



Cite this: *Sustainable Energy Fuels*, 2024, **8**, 3001

Future perspectives in green hydrogen production by catalyzed sono-photolysis of water

Piergiovanni Domenighini, ^a Ferdinando Costantino, ^b Pier Luigi Gentili, ^b Anna Donnadio, ^c Morena Nocchetti, ^c Alceo Macchioni, ^b Federico Rossi^a and Franco Cotana^{ad}

The quest for sustainable energy solutions has led to a growing interest in green hydrogen production, with catalyzed sono-photolysis of water emerging as a promising avenue. This perspective highlights the innovative combination of photocatalysis and acoustic cavitation to enhance the generation of green hydrogen from water splitting. By harnessing the power of semiconductor-based catalysts, the sono-photolysis approach capitalizes on solar radiation to initiate water dissociation. Simultaneously, high-intensity ultrasound waves trigger cavitation, creating reactive microbubbles and localized hotspots that further promote hydrogen evolution. Through systematic experimentation and optimization, researchers are investigating the influence of catalyst type, concentration, and ultrasonic parameters on hydrogen production. Excitingly, early results demonstrated a promising synergistic effect between photolysis and sonolysis. These findings traced a new path that is worth being pursued to open the door to scalable, cost-effective, and environmentally friendly green hydrogen production. In this perspective, catalyzed sono-photolysis holds tremendous potential for meeting the world's energy demands sustainably. Its innovative blend of light and sound-driven water splitting paves the way towards a greener future, offering a viable solution for the large-scale production of clean and renewable hydrogen.

Received 26th February 2024
Accepted 10th May 2024

DOI: 10.1039/d4se00277f
rsc.li/sustainable-energy

Introduction

The global pursuit of sustainable and carbon-neutral energy sources has intensified in recent years, driven by the urgent need to mitigate climate change and reduce dependence on fossil fuels. Among various renewable energy alternatives, hydrogen has emerged as a promising candidate due to its high mass energy density and versatility (especially, non-toxicity and zero-carbon combustion). However, conventional methods of hydrogen production, such as steam methane reforming, require fossil fuels and generate significant carbon emissions. To overcome these limitations, the focus has shifted towards green hydrogen production, which utilizes renewable resources and produces no greenhouse gas emissions.

Among the “green” hydrogen production technologies that utilize energy from renewable systems, the following are worth mentioning:

- **Biomass gasification and pyrolysis:** This process involves partial oxidation of biomass derived from animal waste,

municipal or industrial waste, and agricultural waste. Hydrogen production yields typically range from 14.3 wt% to significant peaks of 17 wt% in the case of lignocellulosic biomass;¹

- **Biochemical processes:** These include direct and indirect bio-photolysis, photo-fermentation, or fermentation in the absence of light;²

- **Water electrolysis:** This method involves the redox splitting of H_2O into H_2 and O_2 induced by an electric current being passed through two electrodes in an aqueous solution.³ It currently represents 4% of the global hydrogen production,⁶ with production efficiencies ranging from 60% (alkaline cells) to 80% (molten carbonate cells);^{4,5}

- **Photoelectrolysis:** This process utilizes photo-catalyzed electrolysis. Depending on the solar radiation intensity and the electrodes made of semiconductors that also act as photocatalysts, water splitting can be achieved with current densities ranging from 10 mA cm^{-2} to 30 mA cm^{-2} and an electrical potential difference of $+1.35\text{ V}^6$.

Catalyzed sono-photolysis of water has emerged as an innovative and promising approach for green hydrogen production. This technique combines the principles of heterogeneous photocatalysis and acoustic cavitation to enhance the efficiency of hydrogen evolution. Heterogeneous photocatalysis involves the use of semiconductor materials to absorb solar radiation and induce the formation of electron–hole pairs that initiate the water-splitting reaction. Concurrently, acoustic

^aDipartimento di Ingegneria, Università degli Studi di Perugia, Via Goffredo Duranti, 93, 06125 Perugia PG, Italy. E-mail: piergiovanni.domenighini@unipg.it

^bDipartimento di Chimica, Biologia e Biotecnologie, Università degli Studi di Perugia, Via dell'Elce di Sotto, 8, 06123 Perugia PG, Italy

^cDipartimento di Scienze Farmaceutiche, Università degli Studi di Perugia, Via del Liceo, 1, 06123 Perugia PG, Italy

^dRicerca sul Sistema Energetico RSE spa, Via R. Rubattino 5420134, Milano MI, Italy



cavitation induces the formation and implosive collapse of microbubbles in the liquid, creating localized hotspots and other extreme physicochemical conditions (such as extraordinarily high pressures) favorable for the dissociation of water molecules into their constitutive elements, which are H_2 and O_2 . The integration of photocatalysis and acoustic cavitation in catalyzed sono-photolysis offers several distinct advantages. First, the use of sunlight as the primary energy source enables a renewable and abundant input for hydrogen generation. Furthermore, the incorporation of ultrasonic waves promotes the cavitation phenomenon, leading to the generation of additional reactive species and accelerating the overall hydrogen evolution process. Advantages over the other popular multifield synergistic catalysis such as photo-electrolysis, or photo-thermal, are mainly due to the extremely high energy contribution of the cavitation field. The cavitation phenomenon allows local temperatures and pressures to be reached which are otherwise inaccessible, which trigger the water-splitting reaction and improve the activity of the heterogeneous catalyst. Great opportunities may be unveiled by optimizing cavitation control, and, for this reason, this method of green hydrogen production has recently gained attention in research planning and financial support from both public-private partnerships in various nations and transnational bodies such as the European Union. Governments and organizations worldwide recognize the potential of green hydrogen as a key component in achieving climate and energy transition goals. As a result, they are actively promoting and investing in research and development efforts related to hydrogen production technologies, including catalyzed sono-photolysis.^{7,8} Partnerships between public and private entities are crucial in advancing the development and deployment of green hydrogen technologies. These collaborations bring together expertise, resources, and funding to support research, innovation, and commercialization efforts. They also foster knowledge sharing, standardization, and policy frameworks to create an enabling environment for the widespread adoption of green hydrogen. At the transnational level, the European Union has identified hydrogen as a strategic priority in its efforts to decarbonize the economy. Through initiatives such as the European Clean Hydrogen Alliance and the European Green Deal,⁹ the EU aims to drive research, innovation, and investment in green hydrogen technologies. This includes funding programs, regulatory support, and the establishment of hydrogen partnerships across member states. The involvement of research planning and financial support from diverse nations and transnational bodies reflects the global recognition of the potential of green hydrogen and its role in the transition to a sustainable and low-carbon future.¹⁰ These initiatives demonstrate a collective commitment to driving the development and adoption of green hydrogen technologies on a global scale.

The preliminary findings from sono-photolysis studies are promising, demonstrating significantly enhanced rates of hydrogen evolution and improved conversion efficiencies compared to conventional water-splitting methods. Moreover, the catalyzed sono-photolysis approach offers scalability, as it can be applied to large-scale systems with relatively simple

reactor configurations. The cost-effectiveness and environmental friendliness of the technique further solidify its potential as a viable solution for green hydrogen production.

In this perspective paper, we aim to provide a comprehensive overview of the current state of heterogeneously catalyzed sono-photolysis of water for green hydrogen production starting from the photolysis and sonolysis processes separately. We will explore the fundamental principles, catalyst design strategies, and optimization approaches employed in this field. Additionally, we will discuss the challenges and opportunities associated with scaling up this technology and its potential integration into existing energy systems. Ultimately, this perspective seeks to shed light on the immense potential of catalyzed sono-photolysis in advancing the transition towards a sustainable, carbon-neutral future powered by green hydrogen.

Water splitting

Water splitting (1) is an endergonic redox reaction:



The protons of water are reduced to H_2 , whereas its dianionic oxygen is oxidized to O_2 . A noticeable amount of free energy is required to perform the reaction: under standard conditions, $\Delta G^0 = 237.4 \text{ kJ mol}^{-1}$. The real energy needed is, nevertheless, much higher than that predicted by thermodynamics, owing to the relevant kinetic barriers present on both sides of the redox process, especially in the oxidative one. The needed free energy to split water into molecular hydrogen and oxygen can be given in the form of UV-visible electromagnetic radiation emitted by the sun and reaching the surface of the earth (photolysis). Alternatively, it can be released by ultrasound (sonolysis) or both UV-visible electromagnetic radiation and ultrasounds (sonophotolysis). Active and robust co-catalysts, capable of offering alternative reaction pathways to water splitting, must be developed, and interfaced with the photocatalytic systems, in order to lower down the unfavorable kinetic barriers and thus maximize the energy conversion efficiency.¹¹

The catalyzed photolysis

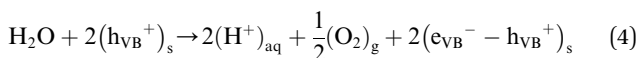
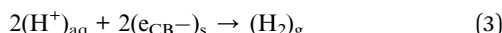
The composition of the solar spectrum corresponds to the thermal radiation emitted by a black body at 5800 K, which is the temperature of the solar surface. The solar radiation is incoherent and polychromatic: its spectrum¹² ranges from UV-C (120 nm) to near-IR (100 μm). The chemical compounds of the atmosphere exert a filter effect and they screen the terrestrial biosphere from UV-C and some bands of the IR. The UV and visible radiations that reach the surface of the earth are not absorbed directly by water. Water molecules absorb some bands of the IR, but IR radiation does not have enough energy to split the chemical bonds of H_2O unless solar concentrators are used to reach very high temperatures.¹³ Therefore, a photocatalyst is needed, *i.e.*, a chemical compound that absorbs some radiation of the solar spectrum with the appropriate energy and transferring it to water, leading to its splitting into H_2 and O_2 , thus



catalyzing the process. From the point of view of large-scale hydrogen production, the use of heterogeneous photocatalysts in the form of particulate semiconductor materials is more practical than other solutions.^{14,15} A semiconductor absorbs photons that have energy equal to or larger than its band gap (BG) (see Fig. 1). Any photon absorbed promotes the jump of an electron from the valence (VB) to the conduction (CB) band of the semiconductor (see eqn (2)): a physically separated electron (e^-) – hole (h^+) pair is generated in the solar radiation harvesting photocatalyst.



These two opposite charges must migrate to the surface of the solid particles without recombining with each other. Then, as soon as they reach the surface, they must participate in the redox reactions required to split water molecules. The electron is supposed to reduce hydrogen whereas the hole converts the oxide to oxygen according to the following two redox semi-reactions (see Fig. 1A):



Reaction (3) is thermodynamically spontaneous if the electric potential associated with the electrons in the CB is more negative than the electric potential of H^+/H_2 , which, under standard conditions, is equal to 0 V. On the other hand, eqn (4) requires that the electric potential of the holes in the VB is more positive than the potential of O_2/H_2O , which, under standard conditions, is equal to 1.23 V.

The sum of eqn (3) and (4) to eqn (2) multiplied by 2 gives the overall photo-induced water splitting:

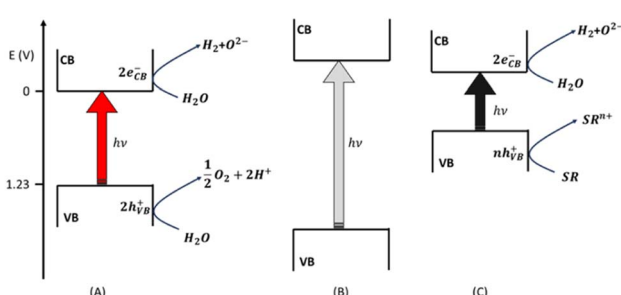
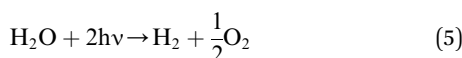


Fig. 1 Water splitting photo-catalyzed by a semiconductor having appropriate electric potential values for the conduction (CB) and valence (VB) bands (A). The semiconductor in (B) has a highly negative CB electric potential and a very positive VB electric potential, guaranteeing the reduction of the over-potentials of the two redox semi-reactions, but generating a broad BG. Graph (C) shows the case of a semiconductor that (1) has a narrow BG, (2) oxidizes a sacrificial reagent (SR), and (3) has a negative CB electric potential assuring the production of H_2 from water.

Based on these considerations, it might appear that two photons having an energy of 1.23 eV (*i.e.*, a wavelength of $\lambda = 1011$ nm) are enough to split one water molecule into H_2 and oxygen.¹⁶ Actually, the energy of an electron–hole pair after photon absorption is internal energy and not free energy. The contribution of entropy must be taken into account. The entropic factor reduces the range of solar wavelengths that can be effective in promoting water splitting to values above 610 nm.¹⁷ Other kinetic factors affect the efficiency of the solar-induced water splitting performed with a heterogeneous photocatalyst.¹⁸ For instance, activation barriers might hinder the migration of the photo-generated charges toward the surface of the solid particles. These barriers can be lowered by synthesizing the semiconductor photocatalysts in the form of nanomaterials having high crystallinity (reducing the internal defects that can act as traps of the photogenerated charges). Moreover, the superficial redox reactions might show overpotentials. This latter issue can be circumvented by loading the photocatalyst with co-catalysts for both the reductive and oxidative steps.^{19–21} Alternatively, the over-potentials can be overtaken if the electric potentials associated with the CB and VB are highly negative and positive, respectively (see Fig. 1B). This choice imposes large BGs on the semiconductors, which become capable of absorbing limited portions of the solar spectrum, almost exclusively restricted to the UV region.^{22–24} A strategy to avoid this blind alley relies upon the use of sacrificial reagents²⁵ when the only water-splitting product that is of interest is the energy vector H_2 (and O_2 is just a by-product). A sacrificial reagent (SR) in the photo-induced hydrogen production from water is any species that can be oxidized more easily than the oxygen of water. Usual SRs are amines, alcohols, carboxylic acids, sulphide and sulphite salts. A farsighted choice consists of using an easily available biomass compound as a sacrificial reducing agent, guaranteeing the possibility of implementing a large-scale sustainable production of H_2 .¹⁶ Whenever a SR is used, the BG of the semiconductor can be shrunk, allowing the capture of a broad portion of the solar spectrum: the only strong thermodynamic requirement is regarding the CB electric potential which must be more negative than the couple H^+/H_2 (see Fig. 1C).

Strategies to optimize the photocatalytic process of H_2 production through water splitting

Engineering the BG of the photocatalyst. The heterogeneous photocatalysts employed so far are almost exclusively metal oxides, sulphides, and nitrides. In these semiconductors, the lowest energy levels of the conduction bands are attributable to empty metal d- or sp-orbitals, whereas the highest energy levels of the valence bands are mainly made of O 2p, N 2p or S 3p orbitals. Generally, metal sulphides and nitrides have smaller band gaps than metal oxides. However, they are more susceptible to collateral photo-corrosion reactions in the water-splitting process. In the literature,¹⁸ there are various examples of metal oxides with broadband gaps, which are capable of photo-inducing the water-splitting reaction. Usually, they have metal cations with d orbitals completely empty (d0), such as



Ti^{4+} , Zr^{4+} , Nb^{5+} , Ta^{5+} , and W^{6+} , or metal cations that have the d orbitals completely full, but empty sp orbitals, such as Zn^{2+} , Ga^{3+} , In^{3+} , Ge^{4+} , Sn^{4+} , and Sb^{5+} . The major drawback of these metal oxides as photocatalysts is their colour: they are white powders that do not absorb visible light. Therefore, three principal strategies have been devised to synthesize colored metal oxides: one is based on doping; another is the extension of their VB, and the third one is the synthesis of solid solutions.

1. Doping: Doping of metal oxides is usually carried out with sulphur or nitrogen atoms or suitable metals. The doping metals must have partially filled d-orbitals (such as Ni^{2+} , Co^{2+} , Fe^{3+} , Mn^{2+} , and Cr^{3+}) or partially filled sp orbitals (such as Sn^{2+} , Bi^{2+} , Pb^{2+}). The dopants introduce electron states in between the VB and the CB, making the material capable of absorbing visible light. However, the dopants are introduced in small percentages. Therefore, the number of visible photons that can be harvested is always limited.

2. Extension of the VB: The extension of the metal oxide VB is feasible by synthesizing metal oxy-nitrides and oxy-sulphides. The introduction of nitrogen or sulphur atoms in amounts comparable to that of oxygen enlarges the top of the VB towards less positive electric potential values. The same effect can be achieved by stoichiometric addition of metal ions having partially filled sp orbitals, such as Sn^{2+} , Bi^{2+} , or Pb^{2+} .

3. Solid solutions: The mixture of white and colored oxides produces colored solid solutions that are good photocatalysts.²⁶ Their synthesis is feasible when acidic and basic oxides are intermixed. Acidic and basic oxides are acceptors and donors of oxide anions, respectively. The acid–base character of a metal oxide depends on the electronegativity of the metal: the more electronegative the metal, the more acidic the metal oxide. The mixture of different metal oxides can give rise to new crystal structures that can be more active as photocatalysts than their parent phases.

Co-catalysts. The presence of co-catalysts plays a crucial role in enhancing the efficiency of the photocatalytic system due to a variety of factors.^{19,20,22–24,27} In a heterogeneously photocatalyzed water-splitting process, a co-catalyst is a solid insoluble compound deposited onto the photocatalyst solid surface capable of assisting the redox reactions in different ways, which can be summarized as follows: (i) enhancing light harvesting properties; (ii) facilitating the charge separation; (iii) adding new catalytically active centres other than those of the photocatalyst, and overall, (iv) reducing the overpotential. Co-catalysts used for both the hydrogen evolution reaction (HER) and oxygen evolution reaction (OER) are commonly based on noble metals such as Pt, Au, Ir and Ru. They exhibit peculiar features in offering alternative and low-energy reaction pathways to both photo-generated electrons and holes to reduce H^+ to H_2 and oxidize O^{2-} to O_2 , respectively. Despite their high efficiency, the main drawback of noble metal-based co-catalysts is their employment in large-scale systems, owing to their high cost and scarce earth abundance. Two strategies are mostly pursued to overcome this problem: (i) search for efficient non-noble metal co-catalysts,²⁸ (ii) minimize the amount of noble-metal atom utilization. As far as the strategy (ii) is concerned,²⁹ it can be accomplished by the exploitation of

molecular catalysts, under the assumption that all metal centers are catalytically active, different from what happens in heterogeneous catalysts.³⁰ Otherwise, it is conceivable to anchor a well-defined molecular catalyst onto the surface of a suitable support, thus obtaining a heterogenized catalyst, combining the positive aspects of homogeneous and heterogeneous catalysis.³¹ Finally, the minimization of metal atoms can be achieved by diluting them in a proper material, having features that maximize metal-accessibility and performance.³²

Transition metal-based compounds, supported on a semiconductor surface, have proven to provide good results for the efficiency of the system.^{33–35} Due to their coloured features, transition metals can help in improving the light-harvesting properties of the system by adding a larger portion of the visible spectrum absorbed and helping the photo-induced charge separation. In addition, the formation of heterojunctions with the photocatalyst surface effectively promotes the charge separation lifetimes by means of two known processes: the formation of Schottky barriers which are electric fields generated by the co-catalyst and placed under the CB of the photocatalyst, which are able to inhibit the backflow of photo-generated electrons, resulting in more electrons to participate in the reaction of H_2 evolution. The second process concerns the semiconducting properties of the co-catalyst, resulting in a band alignment of the CB at lower energy with respect to that of the photocatalysts, thus promoting electron migration upon light excitation. For the oxygen evolution side, the same process can occur with co-catalysts having the VB placed under the VB of the co-catalysts, promoting the hole-migration from photo-to the co-catalyst through the junction interface. In the literature, the most employed co-catalysts for the water splitting reaction based on transition metals belong to several families of compounds such as elemental metals, metal sulphides, nitrides, carbides, borides, phosphides, oxides, hydroxides and oxo-hydroxides.^{36–45} The pure solar-to-hydrogen process, which is solely based on the use of solar radiation without any external current supply or mediated by electrolyte solutions, is limited by the efficiency, being around 3%. As an example, a recent study reported a target value of about 9% by using concentrated solar irradiation and Ga/In nitride nanoparticles as a co-catalyst.⁴⁶

The efficiency of co-catalysts in assisting the photocatalytic process is also related to other relevant factors:

- The microstructural features such as the crystal size distribution, the shape of the crystals and the active facets exposed, the presence of microstructural strain and local defects which can affect the charge separation process;
- The deposition methods such as photoassisted deposition, chemical vapour deposition/sputtering or impregnation with a solution containing the catalyst precursors;
- The dispersibility of the co-catalysts in low boiling solvents able to make homogeneous layers of nanometric size. Some recent papers reported that the use of nanometric-sized crystals, homogeneously distributed and deposited by physical methods such as vapor deposition or photo-deposition, are more effective in inducing stable and durable charge separation and achieving highly homogenous deposition of photo/co-catalysts.^{47,48}



Up-conversion. After engineering the BG of the photocatalyst and introducing co-catalysts, an alternative approach to reduce the waste of low-frequency solar photons is the up-conversion process. Up-conversion allows the deformation of the shape of the solar spectrum because it basically converts two or more low-energy photons into one high-energy photon.⁴⁹ In other words, the up-conversion allows the solar spectrum to be blue-shifted by producing more photons that can be absorbed by the photocatalysts and increase the yield of H₂ production. The up-conversion of the incoherent solar radiation can be achieved through different mechanisms, such as (1) sequential ground state/excited state absorption, (2) energy transfer processes, (3) cooperative processes, (4) photon avalanche, and (5) sensitized adiabatic photoreactions.¹⁶ These five mechanisms require chemical compounds that give rise to metastable electronic excited states, having lifetimes from the micro-second to the milli-second domains. The metastable electronic excited states allow the population of higher levels to be achieved through “ladder-climbing” events based on sequential absorptions and thermodynamically spontaneous or entropy-driven energy transfer processes.^{50,51} After climbing the excited states, the sudden and emissive jump from a very high excited state to the electronic ground state is a source of photons having frequencies higher than those previously absorbed. Rare earth and transition metal ions are the right candidates for obtaining up-conversion through the mechanisms (1), (2), (3), and (4), mentioned above.⁵² They are usually embedded within host matrices having small phonon energies to reduce the probability of non-radiative dissipation processes and prolong the electronic excited states' lifetimes. From this point of view, metal chlorides are the most promising host materials.⁴⁹ The principal drawback of rare earth and transition metal ions is their limited absorption power: their absorption spectra are usually sharp bands harvesting limited portions of the solar spectrum. This limitation can be circumvented by turning to another mechanism for achieving up-conversion: the triplet-triplet annihilation (TTA) process.⁵³ Generally, TTA involves two chromophores: a coordination compound, which shows broad absorption bands and a high triplet quantum yield, and an organic compound with an almost unitary fluorescence quantum yield. The coordination compound is the sensitizer that absorbs low-energy photons and transfers their energy to the fluorescent compound. Ultimately, the fluorophore becomes a source of high-frequency photons after an annihilation of triplet states. The main hindrance in achieving the TTA-based up-conversion comes from the atmospheric O₂, which quenches the metastable triplet states of the coordination and organic compounds. An effective strategy to limit the quenching effect of O₂ is to dissolve the sensitizer and the fluorophore in solvents showing a limited solubility to oxygen and embed these solutions within protecting silica capsules.^{54,55}

The sonolysis

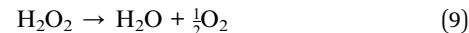
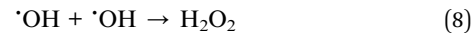
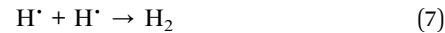
Sonolysis is a process that involves the splitting of molecules within a medium due to the mechanical stimulation caused by the presence of acoustic waves, typically at ultrasonic

frequencies ($f > 20$ kHz).^{56–58} Generally, sonolysis is strictly related to cavitation phenomena in the literature,^{59–61} with bubble generation in the fluid, and it is associated with emission of ultrasonic frequencies from a specific range, known as power ultrasound, with frequencies ranging from 20 kHz to 1 MHz, and a minimum power of 10 W, that can contribute to sonochemical effects.⁵⁶ When ultrasound propagates through a liquid solution, it gives rise to various phenomena:

- Heating of the solution occurs because of the dissipation of mechanical energy and atomization effects.⁵⁹ This can lead to the formation of acoustic fountains at the gas–liquid interface, resulting in localized temperatures of up to 250 °C;⁶²

- Small bubbles are generated, expanding during the pressure rarefaction phases caused by the oscillation of the ultrasound waves. Subsequently, these bubbles collapse during compression, giving rise to cavitation phenomena.⁶⁰ Cavitation events can reach extremely high temperatures of up to 5000 °C and pressures of 2000 atm.⁶³

In the case of water, the sonolysis process involves a series of reactions:⁶⁴



Eqn (6) shows that water sonolysis produces two reactive radical species: H[·] and OH[·]. Then, two H[·] can combine to produce molecular hydrogen (eqn (7)), and two OH[·] can produce hydrogen peroxide (eqn (8)). H₂O₂ is unstable and spontaneously decomposes into water and O₂ (eqn (9)). Finally, H[·] and OH[·] can recombine to form the reagent water molecule (eqn (10)).

Cavitation and sonolysis processes for hydrogen production⁶¹ became popular in recent times because they represent an innovative way for green-hydrogen production (the so-called son-hydro-gen process⁶⁵). Ultrasound power is reported as a conversion process with an overall efficiency of 80–90%, considering the electrical power input at the transducer stage, and a maximum potential loss of 10% for heat at the generator and transducer power conversion respectively.⁶⁵

Combining the state of the art about sonolysis for sono-hydro-gen processes and more general sono-chemical applications, it is possible to list the main parameters which may influence the hydrogen production by sonolysis in water-filled sono-reactors (Fig. 2).

Recent studies provided useful equations to calculate the cavitation energy E_{cav} in terms of input power W_{input} to a sono-trde (type-A sono-reactor, more information below):^{66,67}

$$E_{\text{cav}} = -8 \times 10^{-4} \times W_{\text{input}}^2 + 0.4699 \times W_{\text{input}} \quad (11)$$

And the corresponding sono-chemical energy consumption is:⁶⁶



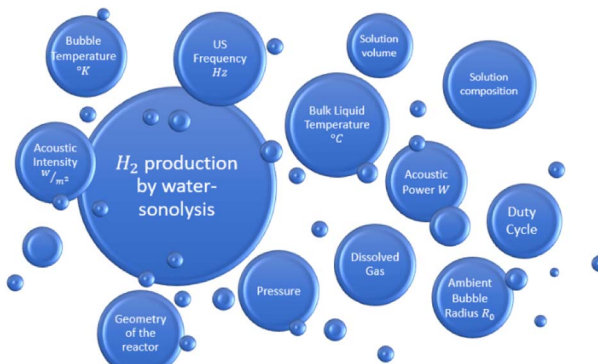


Fig. 2 Factors affecting hydrogen production by water-sonolysis.

$$\eta_{\text{Sono-hydro-gen}} = 3 \times 10^{-4} \times E_{\text{cav}}^2 + 0.014 \times E_{\text{cav}} \quad (12)$$

And the Blake cavitation threshold P_{BL} to individualize cavitation activity in a fluid where static pressure P_0 , water vapor pressure P_v , and initial bubble radius R_0 are known, following the equation:^{66,68}

$$P_{\text{BL}} = P_0 - P_v + \frac{2}{3\sqrt{3}} \left(\frac{2\delta}{R_0} \right)^{\frac{3}{2}} \left(P_0 - P_v + \frac{2\delta}{R_0} \right)^{-\frac{1}{2}} \quad (13)$$

where δ is the surface tension of water. However, eqn (11)–(13) describe a validated method only for a specific kind of sono-reactors and, at the moment, they cannot be considered for a general sonochemistry set-up. To provide a more complete analysis and potentiality of the process, the factors shown in Fig. 2 and their influence on the cavitation, or the sono-hydrogen yield can be discussed point-by-point.

The yield as a function of gas in the reactor. The influence of the gas dissolved in the sono-reactor has been widely studied in the general sonochemistry literature, but the comparison between different gases was not consistently performed in the sono-hydro-gen literature until recent time.^{65,69–72} In general sonochemistry literature, the argon⁷² atmosphere shows an improvement in the cavitation yield in the fluid compared to oxygen O_2 (0.6×10^{-10} vs. 0.5×10^{-10} mol J^{-1} with 20 kHz sonotrode excitation⁷³) and air (with null values under the same conditions⁷⁴), by potassium iodide KI oxidation process⁷⁵ measurements, even if it is not the theoretical best enhancer of cavitation activity. Indeed, the carbon dioxide CO_2 atmosphere experimentally reports the largest production of bigger bubbles in water⁷² due to its higher solubility, heat capacity C_p , and maximum bubbling pressure (or mean driving pressure) compared to other experimented gases (Table 1,^{70,73,76}), enhancing bubble nucleation within the liquid exposed to ultrasound, but engendering a recombination mechanism of C atoms to the set of free radicals in the medium, generating species such as HCO , COOH , or CH_2O ,⁷² reducing the recombination of hydrogen molecules.

Furthermore, no sonochemical products are detected in a pure CO_2 atmosphere.^{72,74}

However, Ar should be considered as a promising dissolved gas in reactors for sono-hydro-gen processes and numerical

work by Merouani *et al.*^{69,70} in dynamics for hydrogen production may prove it: applying the Keller–Miksas equation for cavitation and considering pressure and temperature by the ideal-gas law or van der Waals equation, the authors reported that argon gas shows a hydrogen production rate higher than natural air because of its greater polytropic ratio ($\gamma_{\text{Ar}} = 1.66$ vs. $\gamma_{\text{Air}} = 1.41$) and lower thermal conductivity ($\lambda_{\text{Ar}} = 0.018 \text{ W m}^{-2} \text{ K}^{-1}$ vs. $\lambda_{\text{Air}} = 0.024 \text{ W m}^{-2} \text{ K}^{-1}$), causing bigger bubble radius, higher bubble temperature (6000 K with the van der Waals model⁷¹), and more violent explosion with a higher production of radicals from water. They reported a production of H_2 61 times higher than that obtained in the air atmosphere ($13.6 \mu\text{mol min}^{-1}$ vs. $0.22 \mu\text{mol min}^{-1}$) in a type-A sonoreactor (Fig. 3) at 1 MHz.⁷⁰

The yield depends on the geometry of the sonoreactor, acoustic power, and the number of sonotrodes. The geometry of the sonoreactor and its influence on the sonochemical process is a widely investigated topic. Starting from the three main sonoreactor schemes (Fig. 4), researchers performed numerical studies and experimental measurements on a wide range of cylindrical or rectangular base reactors. For a type-A sonoreactor,⁶² considerable differences in the acoustic pressure distribution are reported with an apparent advantage to the bottom-curved shape geometry at 20 kHz by numerical simulations in COMSOL Multiphysics.⁶⁶ At the same frequency comparable results were obtained by Wei⁷⁷ by simulations, but highlighting the impossibility to implement the bi-phase liquid section represented by the bubbles and the acoustic scattering provided by the bubble clusters as the main limitations to obtain reliable simulated acoustic pressure distributions. Consistently, Wang⁸⁰ confirmed the potential of the sound wave-based propagation simulation approach but highlights lacking in the prediction of the influence of acoustic scattering on pressure distribution, and due to the lack of proof about the cavitation structure, high pressure zones cannot provide data to forecast cavitation activity. Cavitation structures are multiple, quite different from each other, and strictly related to the acoustic pressure P compared to a reference pressure $P_0 \equiv 190 \text{ kPa}$.⁷³ If $P > P_0$, because of primary Bjerknes forces, the cavitation evolves along the pressure lines arranging in different shapes as reported in Table 2, conversely, if $P < P_0$, cavitation evolves along the pressure antinodes. Even the presence of a metal net in the fluid exposed to ultrasound influences the sound propagation by scattering, moving the CBS after the net (artificial conical bubble structure, ACBS).

Correlation between the reactor geometry and cavitation distribution (or structure) is crucial for choosing the reactors and prediction of results by simulation. In multiple studies,^{65,66,73,77–80} there is a reference to the resonance of the acoustic field in the structure and its influence on the cavitation field. Nonetheless, many doubts arise about the real influence of eigenvalues and eigenvectors of geometries: ALF cavitation structure observation as a function of the shape of the reactor (Fig. 3) suggests that the influence of the geometry on the cavitation field is more related to the inclination of reflective surfaces rather than to the eigenvalues of the reactor geometry. Type-B reactors are less investigated. In rectangular shaped



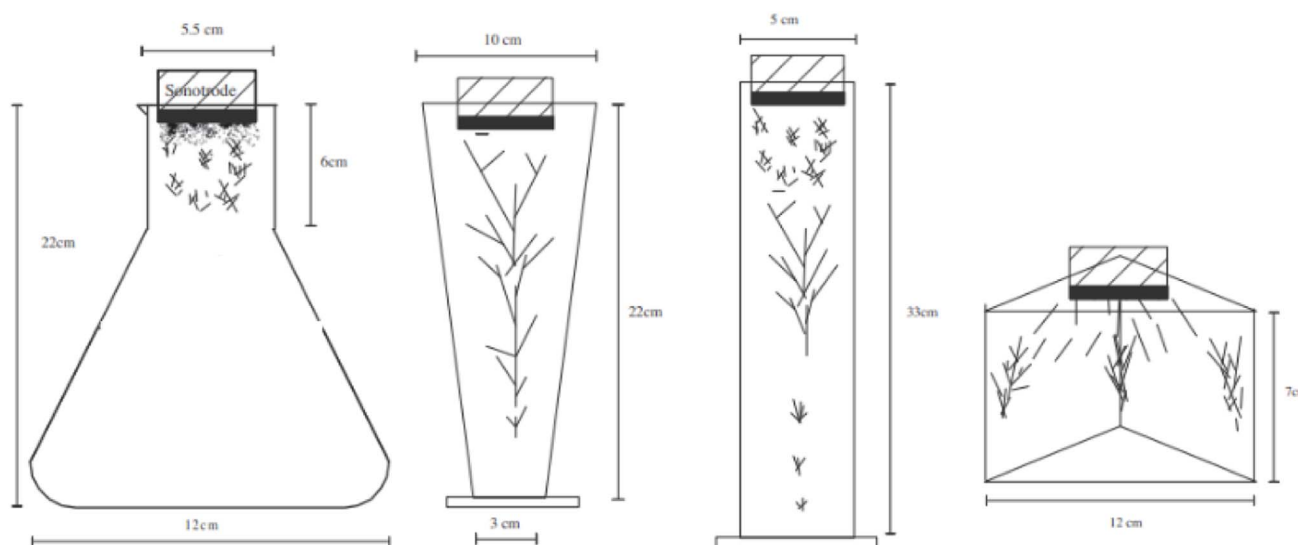
Table 1 Factors affecting hydrogen production by water-sonolysis

Gas	Solubility (mol mol ⁻¹) ⁷³	Maximum pressure [Pa] ⁷⁶	C_p (kJ kg ⁻¹ K ⁻¹) ⁷³
Helium He	7.0×10^{-6}	—	20.9
Argon Ar	2.7×10^{-5}	—	20.8
Nitrogen N ₂	1.5×10^{-5}	1.56×10^4	29.2
Oxygen O ₂	2.5×10^{-5}	1.62×10^4	29.4
Carbon dioxide CO ₂	7.1×10^{-4}	1.7×10^4	37.5
Air	1.524×10^{-5} (ref. 70)	1.61×10^4	—

reactors, Koch⁸¹ found a correlation between the double layer cavitation structure and the acoustic power of ultrasound but there were no geometrical correlations, and comparing simulations and pressure measurements, eigenvalue influence was not proved, and symmetrical distribution of the sound pressure was discussed.⁸²

The number of sonotrodes and the acoustic intensity applied by the transducer are key aspects investigated in the literature. Cavitation power is logarithmically related to the input power of the transducer,^{76,83} and increasing the electrical input power from 23 to 82 W, high values of the cavitation yield ($\sim 3 \times 10^{-7}$ mol W⁻¹ min⁻¹) can be achieved with larger liquid volume (to 0.3 L).⁸⁴ Input power allowed determination of the cavitation threshold in a rectangular type-B reactor. At 45 kHz, with an ELMA 90 W transducer, cavitation structures appear at 14.8 W,⁸¹ with a clear smoking structure at 30.1 W, and a complete double layer from 58.5 W, while in the type-A reactor at 20 kHz the acoustic intensity threshold for cavitation is reported at ~ 0.74 W cm⁻².⁷⁷ Numerically, Merouani *et al.*⁶⁹ showed that with acoustic intensity increasing from 0.5 to 1 W cm⁻², at different US frequencies, the maximum bubble temperature increases reporting an higher hydrogen production rate especially at higher US frequencies (over 1 MHz, on doubling the acoustic intensity, the production rate is 10^{10} higher, while at 20 kHz it is 10^2 higher).

Impedance matching tests were performed to enhance the cavitation energy conversion by Rooze *et al.*,⁷³ measuring in air at 20 °C the US horn using a Hioki 3532-50 impedance analyzer and selecting three frequencies (20–41–62 kHz) as the frequencies with a lower impedance reporting a US-electric yield of 0.46 ± 0.03 , 0.17 ± 0.01 , and 0.21 ± 0.02 respectively, and consistently with the impedance values being measured. Regarding the number of sonotrodes in a type-A reactor,⁶² the cavitation percentage over the sonoreactor volume based on Blake cavitation analysis⁶⁸ is higher for a 3 sonotrode configuration irradiant 20 kHz US in a rectangular reactor with 61.1% of cavitation.⁶⁶ A higher number of sonotrodes for unit volume in a type-A sonoreactor does not provide a higher cavitation volume. The calculated conversion for one sonotrode logarithmically decreases while input power increases with an extreme 44% of energy conversion at 36 W input power to 1% at 580 W. Impedance matching test were performed to enhance the cavitation energy conversion by Rooze *et al.*,⁷³ measuring in air at 20 °C the US horn using a Hioki 3532-50 impedance analyzer and selecting three frequencies (20–41–62 kHz) as the frequencies with a lower impedance reporting a US-electric yield of 0.46 ± 0.03 , 0.17 ± 0.01 , and 0.21 ± 0.02 respectively, and consistently with the impedance values measured. Consistently, the energy conversion efficiency decreases, but a flatter trend is

Fig. 3 ALF structures in different reactors ($P < 180$ kPa).⁵⁷

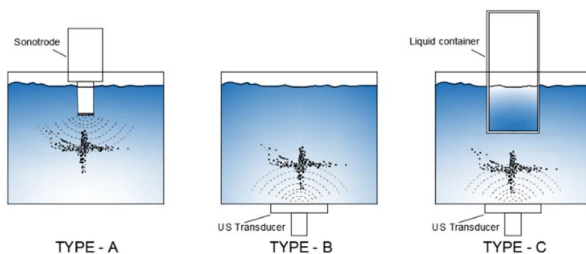


Fig. 4 Common configuration of a sonoreactor.

reported with a decrease of only $\sim 10\%$, from $\sim 43\%$ to 33% from 1 to 5 sonotrodes with each of power 36 W.

The yield as a function of temperature of the bubble and of the solution. Thermal interaction between the cavitation bubble and the fluid has been defined as null, and the surface of the bubble has been defined adiabatic.⁶⁵ In Merouani *et al.*⁷⁰ bubble temperature (BT) during cavitation was numerically calculated as a function of acoustic intensity, frequency, and bubble radius growth ratio, reporting that higher BT (3923–6063 K from 0.5–1 W cm⁻² at 20 kHz) corresponds to more violent bubble explosions, a higher water vapor trapped and a higher hydrogen yield from water cavitation, especially in an argon Ar atmosphere (Theme 1). The BT is inversely proportional to the liquid temperature (from 6063 K at 20 °C to 2255 K at 50 °C), with a corresponding decrement of the hydrogen yield. With the van der Waals's temperature evolution expression into the cavitation numerical simulation model, Kerboua *et al.*⁷¹ confirmed the increase in hydrogen production yield with BT increment under the best temperature conditions at 6000 K. Gogate *et al.*⁷⁶ experimentally and numerically confirmed the yield variations in potassium iodide sonochemical processes as a function of the solution temperature, with a decrease of 80 ppm (from 198.25 to 115.00) of iodine liberated with a temperature increment of the solution from 10° to 30 °C.

The yield as a function of transducer duty cycle. The duty cycle of the transducer is not studied specifically for the production of hydrogen but it was investigated more generally in sonochemical processes by Gogate *et al.*⁷⁶ The authors compared different ranges of duty cycles from 20 to 100% in a time period of 50 s. Surprisingly, the best sonolysis yield is reported at 60%, with the highest iodine liberated as a function of duty time (Table 3) while 100% was not recommended due to the maintenance problem of the transducer.

These are promising results that may be applied to hydrogen production as well.

Table 2 Structure of the cavitation field as a function of the acoustic pressure: Conical bubble structure (CBS), artificial conical bubble structure (ACBS), acoustic Lichtenberg figures (ALFs), smokers, web, tailing and jet bubble structures (TBS and JBS).⁸⁵

	CBS	ACBS	ALF	Smokers	Web	TBS	JBS
$P > P_0$	X			X		X	X
$P < P_0$		X	X		X		

The yield as a function of frequency. Considering the wide range of frequencies reported as power ultrasound,⁵⁷ investigations on the best frequency is an important step to enhance the cavitation efficiency and the potential sono-hydro-gen production. Generally, the computational work by Kerboua,⁷¹ and Merouani *et al.*⁷⁰ reported that by increasing the US frequency the production rate in mol s⁻¹ decreases by order of magnitude doubling the frequency, and this is valid for the single bubbles as a function of different acoustic intensities, gas percentage, and bubble temperatures (Fig. 6). The reason for this decrease can be numerically attributed to the ever-smaller bubble radius before explosion reported with the increase of the US frequency. In experimental measurement conducted in sonochemistry, strategies to find the best frequencies were optimized finding the best frequency as a function of impedance matching between the transducer and fluid or using the first longitudinal modal frequency. Rooze *et al.*⁸⁶ obtained a higher cavitation yield at the frequency with the lowest impedance after the impedance matching test. However Wang⁸⁰ performed US cavitation with the emitted frequency of the first longitudinal modal response. Kerboua *et al.*⁷⁹ numerically investigated the US frequency harmonics on the cavitation yield starting from a sinusoidal signal at 200 kHz, and the first five harmonics, reporting discrete proportionality between the cavitation bubble radius and frequency. Nonetheless, more investigations are required to better define the contribution of longitudinal modal frequencies, whereas the impedance matching test represents a reliable method.

The yield as a function of ambient pressure and solution composition. Few studies were performed to assess the influence of the ambient pressure within the reactor on the cavitation yield. The experimental study conducted by Cotana *et al.*,⁸⁷ reported a decrease in the hydrogen production rate when the ambient pressure increased. Moving from 1 to 2.5 atm, the hydrogen production decreased by 50% in 5 h (from 12 to 6 μmol , Fig. 5). Rashwan⁶⁵ justified this as the impossibility of free oscillations by cavitation bubbles, reducing the amount of heat absorbed, and affecting hydrogen production.

The composition of the solution exposed to ultrasound has been explored as a possible way to enhance the sonolysis by introducing a sacrificial reagent, a chemical species that can be oxidized more easily than oxygen of water, reducing the recombination tendency of electrons and holes and accelerating the rate of hydrogen generation, into the solution.⁸⁸ Bioethanol, a sugar fermentation biomass compound, was tested as a sacrificial agent in 300 mL of distilled water with a 20%

Table 3 Effect of sonotrode duty cycle on iodine liberation

Duty cycle (%)	On time (s)	Off time (s)	Iodine liberated (ppm)
20	10	40	78.75
40	20	30	108.25
50	25	25	141.25
60	30	20	171.00
80	40	10	200.25



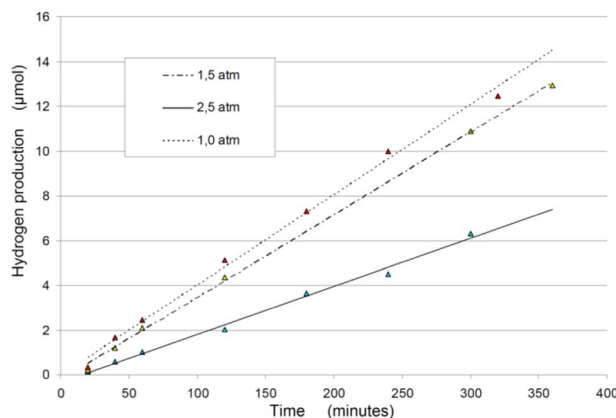


Fig. 5 Hydrogen production rates during exposition time, changing the reactor inner pressure.⁸⁷

volumetric concentration, and the results were compared with sheer water and sheer bioethanol tests.⁸⁹ The presence of bioethanol in water solution resulted in a 40% improvement of the hydrogen production rate compared to sheer water solution (112 ± 3 vs. $80 \pm 2 \mu\text{mol h}^{-1}$). Sheer bioethanol solution reported a lower production rate ($5.5 \pm 0.2 \mu\text{mol h}^{-1}$). The concentration of methanol in the solution for enhancing the sono-hydrogen production has been investigated in the literature.⁹⁰ Henglein and Schultz reported that iodide oxidation by ultrasonic cavitation proportionally decreases with the increase of the methanol concentration in the solution due to sono activity inhibition above a certain concentration (5 mol in 30 mL solution) upon increasing temperature of the solution between 5 and 20 °C. But at 40 °C the presence of methanol did not influence the oxidation anymore. Büttner *et al.*⁹¹ confirmed the effect of methanol⁸³ increments of all species production in a 10% methanol–water solution. Theoretical studies reported the optimal methanol concentration for hydrogen production in water solution as 20%⁷⁹ with an 80% argon concentration in the atmosphere, with an increase of hydrogen production of 4.7 times compared to a sheer water solution.⁹⁰

The sono-photolysis and its synergistic effect

The combined action of solar radiation and ultrasound can exert a synergistic effect on some chemical reactions carried out in the liquid phase in the presence of a heterogeneous photocatalyst. Sono-photolysis was first employed for the degradation of water pollutants, for instance, phenols as reported by Wu and co-authors in 2001 in the absence or in the presence of a catalyst (FeSO_4) using UV irradiation assisted by 30 kHz ultrasonication. The percentage of degraded phenols was slightly higher than that obtained by simple UV irradiation both in the presence or in the absence of dissolved Fe^{3+} ions.⁹² In 2010 a more pronounced synergistic degradation effect of UV/US irradiation (25 kHz, 1 kW) of phenolic solution in the presence of TiO_2 as photocatalyst and H_2O_2 was also reported.⁹³ Other important applications of catalysed sono-photolysis deal with the degradation of pollutants. In 2011

Cui *et al.* investigated a continuous process involving US (28 kHz, 300 W electric power)/UV (254 nm, 30 W, TiO_2) combination, for the degradation of low concentration of methyl orange in water (25 mg L^{-1}).⁹⁴ In a recent paper the degradation of carmoisine was carried out by employing $\cdot\text{OH}$ radicals generated by photolysis and transient cavitation.⁹⁵ A similar process was also reported in 2023 for the sono-photodegradation of carbaryl, a pollutant pesticide, by using a $\text{Ln}@\text{MOF}$ as photocatalyst.⁹⁶ Other organic pollutants as dyes were efficiently degraded by the synergic effect of photo-sonolysis by using $\text{CuO}@\text{TiO}_2/\text{rGO}$ (reduced graphene oxide rGO) as photocatalysts in which the coupled action of light and ultrasound enhanced the degradation kinetics by over 3 times.⁹⁶

Other examples of the use of sono-photolysis as an advanced oxidation process (AOP) for dye degradation, wastewater treatment and for biomass valorization have been also reported in a 2017 concept paper by Chatel and co-authors.⁹⁷

The synergistic effect of sonophotolysis for the water-splitting reaction is a promising field of research although it is still scarcely investigated.

The synergistic effect of solar light and ultrasound in the production of H_2 from water is usually quantified through the equation below:^{16,89,98–100}

$$\text{SynEf} = \frac{k_{\text{hv+US}} - (k_{\text{hv}} + k_{\text{US}})}{k_{\text{hv+US}}} \quad (14)$$

In (14), k_{hv} , k_{US} , $k_{\text{hv+US}}$ are the kinetic constants of H_2 production when simply electromagnetic radiation (hv) or simply ultrasound (US), or both light and ultrasound are applied, respectively. Whenever $\text{SynEf} \neq 0$, there is a synergistic effect.

A recent paper reported on the synergistic and highly beneficial effect of ultrasound irradiation in visible light water photolysis using a Co-phtalocyanine dye, namely Therephthal, as a homogeneous photocatalyst. The effect of ultrasound irradiation in the presence of light resulted in a large increment of oxygen evolution from about 35 (in the absence of ultrasound) to over 1400 μmol after 3 h of US/light irradiation. The authors proposed as a mechanism the US enhanced formation of ROS species such as ($\cdot\text{OH}$, $\cdot\text{OOH}$, $\text{O}_2^{\cdot-}$) able to easily oxidize Co^{2+} to Co^{3+} which was mainly responsible for the oxygen evolution reaction.¹⁰¹ In 2017 Singh and Sinha reported the catalyzed sono-photolysis of water for the HER using CdS as a photocatalyst. The higher activity of H_2 production of horn US connected to a piezoelectric transducer working at 20 kHz and 40 W, with respect to the pure UV irradiation was attributed to a faster removal of bubbles of hydrogen from the surface of photocatalyst due to the mechanical energy associated with ultrasound able to overcome surface tension.¹⁰² The different aspects of the synergy between light and ultrasound still need to be completely understood. It can be due to different factors. Some of them were recently discussed in a review by Wang and co-authors.¹⁰¹

It is evident that the two energy sources play a reciprocal positive feedback effect (see Fig. 7).



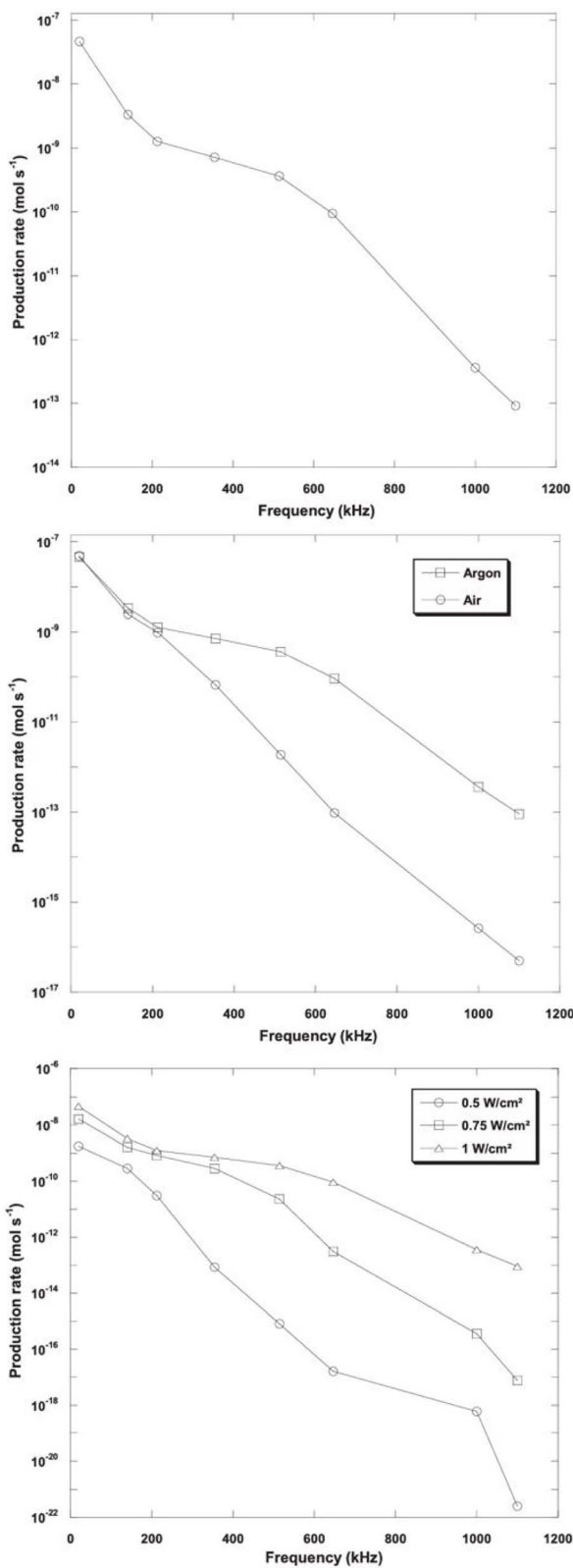


Fig. 6 Hydrogen production as a function of frequency, atmosphere and intensity.⁷⁰

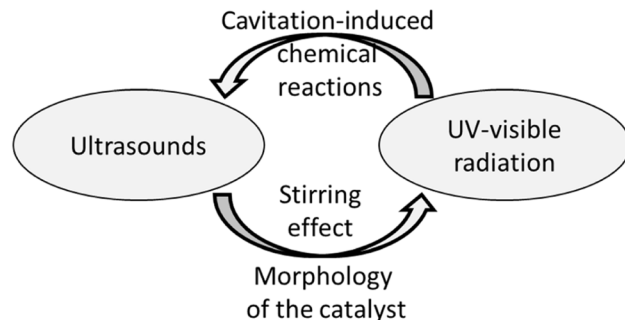


Fig. 7 The reciprocal positive feedback action between the two energy sources. The ultrasound affects the morphology of the heterogeneous catalyst and exerts a stirring effect that favours water splitting. On the other hand, the UV-visible radiation produces chemical species that provide extra nuclei for bubble formation, promoting the cavitation phenomenon.

It is well-known that acoustic cavitation induces the miniaturization of the catalyst's solid particles and an overall increase of the superficial area of the material. Hence, the semiconductor becomes a better photocatalyst.

Furthermore, the ultrasound waves exert a stirring effect in the reaction medium, favouring the physical contact between the liquid and solid phases. Moreover, the stirring effect lifts the catalytic solid particles from the bottom of the reactor up to the top part of the solution, facilitating the capture of light by the catalyst. On the other hand, the chemical species generated by photo-induced water splitting might provide extra nuclei for bubble formation, promoting the cavitation phenomenon. These concerted mechanisms may also affect the electronic structure of the catalyst. In particular, the collapse of cavitation bubbles could produce acoustic luminescence, thus promoting the migration of electrons from the valence band (VB) to the conduction band (CB). In contrast, the holes (h^+) remain in the VB. Consequently, the photogenerated electron–hole pairs react with the system's oxidant to generate reactive oxygen species (ROS), similar to those created by the photocatalytic process.

An additional synergistic effect by ultrasound and photocatalysis combination is reported in the presence of piezoelectric photocatalysts. Besides the photocatalyst properties previously explored, these photocatalysts generate periodic electrical potential into the solution by interaction with ultrasound, resulting in piezoelectric material surface charging.¹⁰³ Those potentials induce flexures of the energy band, enhancing the photogenerated charge separation.¹⁰¹ Furthermore, electronic jumps occur in the VB and CB of these materials providing additional electron–hole sources for the flow separation and transport.¹⁰⁴ Meng *et al.*¹⁰⁵ showed 1.32-fold increase of methyl orange photodegradation by Pt/BaTiO_3 heterojunction fabrication for piezoelectric enhanced photocatalysis, and Liu *et al.*¹⁰⁶ demonstrated 2.44 times higher photocatalysis activity due to the presence of the $\text{BaTiO}_3@\text{ReS}_2$ Schottky heterostructure. Hydrogen production by piezoelectric-assisted sono-photocatalysis was investigated with the $\text{g-C}_3\text{N}_4$ photocatalyst with a 50% H_2 production rate improvement from water sonolysis ($12.16 \text{ vs. } 8 \mu\text{mol (g}^{-1} \text{ h}^{-1}\text{)}$) and $\text{SynEf} \sim 0.32$. Catalytic

Table 4 Effect of photocatalysts on the cinematic viscosity and acoustic absorption of the solution

Photocatalyst	Cinematic viscosity η_c	Acoustic absorption α	Molecular mass g mol ⁻¹	Synergistic H ₂ production improvement after 2 h μmol
LaGa _{0.5} In _{0.5} O ₃	0.0127	0.179	276.15	0.14 ± 0.01
La _{0.8} Ga _{0.5} InO ₃	0.0117	0.166	329.69	0.14 ± 0.01
S:La _{0.8} Ga _{0.5} InO ₃	0.0115	0.161	361.75	0.20 ± 0.01

activity and the piezoelectric field of deformation are positively correlated as well as ultrasound intensity and the catalyst deformation degree.

The presence of photocatalyst powder dispersed into the solution may also affect the ultrasonic field behavior. Indeed, in the literature it has been reported that the cinematic viscosity of the solution varies with a calculable change of the acoustic absorption of the medium.¹⁰⁰ Capillary viscosimeter measurements provided cinematic viscosity data of the solution reporting a variation of 1.7% (Table 4) when the photocatalyst was sulfur-doped. Applying the corrected Stokes formula for the volume viscosity, 3% variations of the acoustic property absorption of the solution were calculated (Table 4). This may be explained because of the decrease in the cinematic viscosity due to the increase in the density of the solution. Those parameters are proved to be inversely proportional to each other.¹⁰⁷ Although this is not a proper declination of the synergistic effect of the photo-sonoysis, and the production improvement is sensible only for a short time period (up to 2 h of US exposition), further investigations are needed to evaluate any beneficial effect of the acoustic absorption variations of the solution on sono-photolysis.

Finally, our experiments¹⁰⁸ have demonstrated that sonophotocatalysis for hydrogen production benefits from using metal oxide solid solutions, which are high-entropy materials.¹⁰⁹ High-entropy materials are expected to be good catalysts because they guarantee the possibility of activating many parallel reactive paths.

For this reason, high-entropy materials are enticing interests in energy conversion and storage.¹¹⁰⁻¹¹²

Conclusions/future perspectives

Sono-photolysis is a phenomenon which involves two different processes for hydrogen production by water splitting using ultrasound and solar radiation exposure of a water-based solution. This process may be the focus of a sustainable green hydrogen production reactor and a possible synergic alternative to electrolyzers. The scientific literature has highlighted the following points:

- The transparent covering of the photo-sono reactor should be made of quartz glass to allow for a broader transmission of light, particularly in the UV spectrum.
- The presence of dispersed photocatalysts in the solution and the use of sacrificial agents are necessary for the proper functioning of the photolysis process. The sulfur doping of catalysts significantly affects the photolysis yields, with up to 17-fold increase. Semiconductor catalysts, particularly S:Y_{0.8}Ga_{0.2}-InO₃, have experimentally shown the highest rates of hydrogen production (0.14 ± 0.05 $\mu\text{ moles (H}_2\text{) per h}$).
- The use of a bioproduct as a sacrificial reagent allows the production of H₂ and avoids the production of O₂, which can partially deplete the desired hydrogen through combustion reactions, and instead produce by-products (such as aldehydes or carboxylic acids) which remain in solution and more easily separated from H₂.
- Introduce capsules of up-converting materials to reduce the waste of low-energy photons.
- The hydrogen production through ultrasound (US) is sensitive to the chemical composition of the solution. Solutions containing 20% vol of water and ethanol exhibit production rates around 40% faster than solutions composed solely of distilled water.
- The volume of the solution and its geometry within the reactor, determined by the peak resonance points of the water volume exposed to the US frequency, significantly enhances the efficiency (up to 112 $\mu\text{ moles (H}_2\text{) per h}$ ⁸⁹).
- Atmospheric pressure conditions for the solution yield the best results in terms of efficiency for sono- and photo-sonoysis.⁸⁷
- Type A or B sonoreactors can achieve the best performances for low-frequency range ultrasounds, with a duty cycle transducer activity of 60%, in an Ar atmosphere to achieve the largest bubble dimensions at the highest temperature.

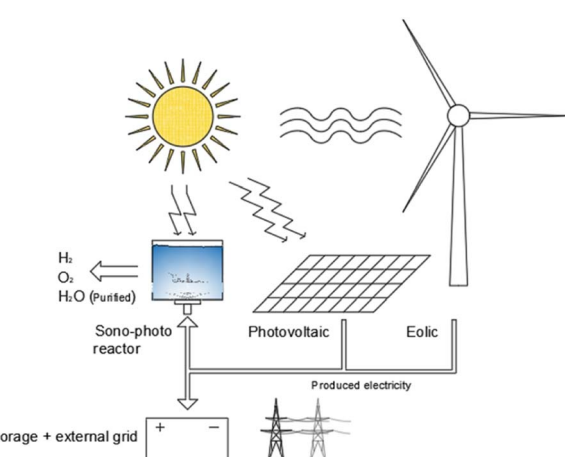


Fig. 8 Sono-photolysis technology integration in existing renewable production systems. The synergistic production of hydrogen, oxygen and purified water is highlighted as a key parameter for the application of this technology.



- Catalysts that reduce the kinematic viscosity of the solution enhance sonolysis yields by decreasing the solution's absorption coefficient.

- Sono-photolysis exhibits synergistic coupling between sonolysis and photolysis, resulting in efficiencies higher than the sum of the individual phenomena. This synergy is approximately 13%, and sulfur doping of catalysts has experimentally shown advantages in efficiency during the initial 3 hours of combined US and solar light exposure.

In envisioning the potential future advancements for sono-photolysis and reactor design, several key areas stand out when compared to the current state of the art. First, there is a growing focus on the environmental sustainability of sacrificial agents in the photolysis process,⁸⁹ with discussions in the literature highlighting bioethanol as a viable option. However, challenges arise as the catalysts explored often involve a 24-hour calcination process at 1373 K or the use of rare earth metals, both of which pose energy-intensive hurdles for large-scale photo-sonolysis applications. Another avenue for progress lies in enhancing photolysis efficiencies through the development of next-generation catalysts. Transition metal hydroxides and coordination polymers, particularly those based on Fe, Co, and Ni, are being explored. Simultaneously, there is an emphasis on extending the photolysis range within the solution by leveraging luminescent materials.¹¹³ Facilitating the electron promotion into the CB by US-activated intermittent electrical fields into the solution by dispersion of photo-catalysts with piezoelectric properties is another promising application that must be further explored. Refinement of the reactor geometry is also a key consideration, with an emphasis on maximizing solution volume based on resonance peaks.⁸⁹ This approach aligns with existing studies in the literature, aiming for optimal conditions for sono-photolysis. However, more investigations are required to predict the contribution of the vessel geometry compared to the modal excitations of the solution volume exposed to ultrasound. The literature⁷⁷ highlighted that the current impossibility of simulating the cavitation-bubbling evolution and spatial distribution precisely may compromise the usefulness of finite volume software approaches due to the absence of acoustic scattering of ultrasound waves by the bubbles. This is a factor which affects the spatial distribution of acoustic pressure into the vessel, rising differences between measurements and mechanical modal analysis.⁸² Furthermore, efforts are underway to investigate the energy efficiency between the mechanical energy induced by ultrasonication in the solution and the subsequent extraction of hydrogen. Although an energy cavitation efficiency in the solution up to 44% has been reported in the literature,⁶⁶ direct measurements of acoustic amplitude in the solution and the optimization of impedance matching between the reactor and the ultrasonic transducer are required. The potential expressed in contemporary literature¹¹⁴ in the production of oxygen through sono-photolysis (500 $\mu\text{mol h}^{-1}$) can only be welcomed as an extremely important point to be further investigated. It represents an additional synergistic effect of the described process, with clear application opportunities. However, the energy conversion of the whole hydrogen production system must be measured to further prove the

potentiality of the method, and to evaluate its margins for improvement. Lastly, efforts are focused on studying the synergistic interaction between photolysis and sonolysis to optimize and maximize hydrogen production efficiency. Researchers aim to uncover how these processes complement each other, seeking to identify conditions that enhance overall efficiency and hold promise for advancing hydrogen production technologies.

The sono-photolysis can play a promising role in the energy transition because of its transversal application potentiality, it can be implemented in existing processes such as water purification by ultrasonic cavitation,¹¹⁵ as well as it can be integrated into existing energy systems such as photovoltaic plants. Differentiating the energy production into electricity and hydrogen for energy storage purposes, sono-photolysis combination with existing energy production technologies systems may be a ready-to-use competitive hydrogen production application. Furthermore, combined with electrical production systems, the water purification process as a by-product of sono-photolysis can play a key role compared with existing hydrogen production technologies such as electrolyzers (Fig. 8), and it may be a synergistic effect extremely promising for applications in areas where there is a high solar radiation intensity magnitude and the necessity to provide off-grid energy and purified water for domestic use.

Conflicts of interest

The authors declare no conflict of interest.

Notes and references

- 1 J. Turner, G. Sverdrup, M. K. Mann, P. C. Maness, B. Kroposki, M. Ghirardi, R. J. Evans and D. Blake, *Int. J. Energy Res.*, 2008, **32**, 379–407.
- 2 D. B. Levin, L. Pitt and M. Love, *Int. J. Hydrogen Energy*, 2004, **29**, 173–185.
- 3 C. M. Kalamaras and A. M. Efthathiou, *Conference Papers in Energy*, 2013, **2013**, 1–9.
- 4 J. Ivy, *Summary of Electrolytic Hydrogen Production: Milestone Completion Report*, Golden, CO (United States), 2004.
- 5 A. Hauch, S. D. Ebbesen, S. H. Jensen and M. Mogensen, *J. Mater. Chem.*, 2008, **18**, 2331–2340.
- 6 S. E. Lindquist and C. Fell, *Encyclopedia of Electrochemical Power Sources*, 2009, pp. 369–383.
- 7 Ministero dello Sviluppo Economico MISE, *Strategia Nazionale Idrogeno – Linee Guida Preliminari*, Italy, 2020.
- 8 Gruppo di Lavoro Idrogeno, *Prime Indicazioni Per Una Strategia Italiana Ricerca Idrogeno (SIRI)*, 2020.
- 9 European Commission, *Communication From the Commission to the European Parliament, the Council, the European Economic and Social Committee and the Committee of the Regions a Hydrogen Strategy for a Climate-Neutral Europe*, 2020.
- 10 Council Regulation (EU Euratom) 2020/2093, *Laying Down the Multiannual Financial Framework for the Years 2021 to 2027*, 2020.

11 L. Baldinelli, G. Menendez Rodriguez, I. D. ' Ambrosio, A. M. Grigoras, R. Vivani, L. Latterini, A. Macchioni, F. De, A. Acde and G. Bistoni, *Chem. Sci.*, 2024, **15**, 1348–1363.

12 K. N. Liou, *Int. Geophys.*, 2002, **84**, 37–64.

13 J. E. Funk, *Int. J. Hydrogen Energy*, 2001, **26**, 185–190.

14 A. J. Bard and M. A. Fox, *Acc. Chem. Res.*, 1995, **28**, 141–145.

15 A. Kudo and Y. Miseki, *Chem. Soc. Rev.*, 2008, **38**, 253–278.

16 M. A. Basile, C. Carfagna, L. Dipasquale, A. Fontana, A. Gambacorta, G. G. d'Ayala, M. Malinconico and P. Cerruti, *Hydrogen Production: Prospects and Processes*, 2015, pp. 411–420.

17 J. R. Bolton, S. J. Strickler and J. S. Connolly, *Nature*, 1985, **316**, 495–500.

18 A. Kudo and Y. Miseki, *Chem. Soc. Rev.*, 2008, **38**, 253–278.

19 M. G. Walter, E. L. Warren, J. R. McKone, S. W. Boettcher, Q. Mi, E. A. Santori and N. S. Lewis, *Chem. Rev.*, 2010, **110**, 6446–6473.

20 S. Wang, A. Lu and C. J. Zhong, *Nano Convergence*, 2021, **8**, 1–23.

21 Z. Chen, W. Wei and B. J. Ni, *Curr. Opin. Green Sustainable Chem.*, 2021, **27**, 100398.

22 K. Maeda, *J. Photochem. Photobiol. C*, 2011, **12**, 237–268.

23 A. Kudo and Y. Miseki, *Chem. Soc. Rev.*, 2008, **38**, 253–278.

24 X. Chen, S. Shen, L. Guo and S. S. Mao, *Chem. Rev.*, 2010, **110**, 6503–6570.

25 V. Kumaravel, M. D. Imam, A. Badreldin, R. K. Chava, J. Y. Do, M. Kang and A. Abdel-Wahab, *Catalysts*, 2019, **9**, 276.

26 P. L. Gentili, M. Penconi, F. Costantino, P. Sassi, F. Ortica, F. Rossi and F. Elisei, *Sol. Energy Mater. Sol. Cells*, 2010, **94**, 2265–2274.

27 Z. Chen, W. Wei, L. Song and B. J. Ni, *Sustainable Horizons*, 2022, **1**, 100002.

28 X. Zou and Y. Zhang, *Chem. Soc. Rev.*, 2015, **44**, 5148–5180.

29 A. Macchioni, *Eur. J. Inorg. Chem.*, 2019, **2019**, 7–17.

30 F. Zaccaria, G. Menendez Rodriguez, L. Rocchigiani and A. Macchioni, *Front. Catal.*, 2022, **2**, 892183.

31 C. Trotta, G. Menendez Rodriguez, L. Tensi, A. V. Yakimov, L. Rocchigiani, A. Donnadio, C. Zuccaccia, C. Copéret and A. Macchioni, *Eur. J. Inorg. Chem.*, 2023, **26**, e202300211.

32 L. Fagioli, F. Zaccaria, F. Costantino, R. Vivani, C. K. Mavrokefalos, G. R. Patzke and A. Macchioni, *Dalton Trans.*, 2020, **49**, 2468–2476.

33 B. Yang, D. Luo, S. Wu, N. Zhang and J. Ye, *Sci. Technol. Adv. Mater.*, 2022, **23**, 587–616.

34 S. Nishioka, F. E. Osterloh, X. Wang, T. E. Mallouk and K. Maeda, *Nat. Rev. Methods Primers*, 2023, **3**, 42.

35 H. Zhao, L. Jian, M. Gong, M. Jing, H. Li, Q. Mao, T. Lu, Y. Guo, R. Ji, W. Chi, Y. Dong and Y. Zhu, *Small Struct.*, 2022, **3**, 2100229.

36 N. N. Rosman, R. M. Yunus, N. R. A. M. Shah, R. M. Shah, K. Arifin, L. J. Minggu and N. A. Ludin, *Int. J. Energy Res.*, 2022, **46**, 11596–11619.

37 L. Chen, J. T. Ren and Z. Y. Yuan, *Green Chem.*, 2022, **24**, 713–747.

38 L. C. Goveas, S. Nayak, R. Vinayagam, R. Selvaraj and A. Pugazhendhi, *Fuel*, 2023, **348**, 128460.

39 E. Heo, S. Lee and H. Yoon, *Mater. Chem. Front.*, 2023, **7**, 6154–6187.

40 A. Sherry, M. Tahir and W. Nabgan, *Int. J. Hydrogen Energy*, 2022, **47**, 862–901.

41 T. Simon, N. Bouchonville, M. J. Berr, A. Vaneski, A. Adrovic, D. Volbers, R. Wyrwich, M. Döbling, A. S. Susha, A. L. Rogach, F. Jäckel, J. K. Stolarczyk and J. Feldmann, *Nat. Mater.*, 2014, **13**, 1013–1018.

42 H. Lin, B. Sun, H. Wang, Q. Ruan, Y. Geng, Y. Li, J. Wu, W. Wang, J. Liu and X. Wang, *Small*, 2019, **15**, 1804115.

43 G. Ge, M. Liu, C. Liu, W. Zhou, D. Wang, L. Liu and J. Ye, *J. Mater. Chem. A*, 2019, **7**, 9222–9229.

44 T. Takata, J. Jiang, Y. Sakata, M. Nakabayashi, N. Shibata, V. Nandal, K. Seki, T. Hisatomi and K. Domen, *Nature*, 2020, **581**, 411–414.

45 Z. Sun, H. Zheng, J. Li and P. Du, *Energy Environ. Sci.*, 2015, **8**, 2668–2676.

46 P. Zhou, I. A. Navid, Y. Ma, Y. Xiao, P. Wang, Z. Ye, B. Zhou, K. Sun and Z. Mi, *Nature*, 2023, **613**, 66–70.

47 X. Shi, M. Fujitsuka, S. Kim and T. Majima, *Small*, 2018, **14**, 1703277.

48 L. Zhang, X. Fu, S. Meng, X. Jiang, J. Wang and S. Chen, *J. Mater. Chem. A*, 2015, **3**, 23732–23742.

49 A. Shalav, B. S. Richards and M. A. Green, *Sol. Energy Mater. Sol. Cells*, 2007, **91**, 829–842.

50 Y. Y. Cheng, B. Fückel, T. Khoury, R. G. C. R. Clady, N. J. Ekins-Daukes, M. J. Crossley and T. W. Schmidt, *J. Phys. Chem. A*, 2011, **115**, 1047–1053.

51 M. Penconi, F. Ortica, F. Elisei and P. L. Gentili, *J. Lumin.*, 2013, **135**, 265–270.

52 F. Auzel, *Chem. Rev.*, 2004, **104**, 139–173.

53 T. N. Singh-Rachford and F. N. Castellano, *Coord. Chem. Rev.*, 2010, **254**, 2560–2573.

54 O. S. Kwon, H. S. Song, J. Conde, H. Il Kim, N. Artzi and J. H. Kim, *ACS Nano*, 2016, **10**, 1512–1521.

55 G. Quaglia, B. Bartolomei, P. L. Gentili and L. Latterini, *J. Mater. Chem. C*, 2022, **10**, 9073–9080.

56 H. Islam, O. S. Burheim and B. G. Pollet, *Ultrason. Sonochem.*, 2019, **51**, 533–555.

57 T. Leong and S. Kentish, *The fundamentals of power ultrasound-A review*, 2011, vol. 39.

58 N. Merabet and K. Kerboua, *Int. J. Hydrogen Energy*, 2022, **47**, 17879–17893.

59 J. L. Laborde, A. Hita, J. P. Caltagirone and A. Gerard, *Ultrasonics*, 2000, **38**, 297–300.

60 K. Yasui, *Acoustic Cavitation and Bubble Dynamics*, Springer International Publishing, Cham, 2018.

61 P. Bruno, *Power Ultrasound in Electrochemistry : from Versatile Laboratory Tool to Engineering Solution Sonochemistry*, Wiley, 2012.

62 M. Legay, N. Gondrexon, S. Le Person, P. Boldo and A. Bontemps, *Int. J. Chem. Eng.*, 2011, **2011**, 1–17.

63 K. S. Suslick, *Science (1979)*, 1990, **247**, 1439–1445.

64 M. D. Archer and J. R. Bolton, *Biopolymers*, 1990, **94**, 975.



65 S. S. Rashwan, I. Dincer, A. Mohany and B. G. Pollet, *Int. J. Hydrogen Energy*, 2019, **44**, 14500–14526.

66 S. S. Rashwan, A. Mohany and I. Dincer, *Int. J. Hydrogen Energy*, 2021, **46**, 15219–15240.

67 Y. Son, M. Lim, M. Ashokkumar and J. Khim, *J. Phys. Chem. C*, 2011, **115**, 4096–4103.

68 Z. Zhang, T. Gao, X. Liu, D. Li, J. Zhao, Y. Lei and Y. Wang, *Ultrason. Sonochem.*, 2018, **42**, 787–794.

69 S. Merouani, O. Hamdaoui, Y. Rezgui and M. Guemini, *Int. J. Hydrogen Energy*, 2015, **40**, 4056–4064.

70 S. Merouani, O. Hamdaoui, Y. Rezgui and M. Guemini, *Int. J. Hydrogen Energy*, 2016, **41**, 832–844.

71 K. Kerboua and O. Hamdaoui, *Ultrason. Sonochem.*, 2018, **40**, 194–200.

72 K. Kerboua, S. Merouani, O. Hamdaoui, A. Alghyamah, M. H. Islam, H. E. Hansen and B. G. Pollet, *Ultrason. Sonochem.*, 2021, **72**, 105422.

73 J. Rooze, E. V. Rebrov, J. C. Schouten and J. T. F. Keurentjes, *Ultrason. Sonochem.*, 2011, **18**, 209–215.

74 H. Harada, C. Hosoki and M. Ishikane, *J. Photochem. Photobiol. A*, 2003, **160**, 11–17.

75 T. J. Mason, J. P. Lorimer and D. M. Bates, *Ultrasonics*, 1992, **30**, 40–42.

76 P. R. Gogate, S. Shah and L. Csoka, *Chem. Eng. J.*, 2014, **239**, 364–372.

77 Z. Wei and L. K. Weavers, *Ultrason. Sonochem.*, 2016, **31**, 490–498.

78 Y. Son, M. Lim, J. Khim and M. Ashokkumar, *Ultrason. Sonochem.*, 2012, **19**, 16–21.

79 K. Kerboua and O. Hamdaoui, *Chem. Phys.*, 2019, **519**, 27–37.

80 Y. C. Wang and M. C. Yao, *Ultrason. Sonochem.*, 2013, **20**, 565–570.

81 C. Koch, *Ultrason. Sonochem.*, 2016, **29**, 439–446.

82 P. Domenighini, A. M. Gambelli, F. Rossi and F. Cotana, *J. Phys.: Conf. Ser.*, 2023, **2648**, 012059.

83 Y. Son, M. Lim and J. Khim, *Ultrason. Sonochem.*, 2009, **16**, 552–556.

84 M. Lim, M. Ashokkumar and Y. Son, *Ultrason. Sonochem.*, 2014, **21**, 1988–1993.

85 L. Bai, W. Xu, J. Deng, C. Li, D. Xu and Y. Gao, *Ultrason. Sonochem.*, 2014, **21**, 1696–1706.

86 J. Rooze, E. V. Rebrov, J. C. Schouten and J. T. F. Keurentjes, *Ultrason. Sonochem.*, 2011, **18**, 209–215.

87 F. Cotana and F. Rossi, in *3rd Int. green energy Conf.*, 2007, pp. 3–8.

88 Q. Li, X. Li and J. Yu, *Interface Sci. Technol.*, 2020, **31**, 313–348.

89 M. Penconi, F. Rossi, F. Ortica, F. Elisei and P. L. Gentili, *Sustainability*, 2015, **7**, 9310–9325.

90 A. Dehane, L. Nemdili, S. Merouani and M. Ashokkumar, *Top. Curr. Chem.*, 2023, **381**, 1–25.

91 J. Büttner, M. Gutiérrez and A. Henglein, *J. Phys. Chem.*, 1991, **95**, 1528–1530.

92 H. Rahmani, M. Gholami, A. H. Mahvi, M. Ali-Mohammadi and K. Rahmani, *Journal of Water Chemistry and Technology*, 2014, **36**, 317–324.

93 I. M. Khokhawala and P. R. Gogate, *Ultrason. Sonochem.*, 2010, **17**, 833–838.

94 P. Cui, Y. Chen and G. Chen, *Ind. Eng. Chem. Res.*, 2011, **50**, 3947–3954.

95 S. Chakma and V. S. Moholkar, *J. Ind. Eng. Chem.*, 2016, **33**, 276–287.

96 S. G. Babu, P. Karthik, M. C. John, S. K. Lakhera, M. Ashokkumar, J. Khim and B. Neppolian, *Ultrason. Sonochem.*, 2019, **50**, 218–223.

97 G. Chatel, S. Valange, R. Behling and J. C. Colmenares, *ChemCatChem*, 2017, **9**, 2615–2621.

98 M. Mrowetz, C. Pirola and E. Sellli, *Ultrason. Sonochem.*, 2003, **10**, 247–254.

99 R. A. Torres, J. I. Nieto, E. Combet, C. Pétrier and C. Pulgarin, *Appl. Catal., B*, 2008, **80**, 168–175.

100 P. L. Gentili, M. Penconi, F. Ortica, F. Cotana, F. Rossi and F. Elisei, *Int. J. Hydrogen Energy*, 2009, **34**, 9042–9049.

101 C. Li, X. Wang, J. Wu, J. Gao, R. Zhao, S. Xia, H. Yang, Z. Chen, L. Li and W. Wang, *Ultrason. Sonochem.*, 2023, **99**, 106584.

102 A. Singh and A. S. K. Sinha, *J. Energy Chem.*, 2018, **27**, 1183–1188.

103 S. Tu, Y. Guo, Y. Zhang, C. Hu, T. Zhang, T. Ma, H. Huang, S. C. Tu, Y. H. Zhang, C. Hu, H. W. Huang, Y. X. Guo, T. R. Zhang and T. Y. Ma, *Adv. Funct. Mater.*, 2020, **30**, 2005158.

104 M. B. Starr and X. Wang, *Nano Energy*, 2015, **14**, 296–311.

105 H. Meng, Z. Chen, Z. Lu and X. Wang, *J. Mol. Liq.*, 2023, **369**, 120846.

106 W. Liu, P. Wang, Y. Ao, J. Chen, X. Gao, B. Jia, T. Ma, W. Liu, P. F. Wang, Y. H. Ao, J. Chen, X. Gao, B. Jia and T. Ma, *Adv. Mater.*, 2022, **34**, 2202508.

107 A. J. Zuckerwar and R. L. Ash, *Phys. Fluids*, 2006, **18**, 47101.

108 P. L. Gentili, F. Rossi, M. Penconi, F. Ortica and F. Elisei, in *Hydrogen Production: Prospects and Processes*, ed. D. D. Honnery and P. Moriarty, Nova Science Publishers, New York, 2011, pp. 413–422.

109 M. Penconi, A. Cesaretti, F. Ortica, F. Elisei and P. L. Gentili, *J. Lumin.*, 2016, **177**, 314–324.

110 C. Oses, C. Toher and S. Curtarolo, *Nat. Rev. Mater.*, 2020, **5**, 295–309.

111 Y. Sun and S. Dai, *Sci. Adv.*, 2021, **7**, eabg1600.

112 Y. Ma, Y. Ma, Q. Wang, S. Schweidler, M. Botros, T. Fu, H. Hahn, T. Brezesinski and B. Breitung, *Energy Environ. Sci.*, 2021, **14**, 2883–2905.

113 M. Ashokkumar and F. Grieser, *Chem. Commun.*, 1998, 561–562.

114 O. V. Kharissova, L. M. Torres Martínez, E. Luevano Hipólito, L. F. Garay-Rodríguez, M. R. Alfaro Cruz and B. I. Kharissov, *J. Photochem. Photobiol. A*, 2023, **437**, 114463.

115 R. Singh and S. Dutta, *Sustainable Fuel Technologies Handbook*, 2021, pp. 541–549.

