosheets [39]. After the preparation of KCa2Nb3O10 by solid state reaction, the material was treated at 800 °C for 5 h under NH3 flow, resulting in the black niobate (N−/Nb4+ -codoped). The previous addition of a certain stoichiometric excess of K2CO3 prevented the formation of oxygen defect, and thus the reduction of Nb5+ ions, which resulted in the yellow niobate (N-doped). Then, the samples were acid-exchanged, followed by the exfoliation reaction with tetrabutylammonium hydroxide (TBAOH). X-ray absorption fine structure (XAFS) data on the Nb K-edge (18.986 keV) of all the samples indicate that doping did not change the perovskite-type crystal structure, as all three samples showed very similar overall profiles. However, an increase of the pre-edge peak (18.985 keV) of black niobate could be attributed to the partial reduction from Nb5+ to Nb4+ and the increased Nb–N bonding for yellow niobate, whereas the black one, showing a simultaneous decrease of both Nb–O and Nb–N, could point out the dominating generation of oxygen vacancies during the ammoniation process without K2CO3 additive. The samples were eValuated for photocatalytic H2 production using 20 vol% methanol aqueous solution under full range irradiation with an Xe lamp. Undoped nanosheets show very low photocatalytic activity (11.4 mmol h−1) compared with the activity of yellow and black niobate nanosheets (42.9 and 154.7 mmol h−1, respectively). In this sense, the self-doping with Nb4+ could significantly promote an enhanced separation efficiency of photogenerated carriers as well as the light absorption beyond single N3− doping. When Pt was loaded at 0.5 wt%, the H2 eVolution rates were three times higher, 190.1 and 429.5 mmol h−1 for yellow and black niobate nanosheets, respectively. The last one could achieve energy conversion efficiency for solar hydrogen production of 2.7%.

Later, the same group could also investigate the combination of elemental doping, liquid exfoliation, and composition control for a series of typical Dion–Jacobson phases KCa2Na_{n−3}Nb_{n}O_{3n+1}
photocatalysts [40]. As seen in the scheme presented in Figure 5, KCa$_2$Na$_{n-3}$Nb$_n$O$_{3n+1}$ were prepared with various values of $n$ in order to tune the thickness of the perovskite layer. Theoretically, using $n = 3$–$6$, the desired thickness of the layer should increase from 2.4 to 4.0 nm. The materials were obtained from the molar ratio of K/Na/Ca/Nb = 1.05/2/1.05($n$–3)/$n$ through calcination at 1200 °C, followed by the N/Nb$^{4+}$-doping via heat treatment under NH$_3$ flow. This method resulted in samples with various colors, varying from white to black. Samples before and after NH$_3$ treatment were analyzed by UV/vis diffuse reflectance, where red-shifts of the absorption edge were observed in all N/Nb$^{4+}$ doped samples. Even after liquid exfoliation of bulk doped samples to obtain nanosheets, their suspension in water kept the colors from the bulk materials. TEM and atomic force microscopy (AFM) images revealed ultrathin sheets with several micrometers in size and thicknesses of 2.4, 2.8, 3.3, and 4.0 nm for the $n = 3$, 4, 5, and 6 homologues, respectively. The photocatalytic performance of the doped nanosheets was evaluated towards hydrogen evolution from 20 vol% methanol solution under full arc irradiation (Xe lamp, 300 W). Considering all samples with and without 0.5 wt% Pt, the $n = 4$ product exhibited the highest H$_2$ production rate. This fact can be related to the migration length, which affects the electron-hole separation step. The higher photocurrent measurement was also found for $n = 4$ with a sequence of $n = 4 > 5 > 3 > 6$, which is consistent with H$_2$ evolution experiments.

The photocatalytic activity of HCa$_2$Nb$_3$O$_{10}$ for hydrogen production was also extended for visible irradiation region by combination with CdS [41]. In this composite, after light excitation of CdS, the photo-generated electrons were transferred to the calcium niobate nanosheets thanks to their matched conduction band position of $-1.32$ eV for CdS and $-1.18$ eV for niobate (V vs. Ag/AgCl, pH = 7.0). So, the observed increase of the electron paramagnetic resonance (EPR) signal after illumination is the result of the efficient electron transfer that further induced the growth of oxygen vacancies. The samples were evaluated for photocatalytic H$_2$ evolution in (0.1 M) SO$_3^{2-}$/S$^2$ solution under visible light, in which the exfoliated niobate was not able to be excited. The best performance was achieved by samples containing 53.6 wt% of CdS with a H$_2$ production rate of 16.5 μmol h$^{-1}$, about four times higher than pure CdS. Further increase of CdS resulted in lower activity, which was related to the aggregation of CdS nanoparticles. Later, a similar approach was applied by Hu and coauthors to design nanohybrid of CdS and HCa$_2$Nb$_3$O$_{10}$ and get photocatalytic H$_2$ production under visible light [42].

Figure 5. Schematic representation of (a) the general structure of KCa$_2$Nb$_3$O$_{10}$ (KCNNO), (b) KCa$_2$Na$_{n-3}$Nb$_n$O$_{3n+1}$ with thickness control of the perovskite layer by varying the $n$ value, and (c) preparation of N/Nb$^{4+}$ codoped nanosheets (CNNO). Reproduced with permission from the authors of [40]. Copyright (2019), John Wiley & Sons, Inc., Hoboken, NJ, USA.
It has been shown that CdS has more negative conduction band energy than the employed niobate, indicating the spontaneous transfer of photoinduced electrons from CdS to niobate nanosheets.

Xiong and coauthors could improve the photocatalytic hydrogen production activity of HNb$_3$O$_8$ nanosheets by loading Cu with a facile photodeposition method [43]. Exfoliated HNb$_3$O$_8$ niobate was dispersed in Cu$^{2+}$ solutions with corresponding loadings of 2%, 1%, 0.5%, and 0.25% which was photodeposited in situ in triethanolamine solution. Firstly, the authors evaluated the samples for H$_2$ evolution activity with 10 vol% of triethanolamine aqueous solution under the simulated solar light using a 300 W Xenon lamp. The photocatalytic performance increased after addition of Cu$^{2+}$ up to 0.5%, reaching the maximum rate of 59.1 μmol h$^{-1}$. With higher amounts of Cu, the shielding effect was detrimental for the photoactivity. In addition, when methanol was used as the sacrificial agent, the photocatalytic hydrogen production activity achieved the value of 98.2 μmol h$^{-1}$. The authors argued that in situ photodeposition induced selective deposition on the photocatalytic reduction sites of HNb$_3$O$_8$ nanosheets rather than arbitrary distribution.

The same type of niobate was recently applied to engineer a stable nanocomposite photocatalyst. Xia and coauthors reported a method to growth NiS in a highly dispersive way on the HNb$_3$O$_8$ surface [44]. In general, the random deposition and easy agglomeration in large particles of NiS are described as being problematic for the photocatalytic H$_2$ evolution. Therefore, an electrostatic adsorption/self-assembly methodology was developed to obtain NiS on the HNb$_3$O$_8$ nanosheet as an efficient photocatalyst for the first time. The photocatalytic H$_2$ evolution was performed with a 300 W Xenon lamp in a 10% triethanolamine (TEOA) aqueous solution, where the bare NiS and layered KNb$_3$O$_8$ showed no or very low H$_2$ production. In this condition, for the composite with 1 wt% NiS, the H$_2$ evolution rate was 1519.4 μmol g$^{-1}$ h$^{-1}$, higher than the physical mixture of the components (582.5 μmol g$^{-1}$ h$^{-1}$). Time-resolved fluorescence decay analysis was applied to calculate the lifetime of the photogenerated electrons and holes. The kinetic data showed a slower decay for the composite, which indicates that the applied methodology resulted in more efficient utilization of photogenerated electrons and holes.

The potential of niobates structures for water splitting has also been demonstrated by some theoretical works. The perovskite-type niobate NaNbO$_3$ was selected by Wang and coauthors to investigate the effect of anionic monodoping with N, C, P, and S dopants, as well as with (N + N), (C + S), and (N + P) codoping pairs by hybrid density functional theory calculations [45]. At first, the direct band gap of pure cubic NaNbO$_3$ was predicted to be 3.30 eV, with the conduction band mainly formed by Nb-4d orbitals and valence band by the O-2p orbitals, and almost no Na-related states around the band edge were found, indicating that the Na atoms have negligible effects on the electronic structures near the Fermi level. In this sense, one of the O atoms in the cell was firstly substituted by one of four different species (N, C, P, and S), corresponding to a doping concentration of 2.5%. In the case of N-doped NaNbO$_3$, a hole would be generated as the N dopant has less valence electrons than the O atom. The higher energy of N-2p states (around 2.0 eV) induced a localized unoccupied impurity state above the Fermi level as well as several occupied impurity states around the valence band of NaNbO$_3$. For C doping, the substitution would result in two holes in the system, however, no unoccupied localized impurity states were found above the Fermi level. As a result, the total magnetic moment was higher and there were several impurity states above the valence band, which shifted up by about 1.58 eV. The doping with P resulted in a spin-polarized ground state as well, and, owing to the P-3p higher energy than the O-2p, several empty impurity states were found near the conductive band. When S was applied as a dopant, no empty states were found within the band gap, but several localized impurity states were localized below the Fermi level, right above the valence band of the niobate, which resulted in a narrowed band gap. This filled impurity states originated from the hybridization of S-3p and O-2p orbitals. Thus, the band gap values for N-, C-, P-, and S-doped NaNbO$_3$ were calculated to be 1.88, 1.61, 1.04, and 2.34 eV, respectively. Despite the narrower band gap energies, some disadvantages were observed as the undesirable empty gap states from N and P monodoping, which may act as trapping and recombination centers, and the elevated
valence band of NaNbO$_3$ after C doping, precluding the oxygen evolution reaction. In this sense, the codoping possibility using pairs of (N + N), (C + S), and (N + P) was also investigated. It resulted in the band values as shown in Figure 6. In comparison with the pristine niobate, for (N + N) codoping, two fully filled states were generated within the band gap and its effective energy was 1.24 eV lower. The (C + S) doping led to mid-gap states above the valence band of host system, and then a narrow band gap of 1.42 eV. For both systems, the band edges matched properly with the redox potentials of water, indicating that the photocatalyst would be suitable for overall photocatalytic water splitting. However, in the case of (C + S) codoping, the conductive band was shown to be longer good for a spontaneous water splitting process even with a band gap value of about 1.33 eV. This study suggested that codoping can be a successful approach to improve the visible light photocatalytic performance of perovskite NaNbO$_3$.

Figure 6. Band edge positions of the pure and codoped NaNbO$_3$. Reproduced with permission from the authors of [45]. Copyright (2019), John Wiley & Sons, Inc., Hoboken, NJ, USA. NHE, normal hydrogen electrode.

Kaneko and coauthors deeply investigated the origin of the visible light absorption and the optical band gap of 1.9 eV in the d1 metallic strontium niobate (SrNbO$_3$) [46]. By computational methods, firstly, they could verify at which electronic states the photoexcitation takes place, from CB → B$_1$ transition or B$_{-1}$ → CB (B$_{-1}$ denotes the band below the conductive band (CB), and B$_1$ denotes the band above the conductive band). By the band structures and projected density of states calculations, it was found that B$_{-1}$ state lies at around ~4 eV and consists of O(p). Thus, from the atomic orbitals constituting each band, it was inferred that the optical gap corresponds to the Nb(d) → Sr(d)/Nb(d) transition, and so the optical gap was attributed to the CB → B$_1$ transition. The shifts of Fermi level as well as the changes in the optical gap concerning the Sr defects and O vacancies were investigated. The authors could conclude that the optical gap hardly depends on the amount of Sr defects. Sr and O are divalent cation and divalent anion, respectively, and the valence of the Nb ion constituting CB did not change even if the Sr defect and the O vacancy were produced in the same amount. Thus, if the valence of the Nb ions constituting CB does not change, the Fermi level will not shift either. Consequently, the optical gap, which is mainly caused by the excitation from the Fermi level to B$_1$, does not change by Sr$_{1-x}$NbO$_{3-x}$. In addition, in the case of Sr$_{0.875}$NbO$_3$ and Sr$_{0.875}$NbO$_{2.875}$, it was found that the latter evidenced the dependence of the optical gap on the amount of Sr defects and its light absorption intensity was larger than that of the former.

Such doped niobates have been prepared and evaluated as photocatalysts. Yu et al. reported at first time the performance of carbon-doped KNbO$_3$ as well as the effect of MoS$_2$ addition in the material for photocatalytic H$_2$ evolution [47]. Both pristine and C-doped KNbO$_3$ presented the orthorhombic phase, however, the latter resulted in peaks of 30–32° shifted to a lower angle, indicating that the carbon atom was doped into the lattice of KNbO$_3$. In addition, the C species were identified as C$^{4+}$ by X-ray photoelectron spectroscopy (XPS) analysis, which meant that it did not replace the position of lattice oxygen. In addition, it was also confirmed that the substitution of Nb$^{5+}$ by C$^{4+}$ occurred and
also generated some oxygen vacancies to compensate the charge balance. The pristine niobate showed only UV light absorption with band gap of 3.07 eV. C-doped niobate presented very similar behavior, however, it showed a difference in the peak tailing in the range of 400–600 nm. The samples after the addition of MoS$_2$ in 0.2 wt% also presented the peak tailing in the visible-light region. H$_2$ eVolution was performed in methanol solution (20% v/v) under 300 W Xe lamp irradiation and in situ photodeposition of metallic Pt (0.37 wt%). The C doping of KNbO$_3$ could enhance the H$_2$ eVolution from 5 µmol g$^{-1}$ h$^{-1}$ to 142 µmol g$^{-1}$ h$^{-1}$ in comparison with the pristine sample. The decoration of MoS$_2$ on C-KNbO$_3$ could further increase the rate of H$_2$ eVolution to 1300 µmol g$^{-1}$ h$^{-1}$. The samples were also evaluated under visible light, where pure niobate did not show activity, while C-KNbO$_3$ and MoS$_2$/C-KNbO$_3$ resulted in rates of 4.2 µmol g$^{-1}$ h$^{-1}$ and 9.3 µmol g$^{-1}$ h$^{-1}$, respectively. These activities were much lower compared with values under UV/vis light, indicating that the composite is more suitable for working in the UV region of irradiation.

Columbites are also found in the literature with further applications in photocatalytic research. The effect of structural distortion of SnNb$_2$O$_6$ in comparison with the crystalline SnNb$_2$O$_6$ regarding the photocatalytic activity was investigated by Huang et al. [48]. The crystalline sample exhibits a band gap energy of 2.10 eV, while the band gap energies for disordered and amorphous samples were 2.33 eV and 2.43 eV, respectively. Furthermore, the electronic band gap could be tuned by the structural disorder engineering, as an increase of structural lattice disorder led to an increase of band gap energies. By XPS and UV/vis absorption analysis, the valence band edge of crystalline SnNbO$_6$ was about 1.42 V with the conduction band of −0.68 V, whereas for the disordered sample, the values were 1.98 V and −0.35 V, respectively. As a consequence, the latter exhibited H$_2$ eVolution rates 11 times and 1.7 times higher than that of crystalline and amorphous samples, respectively, under visible light irradiation (λ ≥ 420 nm). The photinduced charge carrier behaviors under pulse laser at 532 nm were investigated by transient absorption spectroscopy (TAS). The three samples presented broad and continuous absorption in the visible range at a delay time of 100 ns, indicating the separation of charge carries at different trap states. This behavior was especially pronounced in disordered SbNb$_2$O$_6$, which indicated a better charge separation. The transient absorption decay kinetics also expressed the same, with effective lifetimes for the amorphous, disordered, and crystalline samples of 0.52 ms, 0.56 ms, and 0.26 ms, respectively.

The influence of type of polymorph on the photocatalytic activity of CuNb$_2$O$_6$ columbite was further explored by Kamimura et al. [49]. Two different polymorphs, monoclinic and orthorhombic, revealed that the local crystal structure of the CuO$_6$ octahedral was strongly correlated to the optical absorption property. Both samples exhibited the same band gap, 2.7 eV. However, two additional broad bands in the near-infrared (NIR) region were observed for the orthorhombic sample, being ascribed to Cu$^{2+}$ d–d transitions in a distorted octahedral ligand field. In principle, this transition is forbidden by the Laporte selection rule owing to the symmetric CuO$_6$ configuration. But in the case of the orthorhombic phase, it was partially allowed because of the Jahn–Teller effects of orthorhombic CuNb$_2$O$_6$. The photocatalytic H$_2$ eVolution was performed using a 10 vol% methanol solution and an Air Mass (AM) 1.5G solar light irradiation. The two polymorphs of CuNb$_2$O$_6$ produced H$_2$ under these conditions, but the monoclinic sample was more active than the orthorhombic one. Thus, to elucidate if the d–d absorption affects the photocatalytic activity, the previous results were compared to those using a near-infrared (NIR) cutoff filter. When the monoclinic sample was tested, the rate of H$_2$ eVolution rate did not change expressively when the NIR filter was applied. In this case, the loss of 20% of activity could be explained by the low transmittance of the employed cutoff filter in the visible region. In contrast, the active for orthorhombic CuNb$_2$O$_6$ decreased 90% in relation to that observed without an NIR filter. On contrary to the monoclinic phase, which was insensitive to the excitation of the d–d transition in Cu$^{2+}$ ions, for the orthorhombic one, the electron 3d$^9$ configuration of Cu$^{2+}$ readily induces structural distortion by Jahn–Teller effects in an octahedral crystal field. The empty 3d orbitals of Cu$^{2+}$ could work as deep traps for the photo-generated electrons, and thus increase the recombination rate, resulting in a significant decrease in photocatalytic activity in the presence of an NIR cut-off filter. With NIR photons,
the photo-generated electrons filled 3d orbitals of Cu$^{2+}$ via d–d absorption, a state that inhibited the recombination of photogenerated electrons in the conduction band. These distinct mechanisms were schematically represented on Figure 7. So, in the case of orthorhombic structure, the photocatalytic performance was dependent on fully filled Cu 3d-orbitals.

**Figure 7.** Photocatalytic process over orthorhombic CuNb$_2$O$_6$, comparing the presence and absence of near-infrared (NIR)-cutoff filter. C.B. and V.B. correspond to conduction band and valence band, respectively. AM 1.5G defines the employed irradiation conditions following the ASTM G-173-03 Standard. Reproduced with permission from the authors of [49]. Copyright (2019), Elsevier, Amsterdam, Netherlands.

Chun and coauthors employed a two-step hydrothermal process to form nanosized ZnNb$_2$O$_6$ columbite [50]. The samples were applied to photocatalytic H$_2$ eVolution in 20 vol% methanol under irradiation of 500 W high-pressure mercury lamp. The sample prepared by the new method produced an optimal activity of 23.6 µmol H$_2$ h$^{-1}$ g$^{-1}$, while a bulk ZnNb$_2$O$_6$ prepared by solid state exhibited a H$_2$ eVolution rate of 9 µmol h$^{-1}$ g$^{-1}$. This fact was attributed to its appropriate crystallinity and high specific surface area (61 m$^2$ g$^{-1}$). Further, the authors eValuated the effect co-catalyst deposition on nano-ZnNb$_2$O$_6$. In this case, both pristine water and methanol aqueous solution media were applied. In pure water, the bare photocatalyst showed an undetectable amount of H$_2$. The optimal co-catalyst was found to be Pt, as the H$_2$ generation rate achieved up to 680 and 3200 µmol h$^{-1}$ g$^{-1}$ in pure water and methanol aqueous solution, respectively. Moreover, with the modified sample and at optimal conditions, a maximal apparent quantum yield (AQY) value of 4.54% was obtained in pure water. The value increases to 9.25% with the assistance of methanol as the sacrificial agent.

The manipulation of composition, morphological, and surface properties of three-dimensional hierarchical Nb$_2$O$_7$(OH) superstructures was achieved through Ti(IV) incorporation, as reported by Betzler and coauthors [51]. Firstly, the undoped Nb$_2$O$_7$(OH) consisted of blocks of corner-sharing NbO$_6$ octahedra, forming hollow cubic superstructures of nanowire networks (Figure 8a). When an average amount of 5.5% titanium was incorporated into the crystal lattice, it formed a spherical morphology, hollow and built up from nanowires as well (Figure 8b). With a higher amount of Ti(IV) (10.8%–31.2%), a cubic morphology with smaller nanowires arranged to form the walls of hollow cubes was observed (Figure 8c). All the samples presented the same nanowire arrangement; however, it became smaller, shorter, and flatter as the quantity of Ti was increased. Both XRD and energy-dispersive X ray (EDX) measurements indicated a homogeneous distribution of titanium, however, with more than 12% Ti, the crystal lattice of Nb$_2$O$_7$(OH) was not able to accommodate it, and then the formation of niobium-doped anatase TiO$_2$ plates was favored for further titanium excess. The specific surface area was also affected and it changed from 79 m$^2$ g$^{-1}$ for Nb$_2$O$_7$(OH) (morphology a) to 132 m$^2$ g$^{-1}$ and 173 m$^2$ g$^{-1}$ for morphologies b and c, respectively. The three samples showed the same band gap energy of 3.2 eV, but an additional absorbance in the 550–850 nm region was observed, which came from oxygen vacancies in the TiO$_2$ plates. Further, the samples were eValuated to H$_2$ eVolution under an Xe lamp irradiation (600 mW cm$^{-2}$), with 10 vol% methanol and in situ photodeposited Pt at 8 wt%. The Nb$_2$O$_7$(OH) resulted a rate of 870 µmol g$^{-1}$ h$^{-1}$, and then, as the Ti concentration increases, the H$_2$ rates were raised to 1773 µmol g$^{-1}$ h$^{-1}$ for morphology b and 1988 µmol g$^{-1}$ h$^{-1}$ for
morphology c. The same trend was also expressed by transient absorption measurements, where morphology a showed an averaged lifetime of $54 \pm 2\ ps$, while morphologies B and C presented lifetimes of $64 \pm 2\ ps$ and $73 \pm 4\ ps$, respectively. The authors explained this behavior by the fewer amounts of hydroxyl groups required for charge neutrality with the replacement of $\text{Nb}^{5+}$ by $\text{Ti}^{4+}$, as they are known for acting as non-radiative recombination sites. The doubled $\text{H}_2$ production rate of morphology b, compared with morphology a, was most likely the result of a combination of its fewer recombination sites and larger surface area. The further, but less pronounced increase of the $\text{H}_2$ production rate for morphology c came from the presence of $\text{TiO}_2$ plates, which did not have such a strong effect on the photocatalytic reaction.

It is clearly noted that niobates offer the possibility to be combined with innumerable co-catalysts. Previous cited works and other examples from the literature are summarized in Table 1.

### Table 1. Summary of the photocatalytic activity of niobates combined with distinct materials towards $\text{H}_2$ eVolution from aqueous solution.

<table>
<thead>
<tr>
<th>Niobate</th>
<th>Co-Catalyst</th>
<th>Light Source</th>
<th>Sacrificial Reagent</th>
<th>$\text{H}_2$ Formation Rate</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{HCa}_2\text{Nb}<em>3\text{O}</em>{10}$</td>
<td>CdS, Pt</td>
<td>300 W Xe, &gt;400 nm</td>
<td>Lactic acid</td>
<td>52.6 $\mu$mol h$^{-1}$</td>
<td>[41]</td>
</tr>
<tr>
<td>$\text{HCa}_2\text{Nb}<em>3\text{O}</em>{10}$</td>
<td>CdS</td>
<td>300 W Xe, &gt;420 nm</td>
<td>methanol</td>
<td>450 $\mu$mol g$^{-1}$ h$^{-1}$</td>
<td>[42]</td>
</tr>
<tr>
<td>$\text{HNb}_2\text{O}_6$</td>
<td>$\text{Cu}^{2+}$</td>
<td>300 W Xe</td>
<td>methanol</td>
<td>98.2 $\mu$mol h$^{-1}$</td>
<td>[43]</td>
</tr>
<tr>
<td>$\text{HNb}_2\text{O}_6$</td>
<td>NiS</td>
<td>300 W Xe</td>
<td>triethanolamine</td>
<td>1519.4 $\mu$mol g$^{-1}$ h$^{-1}$</td>
<td>[44]</td>
</tr>
<tr>
<td>C-doped $\text{KNb}_2\text{O}_7$</td>
<td>MoO$_2$, Pt</td>
<td>300 W Xe</td>
<td>methanol</td>
<td>1300 $\mu$mol g$^{-1}$ h$^{-1}$</td>
<td>[47]</td>
</tr>
<tr>
<td>$\text{H}<em>2\text{Sr}<em>2\text{Ta}</em>{0.5}\text{Bi}</em>{1.5}\text{Nb}<em>2\text{O}</em>{7}$</td>
<td>Ni-CH$_3$CH$_2$NH$_2$</td>
<td>300 W Xe</td>
<td>methanol</td>
<td>372.67 $\mu$mol h$^{-1}$</td>
<td>[52]</td>
</tr>
<tr>
<td>$\text{KNb}_2\text{O}_4$</td>
<td>g-C$_3$N$_4$</td>
<td>1000 W Hg $\lambda$ &gt; 400 nm</td>
<td>dimethylhydrazine</td>
<td>25.0 $\mu$mol h$^{-1}$ g$^{-1}$</td>
<td>[53]</td>
</tr>
<tr>
<td>$\text{K}<em>2\text{H}<em>3\text{Nb}</em>{10.8}\text{O}</em>{30}$</td>
<td>g-C$_3$N$_4$</td>
<td>1000 W Hg $\lambda$ &gt; 400 nm</td>
<td>dimethylhydrazine</td>
<td>27.0 $\mu$mol h$^{-1}$ g$^{-1}$</td>
<td>[53]</td>
</tr>
<tr>
<td>$\text{HNb}_2\text{O}_4$</td>
<td>g-C$_3$N$_4$</td>
<td>1000 W Hg $\lambda$ &gt; 400 nm</td>
<td>dimethylhydrazine</td>
<td>37.0 $\mu$mol h$^{-1}$ g$^{-1}$</td>
<td>[53]</td>
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<tr>
<td>$\text{Ca}_2\text{Nb}<em>2\text{Ta}</em>{10}$</td>
<td>g-C$_3$N$_4$, Pt</td>
<td>300 W Xe, &gt;400 nm</td>
<td>triethanolamine</td>
<td>43.54 $\mu$mol h$^{-1}$</td>
<td>[54]</td>
</tr>
<tr>
<td>$\text{Ba}_2\text{Nb}<em>2\text{O}</em>{15}$</td>
<td>g-C$_3$N$_4$</td>
<td>3 W LEDs, 420 nm</td>
<td>oxalic acid</td>
<td>2673 $\mu$mol h$^{-1}$ g$^{-1}$</td>
<td>[55]</td>
</tr>
<tr>
<td>$\text{Ag}_2\text{Nb}<em>2\text{O}</em>{7}$</td>
<td>g-C$_3$N$_4$, Pt</td>
<td>300 W Xe, &gt;420 nm</td>
<td>methanol</td>
<td>88.0 $\mu$mol g$^{-1}$ h$^{-1}$</td>
<td>[56]</td>
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<tr>
<td>$\text{KTa}_2\text{Nb}<em>2\text{O}</em>{5}$</td>
<td>g-C$_3$N$_4$, Pt</td>
<td>300 W Xe, &gt;420 nm</td>
<td>methanol</td>
<td>86.2 $\mu$mol g$^{-1}$ h$^{-1}$</td>
<td>[57]</td>
</tr>
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</table>

Besides the previous cited materials, niobium is also flexible in terms of designing different layered compounds with distinct elements. Fujito and coauthors reported a visible light response of layered perovskite oxycarbonate $\text{Bi}_4\text{NbO}_4\text{Cl}$ for water splitting [58]. This material consists of single-layer $\text{NbO}_3$ perovskite units separated by ($\text{Bi}_2\text{O}_2$)$_2\text{Cl}$ blocks. It presented a band gap of 2.39 eV with valence and conductive bands of 2.11 eV and -0.28 eV, respectively. Experimental and theoretical results suggested that the valence band was formed by highly dispersive O-2p orbitals coming from the interactions within and between Bi–O and Nb–O slabs. As expected from the conductive band value, $\text{Bi}_4\text{NbO}_4\text{Cl}$ showed activity for $\text{H}_2$ eVolution from an aqueous methanol solution under UV irradiation (300 W Xe lamp) and Pt (0.5 wt%) loading. However, the obtained rate of ~0.1 $\mu$mol h$^{-1}$ was considered low and a higher performance for $\text{O}_2$ was found. Apparent quantum efficiency for $\text{O}_2$ eVolution was determined as ~0.4% with $\text{FeCl}_3$ as an electron acceptor, under monochromatic light at 420 nm (~25 mW/cm$^2$). So, through a Z-scheme mechanism, the system was coupled to a $\text{H}_2$-evolving photocatalyst of $\text{Rh}$-doped $\text{SrTiO}_3$ and $\text{Fe}^{3+}/\text{Fe}^{2+}$. In this way, as in the schematic shown in Figure 9, a simultaneous eVolution of $\text{H}_2$ and $\text{O}_2$ under visible light was successfully observed.
It is clearly noted that niobates offer the possibility to be combined with innumerous co-catalysts. Previous cited works and other examples from the literature are summarized in Table 1.

Figure 8. SEM images of (a) Nb$_3$O$_7$(OH) and (b,c) Ti(IV) modified samples with the titanium content averaged over the morphology indicated relative to niobium (scale bar of 1 \textmu m). Reprinted with permission from the authors of [51]. Copyright (2019) American Chemical Society, Washington, DC, USA.
Wakayama and coauthors reported a novel compound obtained from a mixture of layered RbNdNb$_2$O$_7$ and Rb$_2$CO$_3$ [59]. The new layered niobium oxynitride, Rb$_2$NdNb$_2$O$_6$N, resulted in a structure as Rb$_2$NdNb$_2$O$_6$N H$_2$O composed by double-layer [NdNb$_2$O$_6$N]$^{2-}$ perovskite slabs separated by two Rb cations and one H$_2$O molecule. The material showed a band gap of 2.5 eV with visible light absorption, whereas the precursor RbNdNb$_2$O$_7$ essentially absorbs UV light with a band gap energy of 3.7 eV. In the former case, additional N 2p orbitals in the valence-band formation caused a valence-band maximum that was much more negative. In contrast, as the conduction band consists mainly of Nb 4d orbitals, its value remained basically unaffected. Thus, the H$_2$ evolution was evaluated in the presence of TEOA as an electron donor in dimethyl sulfoxide (DMSO) containing 1 mL of water. The sample was irradiated with visible light ($\lambda$ > 400 nm) by 300 W xenon lamp, and Pt was previously deposited at 0.5 wt%. As shown in Figure 10, a stable H$_2$ evolution was observed for 20 h from the Rb$_2$NdNb$_2$O$_6$N sample, in contrast to the precursor, which did not show a response under visible light irradiation.

Figure 9. Scheme of Z-scheme water splitting of Bi$_4$NbO$_8$Cl coupled with Ru/SrTiO$_3$/Rh photocatalyst via Fe$^{3+}$/Fe$^{2+}$ redox mediator. Reprinted with permission from the authors of [58]. Copyright (2019) American Chemical Society, Washington, DC, USA.

Figure 10. H$_2$ evolution over Pt/Rb$_2$NdNb$_2$O$_6$N in DMSO/TEOA solution containing 1 mL of water. Reprinted with permission from the authors of [59]. Copyright (2019) American Chemical Society, Washington, DC, USA.

Therefore, it can be observed that niobium layered compounds form a versatile Nb-based materials class that exhibits great activity for H$_2$ evolution reaction. Additionally, several procedures to tuning...
the photocatalytic performance niobium layered compounds were discussed, such as changes in the structural and morphology, doping and co-doping, and combination with other semiconductors.

2.3. Nb-Doped Materials

Two of the main drawbacks in the photocatalysis heterogeneous are the rapid recombination of electron/hole pair and semiconductor activation only under UV irradiation. A strategy to increase the photocatalytic performance of semiconductors by overcoming these challenges is the introduction of impurities able to change the electronic behavior of the material, in one word, the doping. Niobium is well known in the literature as a dopant agent, especially for titanium-based semiconductors, as its higher charged ion can substitute the Ti$^{4+}$ site in the oxide structure, and simultaneously play the role of donor, improving carrier concentration and conductivity [60]. For TiO$_2$, the defect state provided by the doping is located upon the conduction band minimum and contributes electrons to the unoccupied Ti 3d orbital without introducing additional states in the bandgap. However, in order to satisfy the charge compensation, the substitution can be accompanied by the incorporation of other defects as oxygen vacancies or even Ti$^{3+}$ defects below the conductive band, which lead to a visible range absorption [61]. In Table 2, examples of Nb-doped semiconductors for H$_2$ eVolution are listed.

<table>
<thead>
<tr>
<th>Table 2. Summary of Nb-doped photocatalysts employed for H$_2$ eVolution from aqueous solution.</th>
</tr>
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<tbody>
<tr>
<td><strong>Semiconductor</strong></td>
</tr>
<tr>
<td>TiO$_2$</td>
</tr>
<tr>
<td>TiO$_2$</td>
</tr>
<tr>
<td>TiO$_2$</td>
</tr>
<tr>
<td>TiS$_3$</td>
</tr>
<tr>
<td>Cu$<em>5$Ta$</em>{11}$O$_{30}$</td>
</tr>
<tr>
<td>Ta$_3$N$_5$</td>
</tr>
<tr>
<td>KTaO$_3$</td>
</tr>
<tr>
<td>BiVO$_4$</td>
</tr>
</tbody>
</table>

The performance of the Nb-TiO$_2$ photocatalyst was eValuated using various sacrificial agents such as methanol, ethanol, and 2-propanol by Fontelles-Carceller [68]. The photocatalyst was prepared with an Nb/Ti ratio of 0.025:0.975, which was confirmed by XPS measurements. In principle, the presence of Nb does not significantly change the band gap (3.15 eV) or the specific surface area (80.5 m$^2$ g$^{-1}$) of the photocatalyst in comparison with pure TiO$_2$ (3.18 eV of band gap and 82.1 m$^2$ g$^{-1}$ of surface area). The H$_2$ eVolution was investigated by the photocatalysts loaded with 0.5 wt% of Pt in presence of a 3:7 (v/v) alcohol/water mixture under irradiation of an Hg-Xe lamp (500 W). For both TiO$_2$ and Nb-TiO$_2$, the maximum rates were observed for methanol, followed by ethanol and 2-propanol. In all tested conditions, as shown in Figure 11, the doped sample showed a significantly better result than the non-doped sample. Both samples presented maximum rates for methanol, followed by ethanol and 2-propanol. In addition, the products formed during the photocatalytic reaction were investigated and, for methanol and ethanol, basically the same compounds were identified. Both samples generated formic acid and methyl formate by methanol oxidation with the addition of formaldehyde for the Nb-doped sample under UV illumination. When ethanol was applied, in all conditions of irradiation and applied samples, acetaldehyde and ethyl acetate were found. Nevertheless, when 2-propanol was applied as a sacrificial reagent, the Nb-doped sample generated acetone and propanone-diisopropyl-acetal, whereas bare TiO$_2$ produced only acetone. So, it was possible to affirm that the presence of Niobium at the surface eVen in a small proportion was
able to generate a new compound not observed for the pristine sample. In this sense, in situ infrared studies were also carried out under dark and illumination conditions and in the presence of the alcohol in water vapor. In principle, under the dark, it was possible to verify the adsorption of the alcohols on the surface of both samples. Under UV irradiation, the infrared spectra showed that alcohol and alcoxy species were consumed and/or desorbed from the surface of the samples. For methanol and ethanol, carboxylate/carbonate species were found as the main surface product. This could indicate the attack from hole to the alcohol to form oxidized species related to the corresponding alcohol/alcoxy adsorbed species. In the case of 2-propanol, carboxylate species were less present at the surface owing to the low coverage of products from the relatively easy desorption of acetone. Regarding the H\textsubscript{2} evolution, the authors suggested that the differences in activity are correlated quantitatively with the average oxidation potential of the alcohol. When comparing methanol with ethanol using infrared spectroscopy, they showed essentially the same behavior of the titania surfaces against the alcohol and eVolving molecules coming from the hole-attack.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure11.png}
\caption{Reaction rate of the hydrogen photoproduction using different alcohols. Me, Et, and Is stand for methanol, ethanol, and 2-propanol (isopropanol), respectively. Reproduced with permission from the authors of [68]. Copyright (2019), Elsevier, Amsterdam, Netherlands.}
\end{figure}

Different materials that were doped by niobium ions to make them able to split water have been reported in the literature. The improvement of WO\textsubscript{3} for water splitting by photoelectrochemical process was achieved by Nb-doping [69]. The morphology of the material turned from nanorod to nanotriangle, which also increased the porosity of the WO\textsubscript{3} films. Also, the band gap increased to 2.74 eV by Nb-doping in comparison with WO\textsubscript{3} (2.63 eV) and additional oxygen vacancies were induced. Besides that, the carrier densities significantly increased from \(~3.44 \times 10^{19}\) cm\textsuperscript{-3} to \(1.27 \times 10^{22}\) cm\textsuperscript{-3}, indicating that Nb\textsuperscript{5+} ions act as donor dopants in the WO\textsubscript{3} lattice. It could improve the charge separation and electron transport, and then the doped sample exhibited higher photocurrent, incident photon to charge carrier efficiency (IPCE) (39\% to 52\% at 300 nm and 1.23 V vs. RHE), and photoconversion efficiencies.

Also, with the objective to promote photoelectrochemical water splitting, niobium could support the activity of BiVO\textsubscript{4} films [67]. In this case, the authors concluded that Nb\textsuperscript{5+} ions were not placed in the crystalline structure of BiVO\textsubscript{4}, rather as niobium oxide or oxochloride. Firstly, the addition of 10\% of Nb(V) resulted a higher IPCE, which reached 75\% in comparison with pristine BiVO\textsubscript{4} of 55\% at 2.08 V versus RHE (380–550 nm). The H\textsubscript{2} evolution experiments were run by an H-shaped cell with pristine or modified BiVO\textsubscript{4} film as a photoanode coupled with a cathode of Pt and carbon paste. After 16 h of irradiation at 2.08 V, the pristine sample reached a maximum H\textsubscript{2} evolution rate of 0.09 mmol h\textsuperscript{-1}, whereas Nb-modified BiVO\textsubscript{4} increased about two times to 0.18 mmol h\textsuperscript{-1}. The authors argued that the higher performance of the modified sample would be related to the modification of the nanostructure of the film, but also owing to other effects not obvious by the obtained data. A high efficiency for the photoelectrochemical water splitting reaction was achieved by doping of SnO\textsubscript{2} nanotubes (NTs) with Nb and N co-doping [70]. Niobium could be ascribe as an effective dopant.
owing to its abundant electronic states and lower ionic radii of Nb$^{4+}$ (69 pm) compared with Sn$^{4+}$ (71 pm), resulting in minimum lattice strain. As shown in Figure 12a, the addition of Nb could improve the absorption of light at lower wavelengths, suggesting a decrease in the band gap of the doped samples, with an increase in Nb content. Then, a further doping with N could result in a significant decrease of band gap to 1.99 eV from 3.1 eV of (Sn$_{0.95}$Nb$_{0.05}$)$_2$O$_2$. The lowering of the optical band gap for this sample could provide an IPCE of 10% at 500 nm and potential of ~0 V (vs. RHE), which was described by the author as the highest IPCE value obtained for semiconductor materials explored so far for photoelectrochemical (PEC) water splitting. The generated H$_2$ and O$_2$ gases at the cathode (Pt wire) and the photoanode are shown in Figure 12b. The theoretical concentration of H$_2$ from Faraday’s Law was also calculated, which was very similar to the experimental one, suggesting a Faradaic efficiency close to 100%. Besides that, the ratio of produced gases was around 2, showing a stoichiometric decomposition of water into H$_2$ and O$_2$. From these as well as the theoretical results, it was possible to conclude that the Nb and N co-doping in SnO$_2$ afford better light absorption properties, increased carrier density, and facile electrochemical charge transfer.

![Figure 12](image-url)  
(a) UV/vis absorption spectra of SnO$_2$ NTs and (Sn$_{1-x}$Nb$_x$)$_2$O$_2$ NTs (x = 0.05, 0.1) and (b) theoretical H$_2$ gas and measured H$_2$ and O$_2$ gases, during 24 h chronoamperometry test of (Sn$_{0.95}$Nb$_{0.05}$)$_2$O$_2$/N performed in 0.5 M H$_2$SO$_4$ solution at 0.75 V (vs. RHE) and 26 °C. Reproduced with permission from the authors of [70]. Copyright (2019), Elsevier, Amsterdam, Netherlands.

3. Photoreduction of CO$_2$

**Artificial Photosynthesis**

Carbon dioxide (CO$_2$) emission from fossil fuels is the main concern when discussing global warming and related climate changes. Because the global energy demand will continue to increase, the consumption of fossil fuels will continue to be high for the next few decades, which undoubtedly brings the necessity of large-scale CO$_2$ mitigation from the atmosphere [71]. The observation of natural photosynthesis in green plants motivated the development of photocatalytic CO$_2$ reduction (artificial photosynthesis), which is one promising solution to transform the current fossil fuel-based economy into CO$_2$-neutral energy systems using the sustainable “photon” economy and mitigate the greenhouse gas effect [3,72–74]. Another advantage of the photocatalytic CO$_2$ reduction is the indirect solar energy storage on carbon-based molecules. Therefore, the artificial photosynthesis acts as killing two birds with one stone in view of protecting the environment and simultaneously supplying renewable energy. However, CO$_2$ is very stable/inert regarding chemical reactions, meaning that CO$_2$ conversion demands a considerable amount of energy; therefore, it is quite difficult to convert it to reusable chemicals [75].

In the beginning of the 1900s, Giacomo Ciamician published a seminal paper with the first idea about the artificial photosynthesis, with the suggestion that society should shift from consuming fossil fuels to generating sustainable energy from the sun [76]. Later, Inoue et al. reported the
photoelectrocatalytic reduction of CO₂ to form organic compounds such as formic acid, formaldehyde, methyl alcohol, and methane. In the presence of various semiconductors, the dependence between the formed products and the semiconductor conduction band energy was demonstrated [77].

The CO₂ photoreduction process is a multielectron process (2, 4, 6, or 8 electrons) that can yield diverse products and occurs through different steps that involve adsorption, activation, and dissociation of the C=O bond. The adsorption and activation step is especially challenging because CO₂ is highly stable and inert [78]. In Table 3, some of the possible CO₂ reduction reactions are summarized along with their respective redox potentials. Selected potentials are compared with Nb-based photocatalysts at Figure 13. The formation of first intermediate (CO₂•−) state imposes a significant energy to the reaction and is frequently identified as the rate determining step. Once CO₂•− is formed, it may be subsequently reduced via the protonation of its oxygen atom, resulting in the formation of *COOH. This intermediate can be reduced to CO and released from the semiconductor surface. Alternatively, CO₂•− may also be reduced via the protonation of its carbon atom to form HCOO•, which is further reduced to formate (HCOO⁻) [79–81]. Therefore, the CO₂ reduction mechanism is very complex and imposes kinetic and thermodynamic limitations.

### Table 3. Common products from CO₂ reduction, their simplified half reactions, and redox potentials at pH = 7.

<table>
<thead>
<tr>
<th>Reactions</th>
<th>E (V vs. RHE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO₂ + 2e⁻ + 2H⁺ → CO + H₂O</td>
<td>-0.53</td>
</tr>
<tr>
<td>CO₂ + 2e⁻ + 2H⁺ → HCOOH</td>
<td>-0.61</td>
</tr>
<tr>
<td>CO₂ + 4e⁻ + 4H⁺ → HCOH + H₂O</td>
<td>-0.48</td>
</tr>
<tr>
<td>CO₂ + 6e⁻ + 6H⁺ → CH₃OH + H₂O</td>
<td>-0.38</td>
</tr>
<tr>
<td>CO₂ + 8e⁻ + 8H⁺ → CH₄ + 2H₂O</td>
<td>-0.24</td>
</tr>
</tbody>
</table>

**Figure 13.** Band gap (Eg) and band positions of Nb-based semiconductors vs. normal hydrogen electrode (NHE). On the right side, the standard potentials of several redox couples are presented for comparison.

The semiconductors that have been investigated for CO₂ conversion present low activity, selectivity, stability, and high band gap values; therefore, challenges remain on how to make this process feasible. The CO₂ reduction reaction competes with H₂ eVolution in protic media, thus efficient catalysts must exhibit high activity for CO₂ conversion with low or negligible activity for H₂ eVolution. This requires a very high activation energy barrier for H₂ eVolution or a change in the reaction pathway to slow down the kinetics of H₂ formation [82].

Few semiconductors have been identified with high surface affinity to CO₂ and, simultaneously, with suitable electronic properties for CO₂ reduction via a photo-activated mechanism. In this sense, Nb-based materials are promising for application in CO₂ photoreduction owing to their properties, such as high acidity, excellent textural properties, and interesting electronic properties. Additionally, it is well-known that the product selectivity is significantly affected by the nanostructure design and the architecture of the photoactive components [83].
Efficient CO₂ reduction involves the optimization of synthetic procedures and the development of different architectures in order to achieve the structural, electronic, morphological, and surface properties required for this application. Despite that the application of Nb-based semiconductors for CO₂ photocatalytic reduction has only recently been explored, some reports have revealed the potential of these structures for artificial photosynthesis. It can be highlighted that some materials, such as alkali niobates (NaNbO₃ and KNbO₃), columbite niobates (CuNb₂O₆ and ZnNb₂O₆), niobium oxide (Nb₂O₅), and niobic acid (HNb₂O₆). Additionally, Nb-based materials have been involved in some strategies to improve the CO₂ photocatalytic reduction activity and selectivity, such as doping and formation of composites.

Shi et al. [84] evaluated the CO₂ photocatalytic performance of NaNbO₃ with different morphologies for the first time. NaNbO₃ in orthorhombic phase were prepared by a conventional solid-state reaction and hydrothermal method, with micron-sized particles and homogeneous nanowires, respectively. It was observed that NaNbO₃ decorated with Pt was efficient to convert CO₂ in CH₄, while there was almost no CH₄ to be detected over pristine NaNbO₃. Additionally, the CO₂ reduction performance of Pt–NaNbO₃ nanowires was higher than the micron-sized particles, owing to the higher specific surface area of Pt–NaNbO₃ nanowires.

It is well-known that the photocatalytic reactions (oxidative or reductive processes) are particularly affected by the semiconductor crystal structure [85]. Therefore, the CO₂ photocatalytic reduction performance of the cubic and orthorhombic NaNbO₃ was evaluated. The samples were obtained using inorganic and organic Nb-precursors by a furfural alcohol-derived polymerization–oxidation [85]. Both samples exhibited the same morphology features, that is, cuboids particles with a size between 30 and 50 nm, likely owing to the use of poloxamers with surfactant behavior (Figure 14). The cubic Pt–NaNbO₃ presented lower band gap energy than the orthorhombic Pt–NaNbO₃. Cubic NaNbO₃ exhibited a better photocatalytic performance for H₂ evolution and CO₂ reduction for CH₄ than orthorhombic NaTaO₃; the authors attributed this behavior to its unique electronic structure, which benefits the generation and migration of photo-generated electrons and holes. This behavior was confirmed for the same group in another publication [86]. The authors also investigated the effect of annealing temperature in the CO₂ reduction performance, and a deleterious effect with the increase in the temperature was observed.

![Figure 14. TEM images of (a) NaNbO₃–cubic and (b) NaNbO₃–orthorhombic. HR-TEM images of (c) NaNbO₃–orthorhombic and (d) NaNbO₃–cubic. Reproduced with permission from the authors of [85].](image)

On the basis of previous reports, Fresno et al. evaluated the CO₂ photoreduction performance of NaNbO₃ obtained by solid state reaction at 900 °C and compared it with NaTaO₃ photoactivity [87]. Owing to the high annealing temperature, orthorhombic NaNbO₃ with low surface area was obtained.
The main CO₂ reduction products were CO₂, CH₄, and CH₃OH, together with H₂ coming from the concomitant reduction of water competing with that of CO₂. It was observed that NaNbO₃ and NaTaO₃ give rise to similar conversions in the CO₂ reduction reaction. Additionally, they demonstrated that NaNbO₃ was photoactive even without a co-catalyst, although the main product was CO₂; the authors attributed this difference in the selectivity to the absence of a co-catalyst (Pt) [87].

Inspired in the Degussa P25, which consists of anatase and rutile TiO₂ (i.e., heterostructure) and shows higher activities than either pure anatase or rutile in water splitting and gaseous pollutant photodegradation [88], Ye’s group proposed the investigation of heterojunction formation between NaNbO₃ in cubic and orthorhombic phases for the CO₂ photocatalytic reduction [89]. The NaNbO₃ samples were synthesized at 400 °C to 600 °C based on the polymerized complex method. Pure cubic and orthorhombic NaNbO₃ were obtained at 400 and 600 °C, respectively, while the NaNbO₃ samples with mixed phases were formed at the temperature ranging from 400 to 600 °C (Figure 15b,c). Along with the phase transition from the cubic phase to the orthorhombic phase, the absorption edge of NaNbO₃ shows a slight and gradual blue shift, which is consistent with the previous report that cubic NaNbO₃ has a narrower band gap than orthorhombic NaNbO₃ (Figure 14c). Additionally, the presence of heterojunctions between the two phases was verified; the smaller ones less than 10 nm (cubic phase) and the bigger ones about 30 nm (orthorhombic). For the CO₂ photocatalytic reduction experiment, Pt was used as a co-catalyst and CH₄ was the main product. All the mixed-phase samples were more efficient in CH₄ formation than the pure phases. The authors state that the photo-excited electrons migrate from the conduction band of the orthorhombic phase to the trapping sites on the cubic surface, thus avoiding the electron–hole recombination in orthorhombic NaNbO₃ and improving the charge separation efficiency in the mixed-phase NaNbO₃ samples.

Niobates materials are also efficient photocatalysts for CO₂ photoreduction, because a layered materials configuration is favorable for the separation and transportation of photo-excited carriers (i.e., electrons and holes) [90,91]. In this sense, niobic acid (HNb₃O₈) is a promising material, as it exhibited the valence band and the conduction band edges at desirable potential levels. The layered structure facilitates electron transfer, and the protonic acidity is favorable for water adsorption through hydrogen bonding [91]. It is expected that these properties should exert a certain positive impact on CO₂ photoreduction by water [91].
Therefore, Li et al. [92] eValuated the photocatalytic CO₂ reduction performance of HNb₂O₈ obtained through two different methods, hydrothermal (HT) and solid state reaction (SSR), and compared it with that of KNb₂O₈. HNb₂O₈ is isostructural with KNb₂O₈, and both samples were crystallized in an orthorhombic symmetry. The samples obtained through the HT method exhibited thin nanobelts with several micrometers in length, while the samples obtained by the SSR method showed containing irregular particles in micron size. The surface area value increased from 2.7 m² g⁻¹ of KNb₂O₈ (SSR) to 28.8 m² g⁻¹ of HNb₂O₈ (HT), and from 6.5 m² g⁻¹ of HNb₂O₈ (SSR) to 39.4 m² g⁻¹ of HNb₂O₈ (HT). CH₄ was the only one hydrocarbon product detected in the gas phase during CO₂ photoreduction for all samples. The rate of CH₄ formation over KNb₂O₈ (HT) was seventeen times higher than that over KNb₂O₈ (SSR), and the rate of CH₄ formation over HNb₂O₈ (HT) was about twenty times higher than that over HNb₂O₈ (SSR). Additionally, the solid acid sample (HNb₂O₈) exhibited higher activity than the corresponding potassium salt (KNb₂O₈). Also, both samples exhibited higher activity than degussa TiO₂-P25. Therefore, the sample morphology, specific surface area, and protonic acidity of HNb₂O₈ play important roles in the photocatalytic reduction of CO₂ to CH₄.

In order to improve the CO₂ photocatalytic reduction performance of HNb₂O₈, the intercalation of guest components into the interlayer space was eValuated, and lamellar HNb₂O₈ was purposely pillared with silica [93]. The pillarization of HNb₂O₈ with SiO₂ was verified by interlayer distance expansion, from 1.13 nm to 2.85 nm. Owing to the notably expanded interlayer distance, the surface area value of SiO₂-HNb₂O₈ was as large as 197.3 m² g⁻¹, in contrast to 6.5 m² g⁻¹ of the nonpillared HNb₂O₈. All of the samples were loaded with 0.4 wt% Pt by photodeposition to eValuate the CO₂ photoreduction. The yield of CH₄ over the HNb₂O₈ sample was almost three times higher than that achieved over Nb₂O₅. The improved activity for the solid acid material might be ascribed to its layered structure and the stronger adsorption ability to H₂O through hydrogen bonding. The rate of CH₄ formation was further enhanced with the SiO₂-HNb₂O₈ sample. Additionally, the activity of the current SiO₂-HNb₂O₈ photocatalyst was higher than that of previously reported HNb₂O₈ nanobelts [92]. Compared with nonpillared HNb₂O₈, the SiO₂ pillared HNb₂O₈ sample has a notably expanded interlayer distance and much greater surface area. Thus, the reactive active sites at the interlayer space of SiO₂-HNb₂O₈ should be more accessible to the substrate. To investigate the role of protonic acidity for CO₂ reduction, the activity of SiO₂-HNb₂O₈ was compared with that of SiO₂-KNb₂O₈ and anatase TiO₂ in different molar ratios of H₂O/CO₂ (0.06 or 0.25). The SiO₂–Nb₂O₅ sample showed higher activity than SiO₂–KNb₂O₈ and anatase TiO₂; it is noteworthy that the activity of the SiO₂–HNb₂O₈ sample was more significantly enhanced than the other two samples at an elevated water content. This confirms that water molecules could react easily with the protons at the interlayer space of lamellar solid acids through hydrogen bonding.

Despite columbite niobates exhibiting interesting electronic properties and suitable conduction and valence band energies for CO₂ photoreduction [6], this class of material has been explored only in a limited manner. Kormányos et al. eValuated the CO₂ photoelectrochemical reduction activity of monoclinic type-p CuNb₂O₆ and orthorhombic type-n ZnNb₂O₆ semiconductors [94]. The CuNb₂O₆ and ZnNb₂O₆ samples showed an indirect bandgap of 1.77 eV and 3.55 eV, respectively. The photoelectrochemical (PEC) performance of the p-CuNb₂O₆ sample was tested in CO₂- and N₂-saturated 0.1 M NaHCO₃ solution. The photocurrents are two times higher (at E = −0.4 V) when the solution was saturated with CO₂. According to the authors, the dramatic increase in the presence of CO₂ is clearly attributable to the ability of CuNb₂O₆ to photoelectrochemically reduce CO₂. However, the CuNb₂O₆ sample’s efficiency and selectivity was not eValuated, as the products formed from CO₂ photoelectrochemical reduction were not monitored.

Recently, da Silva et al. investigated the role of Nb₂O₅ surface acidity for CO₂ reduction performance [95]. Nb₂O₅ catalysts were prepared through a modified peroxide sol–gel method using different annealing temperatures, and it was verified that the increase in the annealing temperature decreases the surface acidity. The authors related the activity and selectivity of the Nb₂O₅ samples to their surface acidity; high surface acidity prompted the conversion of CO₂ to CO, HCOOH, and
CH$_3$COOH, whereas low surface acidity led to the conversion of CO$_2$ to CH$_4$. It was verified that CO is the main intermediary during the CO$_2$ photoreduction in all conditions. This work unveiled the importance of surface acidity for CO$_2$ photoreduction.

Despite the remarkable properties of NaNbO$_3$ and interesting performance for CO$_2$ photocatalytic reduction, the relative wide band gap (3.4 eV) limits its photocatalytic activity only to be active in the ultraviolet light region (4% of the solar spectrum). Therefore, there is a growing necessity to develop photocatalysts with an efficient response under visible-light irradiation. In this sense, Shi et al. design a heterostructure between NaNbO$_3$ and C$_3$N$_4$ in order to extended the absorption spectral range for visible region [96]. The NaNbO$_3$ sample was obtained through hydrothermal synthesis, and the heterostructure was tailored by annealing NaNbO$_3$ with melamine (C$_3$N$_4$ precursor) at 520 °C for 4 h. The XRD patterns of g-C$_3$N$_4$/NaNbO$_3$ revealed coexistence crystalline phase of NaNbO$_3$ and g-C$_3$N$_4$ in the composite. NaNbO$_3$ nanowires are found to randomly deposit and distribute on the surface of g-C$_3$N$_4$ sheets, which results in forming a heterostructured g-C$_3$N$_4$/NaNbO$_3$ material. The CO$_2$ photoreduction under visible irradiation showed that g-C$_3$N$_4$/NaNbO$_3$ improved over eight-fold the activity than that of bare C$_3$N$_4$ in the CH$_4$ formation. Such a remarkable enhancement of photocatalytic activity was mainly attributed to the improved separation and transfer of photogenerated electron-hole pairs at the intimate interface of g-C$_3$N$_4$/NaNbO$_3$ heterojunctions. The same group evaluated the CO$_2$ photoreduction performance of g-C$_3$N$_4$/KNbO$_3$, and the photocatalytic activity of g-C$_3$N$_4$/KNbO$_3$ for CO$_2$ reduction was almost four times higher than that of individual g-C$_3$N$_4$ under visible light irradiation [97].

Another Nb-based material that is worth mentioning is Sr$_2$Bi$_2$Nb$_2$TiO$_{12}$, an Aurivillius-type perovskite. Yu et al. investigated, for the first time, the Sr$_2$Bi$_2$Nb$_2$TiO$_{12}$ as photocatalyst for CO$_2$ photoreduction [98]. The introduction of oxygen vacancies on the surface of Sr$_2$Bi$_2$Nb$_2$TiO$_{12}$ extends photoreponse to cover the whole visible region and also tremendously promotes the separation of photoinduced charge carriers. Compared with the bulk material prepared by traditional solid-state reaction, the Sr$_2$Bi$_2$Nb$_2$TiO$_{12}$ nanosheets with the optimal oxygen vacancies concentration yields a substantially high CO evolution rate of 17.11 μmol g$^{-1}$ h$^{-1}$.

Niobium has been used also as dopant, as it can improve the photocatalytic activities of anatase TiO$_2$ [61,99–101], by changing the crystal structure, electrical properties, and absorption characteristics of TiO$_2$. It was proposed that Nb would dope into TiO$_2$ lattice and decrease the band gap energy of the photocatalyst, and thereby Nb substitution on the Ti site creates a Nb$^{4+}$ defect state located upon the conduction band minimum. Therefore, Nb doped TiO$_2$ would work as a visible light-driven photocatalyst [61,100,101].

Nogueira et al. investigated the role of Nb-doped TiO$_2$ for CO$_2$ photocatalytic reduction to methanol [102]. The samples were obtained through the Pechini method, which promoted the substitution of Ti$^{4+}$ for Nb$^{4+}$ at different concentrations. It was observed that the surface area tends to increase with the concentration of Nb. The concentration of formed methanol was directly proportional to the Nb concentration in the TiO$_2$, the authors attributed the formation of methanol to the presence of Nb in the lattice of TiO$_2$.

The evaluation of CO$_2$ photoreduction performance of Z-scheme formed between Nb–TiO$_2$ and g-C$_3$N$_4$ was also evaluated [103]. It was confirmed that the band gap energy of the Nb–TiO$_2$ (2.91 eV) was lower than that of the TiO$_2$ (3.2 eV). The formation of Z-scheme Nb–TiO$_2$/g-C$_3$N$_4$ was evidenced through XRD and TEM analysis. CO, CH$_4$, and O$_2$ were formed from CO$_2$ photoreduction when the Nb–TiO$_2$ sample was used. Nb–TiO$_2$/g-C$_3$N$_4$ not only converted CO$_2$ to CO, CH$_4$, and O$_2$, but also generated HCOOH. Additionally, 50Nb–TiO$_2$/50g-C$_3$N$_4$ was the best material for the CO$_2$ reduction.

It is well known that, while metals and semiconductors usually exhibit great stability, but poor selectivity for CO$_2$ reduction, enzymes and metal complexes can selectively convert CO$_2$ into CO, but they lack long-term stability. Therefore, Faustino et al. evaluated the role of hexaniobate nanoscrolls (K$_x$H$_{4-x}$Nb$_6$O$_{17}$) as suitable substrates for the immobilization of two kinds of Re(I)-based molecular catalysts (namely I and II) for CO$_2$ photoreduction [104]. Hexaniobate nanoscrolls were obtained
through solid state reaction with subsequent proton exchange and exfoliation with \textit{tert}-butylammonium hydroxide. The Re(I) complexes were immobilized by an adsorption process that was verified for UV/vis absorption and Fourier-transform infrared (FTIR) spectroscopies. As a result of the hexaniobate exfoliation, the specific surface area was increased from 2 m$^2$ g$^{-1}$ to 110 m$^2$ g$^{-1}$. The sensitized oxides were employed as visible-light photocatalysts for CO$_2$ reduction in 5:1 dimethylformamide (DMF)/TEOA exposed to a 300 W Xe lamp ($\lambda = 420$ nm). The possible gaseous products were analyzed by mass spectrometry with CO being the only detected photoproduct. The complexes immobilized on the surfaces of the oxides exhibited much higher turnover number (TON$_{CO}$) than those in solution. For comparative purposes, the complexes were immobilized on TiO$_2$ as well. While for complex I, the immobilization on the niobate nanoscrolls led to 20% higher TON$_{CO}$ in comparison with that observed on TiO$_2$, for complex II, the opposite behavior was observed; that is, the immobilization on TiO$_2$ led to better photoactivities. Such behavior was rationalized based on the recombination kinetics of each composite and on the reductive potential of the injected electrons on the niobate or TiO$_2$ conduction bands. For comparison, a summary of the photoactivity of Nb-based photocatalysts towards CO$_2$ reduction is presented in Table 4.

Table 4. Summary of the photocatalytic activity of Nb-based compounds combined with distinct materials towards CO$_2$ reduction.

<table>
<thead>
<tr>
<th>Nb-Based Material</th>
<th>Co-Catalyst</th>
<th>Light Source</th>
<th>Main Products</th>
<th>Observed Rate</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaNbO$_3$</td>
<td>Pt</td>
<td>300 W Xe</td>
<td>CH$_4$</td>
<td>653 ppm h$^{-1}$ g$^{-1}$</td>
<td>[84]</td>
</tr>
<tr>
<td>NaNbO$_3$</td>
<td>Pt</td>
<td>300 W Xe &gt; 300 nm</td>
<td>CH$_4$</td>
<td>10.4 µmol h$^{-1}$ m$^{-2}$</td>
<td>[85]</td>
</tr>
<tr>
<td>NaNbO$_3$</td>
<td>None</td>
<td>300 W Xe &gt; 420 nm</td>
<td>CO, CH$_4$, CH$_3$OH, H$_2$</td>
<td>NA</td>
<td>[87]</td>
</tr>
<tr>
<td>NaNb$_5$O$_9$ heterojunction</td>
<td>Pt</td>
<td>300 W Xe &gt; 300 nm</td>
<td>CH$_4$</td>
<td>-10 µmol h$^{-1}$ m$^{-2}$</td>
<td>[89]</td>
</tr>
<tr>
<td>HNb$_3$O$_8$</td>
<td>None</td>
<td>350 W Xe</td>
<td>CH$_4$</td>
<td>3.5 µmol g$^{-1}$ h$^{-1}$</td>
<td>[92]</td>
</tr>
<tr>
<td>HNb$_3$O$_8$</td>
<td>Pt</td>
<td>350 W Xe</td>
<td>CH$_4$</td>
<td>2.90 µmol g$^{-1}$ h$^{-1}$</td>
<td>[93]</td>
</tr>
<tr>
<td>Nb$_3$O$_5$</td>
<td>None</td>
<td>UVC lamp, 0.167 mW cm$^{-2}$</td>
<td>CO, CH$_4$, CH$_3$COOH, HCOOH</td>
<td>-10 µmol L$^{-1}$ g$^{-1}$</td>
<td>[95]</td>
</tr>
<tr>
<td>C$_3$N$_4$/NaNbO$_3$</td>
<td>None</td>
<td>300 W Xe $\lambda &gt; 420$ nm</td>
<td>CH$_4$</td>
<td>-6 µmol h$^{-1}$ g$^{-1}$</td>
<td>[96]</td>
</tr>
<tr>
<td>C$_3$N$_4$/KNbO$_3$</td>
<td>Pt</td>
<td>300 W Xe $\lambda &gt; 420$ nm</td>
<td>CH$_4$</td>
<td>0.25 µmol h$^{-1}$</td>
<td>[97]</td>
</tr>
<tr>
<td>Re(I) poly(pyridyl complexes) (K$_3$H$_4$-Nb$_5$O$_17$)</td>
<td>None</td>
<td>300 W Xe $\lambda &gt; 420$ nm</td>
<td>CO</td>
<td>2.9 *</td>
<td>[104]</td>
</tr>
</tbody>
</table>

* Turnover frequency (TOF), h$^{-1}$.

4. Concluding Remarks and Perspectives

Over the past few years, great efforts have been spent by the scientific community to develop clean renewable energy sources. Photocatalysis can be described as an efficient method to produce solar fuels from available sources as H$_2$O and CO$_2$. Nevertheless, it is always a challenge to develop efficient materials able to promote solar-to-fuel conversion. In this sense, niobium-based materials offer diverse possibilities to engineer compounds showing important characteristics for photocatalytic processes such as efficient light absorption, charge separation, and charge transfer reaction. Herein, we could provide a glimpse from the last years on the extensive flexibility offered by niobium-based compounds for CO$_2$ reduction and notably for H$_2$ eVolution. Niobium oxides as Nb$_2$O$_5$ and the wide range of niobates offer untold possibilities of structure, morphologies, and electronic configuration, which can supply desired properties for photocatalytic reactions. Moreover, niobium cations as a dopant have provided the opportunity to tune band energy positions and increase the efficient of other materials. Thus, niobium-based compounds are promising materials that still offer opportunities of new investigations and insights regarding photocatalytic applications.
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**Conflicts of Interest:** The authors declare no conflict of interest.

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