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1. Introduction

Due to the gradual, but constant and inescapable depletion of fossil fuels and the increasing energy demand to support the current model of economic growth, mankind is facing a global energy problem.^{1,2} Amongst the number of alternative resources, renewable energies are rapidly becoming the leading solution to fulfil the growing needs of power sources.^{3,4} In this respect, wind power, solar energy, hydropower, geothermal energy, biomass and biofuel are extensively investigated for a few decades both from the scientific/academic and industrial viewpoints.⁵

At present, solar energy is considered the most promising renewable resource: every day, the Sun shines on the Earth, thus providing around 3×10^{24} J of green energy per year, which exceeds by a factor of 10^4 the present global population consumption.^{6–8} A simple calculation leads self-evidently to the conclusion that covering only around 0.1% of the Earth's surface

Aqueous dye-sensitized solar cells

Federico Bella,*^a Claudio Gerbaldi,^a Claudia Barolo^b and Michael Grätzel*^c

Nowadays, dye-sensitized solar cells (DSSCs) are the most extensively investigated systems for the conversion of solar energy into electricity, particularly for implementation in devices where low cost and good performance are required. Nevertheless, a key aspect is still to be addressed, being considered strongly harmful for a long time, which is the presence of water in the cell, either in the electrolyte or at the electrode/electrolyte interface. Here comes the present review, in the course of which we try our best to address the highly topical role of water in DSSCs, trying to figure out if it is a poisoner or the keyword to success, by means of a thoroughly detailed analysis of all the established phenomena in an aqueous environment. Actually, in the last few years the scientific community has suddenly turned its efforts in the direction of using water as a solvent, as demonstrated by the amount of research articles being published in the literature. Indeed, by means of DSSCs fabricated with water-based electrolytes, reduced costs, non-flammability, reduced volatility and improved environmental compatibility could be easily achieved. As a result, an increasing number of novel electrodes, dyes and electrolyte components are continuously proposed, being highly challenging from the materials science viewpoint and with the golden thread of producing truly water-based DSSCs. If the initial purpose of DSSCs was the construction of an artificial photosynthetic system able to convert solar light into electricity, the use of water as the key component may represent a great step forward towards their widespread diffusion in the market.

> by means of energy conversion devices having an efficiency of about 10% would satisfy the present global energy needs. These encouraging numbers are inducing the scientific community to make even greater efforts towards the direction of improving solar energy conversion technologies as well as proposing new intriguing solutions.^{9,10} To have an idea of the magnitude of this effort, a rapid search on a scientific database (*e.g.*, Scopus), using the simple keyword "solar energy", provides more than 10 000 publications for each of the recent years.

> The creation of a potential difference, or electric current, in a material upon exposure to light is possible because of the photovoltaic (PV) effect, and - depending on the materials used three generations of solar energy devices have been developed to date.¹¹⁻¹³ Among all, the mostly studied device of the last generation is the dye-sensitized solar cell (DSSC), being a lowcost and high efficiency solar energy-to-electricity converter. Firstly assembled and demonstrated by O'Regan and Grätzel in 1991,¹⁴ they are composed of only five components (Fig. 1): a transparent conductive oxide (TCO) substrate,15 a nanostructured n-type semiconductor,¹⁶ a visible-light absorber dye,¹⁷ an electrolyte¹⁸ and a counter electrode.¹⁹ With the key idea of mimicking the natural photosynthesis, DSSCs are being pursued as eco-friendly devices, which could be easily fabricated.^{20,21} In a dye-sensitized solar cell, upon absorption of light, dye molecules reach an excited state and, with an appropriate energy level alignment of all the

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pioneered the use of mesoscopic

materials in energy conversion

and storage systems, in particular

photovoltaic cells, lithium ion

batteries and photoelectrochemical

devices for water splitting by

sunlight. He has received the

Albert Einstein World Award of

Science, the Gutenberg Research

Award, the Galvani Medal, the

de Lausanne.

as

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nanostructured electrodes,

components, charge separation occurs at the interface between the dye-sensitized semiconductor and the electrolyte. While electrons are injected into the conduction band (CB) of the semiconductor (usually titanium dioxide, TiO₂) and transported to the conductive electrode, the regeneration of the oxidized dye takes place at the counter electrode by means of an electrondonor species (typically a liquid electrolyte based on the redox couple iodide/triiodide, I^-/I_3^-).²² Multidisciplinarity, academic and industrial interests have boosted the number of publications per year (Fig. 2), resulting in about 4000 international peer-reviewed0papers in the 2013–2014 period of time. Some of these articles, which led to the identification of new materials with concrete applications also in other research fields, 23,24 are highly cited: each of them counts even more than 1000 citations. $^{25-28}$

The most performing DSSCs (efficiencies up to 13% have been recently demonstrated²⁹) use organic solvent-based liquid electrolytes. These have the relevant drawback of high vapour pressure and a severe environmental impact. Moreover, several organic solvents are toxic and/or explosive, thus seriously limiting their practical applications in DSSCs because of safety issues. Although several alternatives to organic solvents have been proposed (*i.e.*, plastic crystals and solid-state conductors^{30,31}), the biggest – sometimes ignored – issue still remaining unsolved



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field of sensitizers for solar cells, she is the recipient of several

collaborations, and, since 2011 she has been a member of the

Technical and Scientific Committee of Dyepower consortium. She

has published more than 55 ISI articles with \sim 200 annual citation

research grants,



Claudio Gerbaldi

and innovative polymer electrolytes for the development of ecofriendly energy storage and conversion devices. Among others, he has received the "UniCredit Award" for the Young Innovation in Research (2007) and has published \sim 75 ISI articles with \sim 250 annual citation counts in recent years.

Fédérale



Claudia Barolo

national and international

counts in recent years.

Claudia Barolo received her PhD in Chemistry from Università di Torino in 2001. In 2006, she became an assistant professor and, in 2014, an associate professor in Industrial Chemistry at the Department of Chemistry of the same university. Her research activity is mainly focused on the synthesis and characterization of functional molecules and hybrid materials for technological applications (photonics, nanotechnology, biotechnology). As an expert in the

industrial



Michael Grätzel

Faraday Medal and the Millenium Technology Grand Prize. He has published ~ 1100 papers, 50 reviews/book chapters and is the inventor/ co-inventor of >50 patents. His studies have been cited over 140000 times (h-index 170), making him one of the 10 most highly cited chemists in the world.

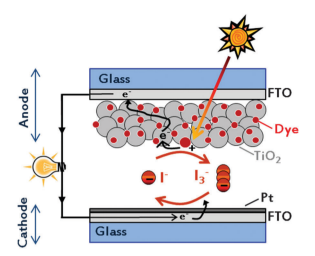


Fig. 1 Schematic representation of the components and of the basic operating principle of a DSSC.

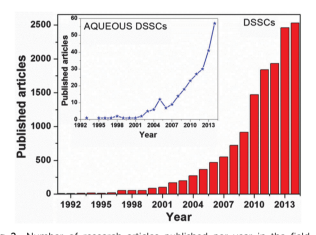


Fig. 2 Number of research articles published per year in the field of DSSCs. Inset: data for aqueous DSSCs. Source: SCOPUS Database.

in standard aprotic DSSC systems is the contamination of the cell by means of moisture/water. This often affects both the cell performance and the long-term stability, even if robust encapsulation systems and specifically conceived barrier materials are utilised. In fact, traces of water are always present in the electrolytic solution and in the pores/voids of the semiconductor electrode film, being eventually introduced during cell assembly or upon operation under ambient conditions (tiny defects in the encapsulation are often present). This phenomenon is even more accentuated when the production of flexible DSSCs is envisaged,^{32,33} quite rapidly permeated by water molecules (*i.e.*, $0.01 \text{ g m}^{-2} \text{ day}^{-1}$) even in the presence of rather expensive barrier layers; in fact, it has been calculated that the water content in the electrolyte may be >10% after one year of real outdoor use.³⁴ These results have led to the general opinion that water is detrimental, in fact a poisoner, for DSSCs.

The aim of this review article is to thoroughly unravel the specific role of water as an electrolyte component in DSSCs. First of all, it is fundamental to understand how the presence of water can influence the device performance; many studies have

been proposed over the years in this respect. Crossing experimental data obtained by using very different techniques as well as theoretical simulations would help in identifying the interactions between water molecules and DSSC components. Then, we consider very much interesting and surprising to show how, in the last few years, the scientific community has turned its efforts in the direction of DSSCs fabricated with water-based electrolytes. Indeed, being less expensive, non-flammable and more environmentally friendly, these novel devices would also not suffer from water contamination issues, with the added value of easily solvating many potential redox mediators. Photoanode modifications, introduction of novel additives and surfactants, selection of specifically conceived redox couples, preparation of suitable cathodes and stabilization of electrolytes have progressively led to the fabrication of 100% aqueous DSSCs. Achieving high performance in a fully aqueous cell represents the Holy Grail for DSSCs, and brings back the attention to their initial purpose: the realisation of an artificial photosynthetic system able to convert solar visible light energy into electricity, by using a unique solvent, water, the solvent of life.

2. Electrolytes for DSSCs: state of the art

The electrolyte has been identified as one of the component which most significantly influences DSSC efficiency and stability, since it has a key role in the regulation of the electron transfer kinetics. According to their physical consistency, electrolytes may be classified into three main classes: liquid, quasi-solid and solid.

Traditional liquid electrolytes consist of a redox couple and a few additives dissolved into an organic solvent. As previously stated, the standard redox mediator is the I⁻/I₃⁻ couple, but several valid alternatives have been proposed. In particular, cobalt complexes³⁵ have exceeded the performance of the standard redox pair, achieving efficiencies up to 13%.29 Other redox mediators, such as SCN⁻/(SCN)₃⁻,³⁶ SeCN⁻/(SeCN)₃⁻,³⁷ Br⁻/Br₃⁻, ³⁸ sulphur-based systems, ³⁹ copper complexes, ⁴⁰ ferrocene derivatives⁴¹ and stable nitroxide radicals,⁴² have also been proposed. Each of these systems is characterized by peculiar thermodynamic and kinetic properties, which must be well matched with the other cell components. Moreover, also counterions of redox mediators must be properly selected, since it has been reported that the photocurrent decreases and the photovoltage increases while increasing the cation radius, due to the variation of the TiO_2 CB energy (E_c) and the associated influence on the electron injection efficiency.⁴³

Additives are often introduced in appropriate amounts to increase the PV parameters; among them, nitrogen-containing heterocyclic compounds and guanidinium thiocyanate (GuSCN) are the most frequently used. While the former improve cell photovoltage by shifting the band edge of TiO_2 and increasing the electron lifetime,⁴⁴ the latter enhances the photocurrent by positively moving the flatband potential of the photoelectrode, thus increasing the electron injection yield.⁴⁵

As regards organic solvents, acetonitrile (ACN), valeronitrile (VAN), 3-methoxypropionitrile (MPN), methoxyacetonitrile (MAN), ethylene carbonate (EC), propylene carbonate (PC), γ-butyrolactone (GBL) and N-methylpyrrolidone (NMP) are the most commonly used, due to their high dielectric constant and low viscosity. Since all these solvents can be considered as electron acceptors, a classification of their donor number (i.e., the enthalpy of reaction of the solvent with a strong electron acceptor) was proposed,46 and it was proved that an increase in the electrondonor (or basic) character led to higher cell potentials and lower photocurrents.⁴⁷ The worst drawback of organic solvents is their high volatility, thus room temperature ionic liquids (RTILs), being non-flammable and non-volatile organic compounds, are widely used as additives or solvents in order to guarantee good chemical and thermal stabilities.⁴⁸ However, RTILs often show high viscosity values, and specific improvements of their rheological characteristics are necessary to ensure an efficient ion transport. The enormous research efforts carried out in the field of liquid electrolytes have been already thoroughly discussed in several review articles.49-53

Limited long-term stability, difficulty in hermetic sealing and leakage of the liquid electrolyte have been identified as relevant drawbacks in view of real indoor and outdoor applications of DSSCs.^{54–56} A possible solution is represented by the entrapment of the liquid electrolyte by means of a polymeric or inorganic network. Polymeric quasi-solid electrolytes can be prepared in the form of gels⁵⁷ or membranes,^{58,59} where a huge amount of crosslinked matrices has been investigated.^{60–62} As an alternative, inorganic nanoparticles (NPs) have been adopted to jell liquid electrolytes, creating a quasi-solid (paste-like) network containing self-assembled channels able to ensure sufficient ionic transport.^{63–65} To date, quasi-solid electrolytes are considered the optimum compromise between efficiency and durability,⁶⁶ and the recent advances in this subject have been collected in some review articles.^{67–69}

Solid electrolytes are based on a completely different working principle: holes transferred from the dye are transported via hopping between the electronic states of the solid electrolyte to the metallic or polymeric counter electrode. Crystalline p-type semiconductors,⁷⁰ organic hole-conducting molecules,⁷¹ as well as conductive polymers⁷² have been widely studied.⁷³⁻⁷⁵ However, the poor intrinsic conductivity, the low electron transfer from the dye molecules, the poor penetration into the mesoporous TiO₂ and the faster recombination phenomena do not allow solid cells to be more efficient than their liquid and quasisolid counterparts. Studies on solid DSSCs have recently led to the development of perovskite solar cells (PSCs), in which compounds like CH₃NH₃PbI_xCl_{3-x} act as the light harvester as well as the conductor of both electrons and holes.^{76–78} PSCs have achieved efficiencies above 19%79 (certified value: 17.9%,80) and, although very young, will surely be able to establish themselves as a leading device for solar energy conversion.

The introduction of water in liquid, quasi-solid and solid electrolytes causes a very complex set of effects on the PV performance of the resulting DSSCs. The initial section of this review will focus on their detailed investigation, while the rest of the manuscript will deal with cells in which water is purposely added to the electrolyte with gradually increasing percentages, giving rise to so-called "aqueous DSSCs".

3. Harmful effects due to water contamination

As anticipated in the Introduction, the presence of water was initially viewed as a strongly negative aspect for DSSCs. Where does water in cells come from? Unless DSSCs are assembled in a N2- or Ar-filled glove-box (procedure almost never encountered in the literature), water is present during all the manufacturing steps. Indeed, it can be adsorbed on the anodic semiconductor, be present in the solvents and in the solutions used to make the dve-uptake or the liquid electrolyte filling, permeate the polymer films used for device sealing. The effects of water on the PV performance, and the stability and the visual aspects of DSSCs have been extensively studied by adopting several experimental techniques and/or through computational studies, which are reported in the following Sections (3.1-3.5). For ease of understanding, we will report the water content as percentage by weight (wt%) or volume (vol%); however, in the case of very low quantities we will adopt the molar concentration (for H₂O, 1.0 M corresponds to 1.8 wt%).

3.1 Effects on photovoltaic parameters

A couple of years after inventing the DSSCs, Grätzel and coworkers proposed the first study on the stability of *cis*-X₂bis(2,2'-bipyridyl-4,4'-dicarboxylate)-ruthenium(II) (X = Cl⁻, Br⁻, I⁻, CN⁻ and SCN⁻) sensitizers.⁸¹ All of the cells showed an initial 20–30% decay in short-circuit current density (J_{sc}), which was attributed to the absorption of residual UV light as well as to the contamination by water molecules. Such detrimental effects were attributed to the electronic and chemical equilibration of the dye-loaded TiO₂ with the liquid electrolyte, the water uptake from ambient air being envisaged as their main cause. Indeed, when a water-tight cell sealant was used, the initial decay in the performance of the cell was suppressed.

The first analytical investigation of water contamination dates back to 1998, when Lindquist and coworkers showed that $J_{\rm sc}$ decreased (Fig. 3A) and open-circuit voltage ($V_{\rm oc}$) increased (Fig. 3B) when the water content in the electrolytic solution (LiI 0.10 M and I_2 10 mM in MAN) increased from 0 to 2.2 M.⁴³ It was proposed that water molecules, being strongly adsorbed onto the TiO₂ surface, coordinated with the surface Ti atoms⁸² and blocked the reaction of I_3^- with the electrons in the TiO_2 CB. As a result, V_{oc} increased proportionally to the amount of water introduced in the cell. Such a phenomenon was so strong as to balance and quantitatively overcome the negative shift of the CB which was caused by the addition of water to the aprotic electrolytic solution.⁸³ Anyway, despite the improvement in $V_{\rm oc}$, cell performance decreased due to the neat reduction of J_{sc} values. In their preliminary studies, Lindquist and coworkers motivated this effect by arguing the following hypotheses: desorption of the N3 dye from the TiO₂ surface, weakening of

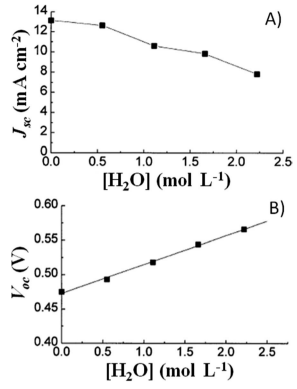


Fig. 3 Variation of J_{sc} (A) and V_{oc} (B) values, measured a 0.82 sun, due to water addition in the liquid electrolyte. Electrolytes: Lil 0.10 M and I_2 10 mM in MAN/H₂O. Dye: N3. Adapted and reprinted from ref. 43.

the TiO₂-dye interaction, photoinduced substitution of the -NCS ligand of the dye and change in the absorption properties of the sensitizer molecules due to the variation in the solvent polarity. For sure, the most intriguing phenomenon was observed when the contaminated electrolyte was replaced by a freshly prepared water-free one: $J_{\rm sc}$ was fully restored (from 1.93 to 2.62 mA cm⁻²), while $V_{\rm oc}$ retained a higher value (0.535 V vs. the initial 0.453 V). As a result, the overall cell efficiency (η) increased (from 5.2% to 6.4% at an irradiation intensity $P_{\rm in}$ of 0.15 sun), and Lindquist's group compared the effect observed in the water-contaminated systems to that provided by the traditionally used pyridine derivative additives.⁸⁴

3.2 Spectroscopic investigation of water-promoted effects

Fourier transform infrared (FTIR) studies reported in Fig. 4A–C showed that when the working electrodes were immersed in an electrolyte (LiI 0.50 M and I₂ 50 mM in MPN) containing 5 vol% H₂O, the broadening as well as decreased intensity of the –NCS ligand absorption band (2100 cm⁻¹) was observed. In the same study, Hagfeldt and coworkers found a small absorption peak at 1575 cm⁻¹, which was assigned to the symmetric bending of the water molecules, either bonded to the N719 dye or adsorbed onto the nanostructured TiO₂ film.⁸⁵ UV-Vis spectra (Fig. 4D–F) showed blue-shifted absorption maxima with respect to the reference ($\lambda_{max} = 551$ nm), both under dark ($\lambda_{max} = 529$ nm) and under normal illumination conditions ($\lambda_{max} = 500$ nm); these shifts were probably caused by the exchange of the –NCS

ligands of the dye with water molecules and/or OH⁻ anions in the contaminated electrolyte.

To ascertain the eventual influence of hydrogen bonds and hydrophobic interactions on the water-contaminated cell components, the use of water isotopes (*i.e.*, H₂O, D₂O and H₂¹⁸O) has been proposed by Yang and coworkers,⁸⁶ who exploited the potential prospects of this procedure usually applied in geology, biology and medicine.^{87,88} By monitoring the time-dependent surface reaction occurring on the photoelectrodes in the presence of water isotopes, the diffuse-reflectance infrared Fourier transform (DRIFT, Fig. 5) technique showed that the intensity of the SCN⁻ ligand remarkably decreased when the water content in the electrolyte (LiI 0.50 M, I2 50 mM and TBP 0.50 M in ACN; TBP = 4-tert-butylpyridine) increased (from 0 to 5 wt%). Additionally, the peak intensity of SCN⁻ decreased with prolonged soaking time, due to the progressive replacement of the -SCN ligand with water molecules or OH⁻ ions.⁸⁹ This was confirmed by the appearance of a strong band at $3000-3600 \text{ cm}^{-1}$, due to water adsorption on the nanostructured TiO₂ lattice.⁹⁰ Moreover, the interaction of Li⁺ cations (from LiI) with the free SCN⁻ anions led to the formation of LiNCS, as indicated by the peak at 2079 cm⁻¹.⁹¹ Considering the sensitizer structure as a whole, it resulted that only the bipyridine ligands of the sensitizer did not take part in the degradation process. When water isotopes were used, Yang and coworkers measured that the degradation rate increased in the order of $H_2O > D_2O > H_2^{18}O$, fully consistent with the diffusion rate of these isotopes in the electrolyte, which is inversely proportional to the square root of their molecular masses.⁹² As regards the PV performance, the bonding between Li⁺ and SCN⁻ ions gave rise to negative shifts in the TiO₂ Fermi level, which resulted in lower $V_{\rm oc}$ values for aqueous DSSCs, in contrast to the previously discussed results by Lindquist and coworkers.43

3.3 Imaging techniques and visual aspects: how cells change over time

Photocurrent imaging techniques represent a valuable investigation method to provide information on problems related to DSSC efficiency as well as long-term stability.^{93,94} Cell mapping is usually carried out by means of a He-Ne laser (632 nm), inducing a photocurrent point by point, which is registered through a lock-in amplifier. Using such a technique, Tributsch and coworkers observed the bleaching of the electrolyte, as a result of the disappearance of I₃⁻ ions.⁹⁵ This resulted in the increase of the internal resistance, because the I_3^- reduction reaction at the counter electrode stopped. Interestingly, this was the first work to go in contrast to the others: the effect of water was not geared solely to the dye stability, but also the electrolyte underwent major changes. Even if the formation of iodate (IO₃⁻) from I₃⁻ was expected by Tributsch and coworkers, FTIR analysis revealed the absence of iodate in the electrolyte. Nevertheless, the FTIR spectrum of the electrolyte in the region of the OH-stretching mode showed increased water concentration values in the solvent (ACN). Moreover, the water content increased with time, and aging studies showed that the corresponding OH-stretching mode became the main signal in the spectrum after 84 days. Thus, it was proposed that water could

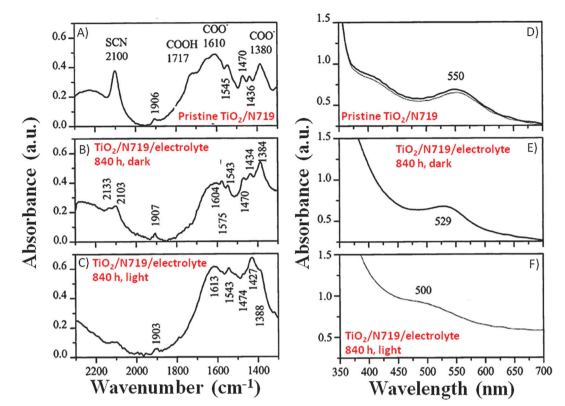


Fig. 4 FTIR (left side) and UV-Vis (right side) spectra of: (A, D) non-treated working electrodes in an Ar atmosphere; (B, E) working electrodes being soaked in a liquid electrolyte (Lil 0.50 M and I_2 50 mM in MPN) with 5 vol% H_2O under dark for 840 h; (C, F) working electrodes being soaked in the same liquid electrolyte under visible light illumination for 840 h. FTIR spectra of the treated samples were normalized based on the TiO₂ absorption maximum of the reference sample at 832 cm⁻¹. Adapted and reprinted from ref. 85.

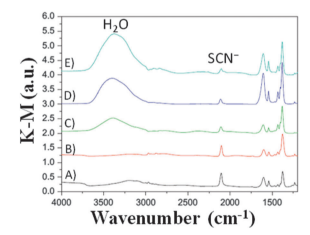


Fig. 5 DRIFT spectra of N719/TiO₂ films immersed in liquid electrolytes (Lil 0.50 M, I₂ 50 mM and TBP 0.50 M in ACN) having different water contents for 192 h: (B) 0 vol% H₂O; (C) 1 vol% H₂O; (D) 2 vol% H₂O; (E) 5 vol% H₂O. The response of the photoelectrode before immersion is also shown (A) for comparison purposes. Adapted and reprinted from ref. 86.

activate as a weak base TBP, commonly used as a $V_{\rm oc}$ improver.⁴⁴ This would have caused the instability of the I₃⁻ anions: as wellknown from basic ion chemistry, I₃⁻ ions become unstable and IO₃⁻ is formed when the pH increases up to 8–10.⁹⁶ However, the expected absorption of IO₃⁻ at 800 cm⁻¹ was not detected,⁹⁷ thus its adsorption on the TiO₂ electrode was proposed.

Spatially resolved photocurrent mapping⁹⁸ has also been proposed by Cameron and coworkers, who coupled Z907 dyesensitized photoelectrodes with ACN- as well as water-based electrolytes.⁹⁹ When vacuum-filled, the ACN-containing cells showed a homogeneous performance all over their volume (Fig. 6A), with the exception of an area of high photocurrent at the top of the device, where dye coverage was higher. As regards aqueous cells, the map showed an area with high photocurrent spreading out from the filling hole (Fig. 6B), where maybe electrolyte filling was relatively poor. To overcome such a defect, aqueous cells were vacuum treated at 80 °C: homogeneous current densities were obtained across the whole area of the cell (Fig. 6C), due to an improved electrolyte penetration. Anyway, all of the aqueous cells underwent a loss of photocurrent (and dye as well) nearby the filling holes; Cameron and coworkers hypothesized that TBP-based alkali solutions could desorb dye from TiO₂, especially around the filling holes. After the previous study by Tributsch and coworkers,95 this was the second research article where a synergistic effect between water and TBP was observed to have detrimental effects on the device stability.

3.4 Modelling interactions between water and cell components

After being experimentally consolidated that water interacts with the molecules of dye adsorbed onto the semiconductor

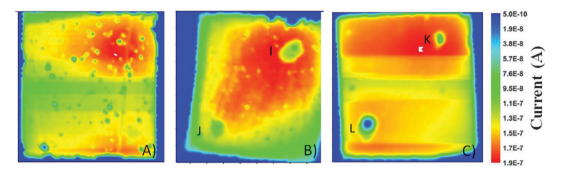


Fig. 6 Photocurrent maps for: (A) vacuum-filled ACN-based cell; (B) vacuum-filled aqueous cell; (C) high temperature vacuum-filled aqueous cell. The positions of the filling holes and the corresponding cold spots are indicated by the letters I–L. The side of each image is 1 cm. Adapted and reprinted from ref. 99.

active material particles, several mechanistic studies have been proposed to investigate this phenomenon. De Angelis and coworkers performed an ab initio molecular dynamics simulation and an extensive time-dependent density functional theory (TDDFT) excited state calculation of a squaraine dye^{100,101} anchored to an anatase slab exposing the (101) surface,¹⁰² surrounded by 90 water molecules.¹⁰³ Water was observed to adsorb, also in overlayer configuration,¹⁰⁴ at the TiO₂ surface. As a result, the blocking function toward recombination of injected electrons with oxidized species in the electrolyte proposed by Lindquist and coworkers⁴³ was confirmed. As regards the squaraine dye, the TDDFT study demonstrated that an intensive solvent reorganization around the TiO₂-bonded carboxylic group led to a transition from the starting bridged-bidentate to the monodentate adsorption configuration. The latter was stable for ~ 6 ps; then, the proton anchored at the surface was transferred to the anionic carboxylic group, resulting in the desorption of the dye, which is illustrated in Fig. 7.

Apart from the dye to water interaction, the concurrent presence of TiO₂ to water as well as organic solvent to water interactions should also be considered in DSSC modelling. It is important to recall that aprotic solvents (e.g., ACN and MPN) are responsible for the high DSSC efficiency,¹⁰⁵ due to their high dielectric constant and solubilisation ability versus many inorganic and organic salts and additives.^{106,107} Furthermore, transient adsorption spectroscopy (TAS) demonstrated that, when photoanodes were immersed in ACN, a 50% increase in the signal intensity occurred, indicating that aprotic solvents increased the electron injection efficiency. In this respect, Tateyama and coworkers modelled the effect of ubiquitous water contamination in an ACN-based electrolyte solution¹⁰⁸ by means of DFT methods implemented in Car-Parrinello molecular dynamics (CPMD).^{109,110} Surprisingly, considering the solvation shell $(ACN)_{46}(H_2O)_1$ ([H₂O] = ~0.50 M), it was illustrated that ACN molecules prevented water from coming in contact with the (101) TiO₂ surface by means of a network of hydrogen bonds. However, CPMD also indicated that water molecules that were already adsorbed onto the photoanode before cell assembly were hardly removed because of their strong interaction with the anatase (101) surface. This strong water to TiO₂ interaction (Fig. 8) was observed to be stable up to

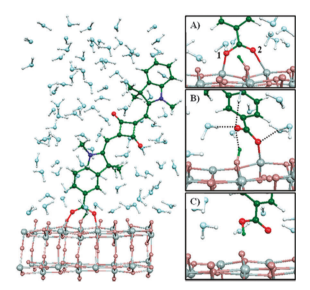


Fig. 7 Squaraine dye adsorbed onto an anatase slab, surrounded by 90 water molecules. In the right hand-sided images, relevant configurations sampled during the TDDFT study are shown: (A) initial bridged-bidentate configuration with labeled oxygen atoms; (B) monodentate configuration; (C) dissociated dye. Note the different proton positions (green). Dashed lines in (B) represent hydrogen bonds to the carboxylic group. Reprinted from ref. 103.

400 K, and could be detrimental for DSSC performance. In fact, photogenerated holes could migrate to water molecules (due to the oxygen atoms charges), thus producing water cation radicals able to interact with the dye molecules onto the TiO_2 surface. Therefore, the common procedure to operate in an anhydrous environment during DSSC fabrication was strongly recommended by the authors.

3.5 Effects of sealing conditions and device area

The experimental conditions of DSSC manufacture have been studied by Dittrich and coworkers,¹¹¹ who focused their attention on oxygen- and water-related surface defects on the photoanode, such as oxygen vacancies (or Ti^{3+} donor states) and hydroxy surface complexes. Cells were assembled either with or without pre-treatment under vacuum. It was observed that J_{sc} values decreased when the TiO_2 surface was pre-treated under vacuum, due to the

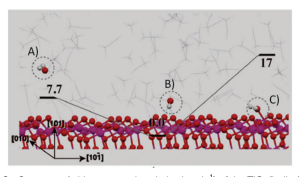


Fig. 8 Structures (with energy values in kcal mol⁻¹) of the TiO₂/bulk-ACN interfaces contaminated with water. (A) Water molecules exist in bulk ACN; (B) the most stable structure of water molecule in bulk ACN and water adsorbed on the TiO₂ surface *via* a hydrogen bond through the H atom in water; (C) water adsorbed onto the TiO₂ surface *via* a hydrogen bond through the O atom in water. Adapted and reprinted from ref. 108.

formation of recombination paths for injected electrons.^{112,113} The decrease of J_{sc} was observed to be more pronounced when the cells were pre-treated under a mixed vacuum/water environment, the interaction of –OH groups with TiO₂ being very intense on defective sites.¹¹⁴ In contrast, chemisorption of water improved V_{oc} values in both vacuum and oxygen pre-treated cells, as a result of the increase of the barrier height between the TiO₂ CB and the redox mediator potential (*E*).

All of the studies so far reported focusing on water contamination highlighted several detrimental phenomena occurring in different DSSC components: negative shift of the TiO₂ CB, weakening of the TiO₂-dye interaction up to dye desorption, photoinduced substitution of the -NCS ligand and the related change in the absorption properties of the sensitizer, disappearance of I₃⁻ ions, unwelcome interaction of water with both additives (TBP) and salts (LiI). Moreover, thinking about DSSCs large-scale applications it is important to consider that all of these drawbacks are exponentially emphasized with the increase in the overall volumetric dimension of the device. Considering 5 cm \times 5 cm cells assembled under an ambient atmosphere (50-60% RH), Kitamura and coworkers observed the progressive decolouration of the photoelectrodes, accompanied by a 40% decrease in $J_{\rm sc}$ after 800 h at ambient temperature and under dark conditions.¹¹⁵ This effect was prevented by assembling the cells in the inert atmosphere of an Ar-filled dry glove box (0.1% RH).

At present, two options are available. Specific technologies that may 100% avoid water contamination during the whole manufacturing process of DSSCs as well as during their entire operational life have to be adopted. Alternatively, scientists have to propose dyes, electrodes and electrolytes which can properly operate under a partial or fully aqueous environment, taking advantage of the amount of positive aspects that water would guarantee to third generation PV devices, as stated in the introductory section of this review. We believe that the first option is hardly feasible in terms of cost as well as manufacturing issues (especially because DSSCs are considered promising also for their ease of manufacture), while the development of waterbased or fully aqueous DSSCs is certainly the most exciting and promising prospect as will be detailed and analysed in the following sections.

4. Back to 1988: aqueous photoelectrochemical cells

Before discussing the recent findings concerning aqueous DSSCs, it is important to move a step back to a few years before their invention (1991). Indeed, it must be enlightened that the photoelectrochemical cells developed in the 1980–1990 decade were intended to operate with fully aqueous electrolytes.^{116,117}

In 1988, Grätzel and coworkers reported on the surprisingly broad band sensitization of TiO₂ to visible light by means of *cis*-Ru(n)L₂(H₂O)₂²⁻ (L = 2,2'-bipyridyl-4,4'-dicarboxylate).¹¹⁸ A high surface area fractal anatase film deposited onto a Ti sheet acquired an intense violet coloration due to adsorption of the dye from an acidic aqueous solution. The regenerative photoelectrochemical cell containing a KI 0.10 M and I₃⁻ 1.0 mM aqueous electrolyte provided an efficiency of 2% ($J_{sc} = 0.38$ mA cm⁻², $V_{oc} = 0.520$ V, fill factor FF = 0.70; $P_{in} =$ 0.7 sun). Interestingly, the photocurrent remained stable after 4 days of illumination, thus confirming the remarkable stability of the sensitizer; moreover, a maximum incident photon to current efficiency (IPCE) value of 62% was observed. In this study, water was present in the sensitizer molecule, in the dyeuptake solution and in the electrolyte as well.

In 1988, RuL_3^{4-} (L = 2,2'-bipyridyl-4,4'-dicarboxylate) was proposed by Grätzel and coworkers as another promising sensitizer for aqueous photoelectrochemical cells;¹¹⁹ it was also coupled with a new Br-based redox couple (LiBr 1.0 M, Br₂ 1.0 mM and HClO₄ 1.0 mM). Upon irradiation with a 1.58 W m⁻² monochromatic light (470 nm), the cell demonstrated a remarkable efficiency of 12% (short-circuit current I_{sc} = 135 µA, V_{oc} = 0.720 V, FF = 0.74; IPCE = 56%); at that time, this was the highest monochromatic conversion yield achieved using a dye-sensitized regenerative photoelectrochemical cell.

With the advent of DSSCs in 1991,¹⁴ aprotic organic solvents replaced water, and cells operating in an aqueous environment were no longer investigated for the following 10–15 years.

5. The trend is being reversed: aqueous electrolytes

In the twenty-first century, scientists started again to focus their research efforts on more deeply investigating the effects of water in DSSCs, the final challenging goal being the full replacement of the traditionally used organic solvents. The seminal paper published in 2010 by O'Regan and coworkers may be undoubtedly considered the fundamental contribution for the scientific community in this context.¹²⁰ The authors varied the relative fractions of MPN and water to prepare electrolytes (PMII 2.0 M, I₂ 50 mM, GuSCN 0.10 M and TBP 0.5 M; PMII = 1-propyl-3-methylimidazolium iodide) having 0, 20, 40, 60, 80 and 100 vol% H₂O with respect to MPN. Three noticeable insights were proposed in this work: the use of a

hydrophobic dye (TG6), the introduction of 1% Triton X-100 (see Section 7) to avoid phase separation in the aqueous electrolyte and the selection of a high PMII concentration, which concurrently acted as an iodide source and a surfactant. As shown in Table 1, being iodide more soluble in water than tri-iodide, the addition of water shifted *E* towards positive values. Moreover, no detrimental effect on cell performance was observed up to 40 vol% H₂O. The basic functions of DSSCs (injection, regeneration and transport) worked properly even when using electrolytes having high water contents. Furthermore, the 80 vol%-based cell showed only 7% and 8% losses in $J_{\rm sc}$ and $V_{\rm oc}$, respectively, after 1000 h under 1 sun (35 °C with UV-filter). The desorption of the hydrophobic dye TG6^{121,122} was not detected.

Interestingly, it was observed that, by reducing the irradiation intensity, the performance mismatch between aqueous and non-aqueous electrolytes decreased. As shown in Fig. 9, O'Regan and coworkers observed that the fraction of photocurrent lost at low $P_{\rm in}$ and the saturation photocurrent at higher $P_{\rm in}$ values were correlated, and an increase in performance variation between identical cells at high water content was present. This represented a key issue: if in Section 3 we reported that the presence of water affected the PV parameters, the stability over time and the visual appearance of DSSCs, now also the worsened reproducibility emerges. Such a scenario opened the debate in the scientific community, and the following arguments were proposed:

– A reduction in the dye excited-state lifetime occurred due to the increase in the dielectric constant when moving from organic solvents towards water.¹²³ However, even if losses at the injection step could be independent of $P_{\rm in}$, they were not expected to strongly vary when identical cells were considered.

– Electrons could be lost by increased recombination with the oxidized dye molecules, due to the increase of *E* (see Table 1), which reduced both the driving force and the rate constant for dye regeneration. However, even if losses from this route are usually strongly dependent upon $P_{\rm in}$, O'Regan and coworkers performed photovoltage transient experiments which demonstrated that the variation in the electron recombination lifetime was reduced by a factor of two over all the different water contents.

– The real explanation of the behaviour showed in Fig. 9 involved phase segregation inside the pores of TiO_2 , due to non-homogeneity in the pore sizes as well as in the dye coverage. In fact, the concurrent presence of (i) pores having high I_2/TBP

Table 1 *E* values for aqueous electrolytes and PV parameters ($P_{in} = 1 \text{ sun}$) of the corresponding DSSCs. Electrolytes: PMII 2.0 M, I₂ 50 mM, GuSCN 0.1 M and TBP 0.5 M in H₂O/MPN. Dye: TG6. Adapted and reprinted from ref. 120

H ₂ O (vol%)	$E(\mathbf{V})$	$J_{\rm sc}~({ m mA~cm^{-2}})$	$V_{\rm oc}$ (V)	FF	η (%)
0	0.058	11.3	0.73	0.67	5.5
20	0.077	11.8	0.73	0.67	5.7
40	0.103	11.1	0.73	0.68	5.5
60	0.117	8.9	0.75	0.67	4.5
80	0.129	6.5	0.75	0.68	3.3
100	0.136	4.7	0.74	0.69	2.4

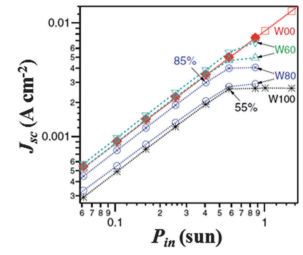


Fig. 9 J_{sc} vs. P_{in} for selected water-free and water-based DSSCs (see Table 1). Percentages are J_{sc} relative to the water-free cell at the same P_{in} value. Adapted and reprinted from ref. 120.

phases, (ii) high H_2O/I^- phases and (iii) empty pores resulted in a reduced redox couple diffusion throughout the photoanode. Since phase segregation was sensitive to the exact pore structure, the variation in performance within identical cells at high water content was explained.

Following the insights proposed in the work by O'Regan's group, it is reasonable to assume that the above reported segregation phenomena occurring in water electrolytes should not occur when non-mesoporous films are considered. Nevertheless, to the best of our knowledge and the present literature reports, no scientific group has undertaken this research path so far.

Other literature reports showed as well that the presence of a specifically selected amount of water in the electrolyte can increase the PV conversion efficiency. Weng and coworkers observed that, by increasing the water concentration up to 2.2 M, $V_{\rm oc}$ and FF values increased monotonically, while $J_{\rm sc}$ showed a continuous decrease, as reported in Table 2.124 Efficiency increased from 3.8% to 4.5% in the presence of a 1.7 M H₂O solution in a standard liquid electrolyte (LiI 0.50 M and I_2 50 mM in MPN/H₂O). The authors made the following hypotheses regarding the reasons for the $V_{\rm oc}$ improvement: an increased electron injection efficiency, a decreased electron density and a retardation of decay rate. Time-resolved infrared absorption spectroscopy, a useful technique to directly detect the CB as well as the trapped electrons,¹²⁵⁻¹²⁷ showed that - in the presence of water - the optical phonon scattering, which indicates surface trapped states, was observed. It revealed that hydroxide group interaction with TiO2 gave rise to an increase in the surface-trap states.¹¹⁴ The interfacial back electron transfer process was retarded in an aqueous environment, and the electron injection efficiency increased up to 1.7 M of water.

Mixtures of water and organic solvents have been recently investigated by Frank and coworkers.¹²⁸ Regardless the water content, Table 3 shows that V_{oc} , FF and η values were higher for the ACN:VAN-based DSSCs than for the MPN-based counterparts,

Table 2PV parameters ($P_{in} = 1 \text{ sun}$) of DSSCs assembled with electrolytesprepared at various concentrations of water. Electrolytes: Lil 0.50 M and I250 mM in MPN/H2O. Dye: N3. Adapted and reprinted from ref. 124

$H_2O(M)$	$J_{\rm sc}~({ m mA~cm^{-2}})$	$V_{\rm oc}$ (V)	FF	η (%)
0	16.5	0.47	0.49	3.8
0.6	15.8	0.49	0.53	4.1
1.2	15.1	0.52	0.55	4.3
1.7	14.1	0.54	0.59	4.5
2.2	12.1	0.55	0.62	4.1

Table 3 PV parameters ($P_{in} = 1 \text{ sun}$) of DSSCs assembled with electrolytes containing different amounts of water and organic solvents. Electrolyte: PMII 1.0 M, I₂ 0.15 M, GuSCN 0.10 M and NBBI 0.50 M. Dye: Z907. Adapted and reprinted from ref. 128

Organic solvent	H ₂ O (vol%)	$J_{\rm sc}~({\rm mA~cm^{-2}})$	$V_{\rm oc}$ (V)	FF	η (%)
MPN	0	15.77	0.647	0.658	6.72
MPN	10	16.15	0.662	0.643	6.87
ACN:VAN	0	14.63	0.661	0.725	7.01
ACN:VAN	10	15.67	0.680	0.706	7.52

while J_{sc} followed the opposite trend. As regards water addition in the liquid electrolyte (PMII 1.0 M, I2 0.15 M, GuSCN 0.10 M and NBBI 0.50 M; NBBI = N-butylbenzimidazole), it enhanced both J_{sc} and V_{oc} for any of the organic solvents used, but lowered FF values. Such a behaviour was quite different with respect to what reported by Weng and coworkers,124 who used N3 dye instead of Z907. Further investigation was carried out by means of electrochemical impedance spectroscopy (EIS) under bias light at 680 nm. It showed that the addition of water decreased the difference in the potential values between $E_{\rm c}$ and E of about 28 mV: this resulted from a downward shift of 10 mV for E toward positive electrode potentials, along with a 38 mV downward shift of the TiO₂ CB. All of these changes in the potential values of the different energy levels were attributed to an increased proton concentration at the photoanode surface.^{129,130} By combining transport and recombination measurements, it was evidenced that the addition of water increased the chargecollection efficiency.^{131,132} This effect, together with the higher charge-injection efficiency derived from the downshift of the TiO_2 CB upon water addition,¹³³ led to improved J_{sc} values. Frank and coworkers also expected a reduced $V_{\rm oc}$ due to the lower gap in the potential between E_c and E; however, the addition of water increased the photovoltage of about 15-19 mV (Table 3). Such a behaviour was justified by considering the decreased recombination in aqueous cells by a factor of 4-5.86,134 As regards the decreased FF values, an increase in the dark exchange current was detected, which was attributed to a larger dark electron concentration in the presence of water.⁸⁴ Outstandingly, the J-V characteristics of the DSSCs assembled using electrolytes prepared both with and without the addition of water showed essentially the same aging behaviours upon 1000 h of continuous light soaking. This represents a key issue, quite in contrast with what reported in Section 3: does the presence of water have an influence on the cell stability or not? Dye nature, electrolyte composition or dye-electrolyte combination: what the preponderant factor is? From Frank and coworkers study,

 $\begin{array}{lll} \textbf{Table 4} & \text{PV} \text{ parameters} \ (P_{in} = 1 \ \text{sun}) \ \text{of DSSCs} \ \text{assembled} \ \text{with electrolytes} \\ \text{based on water and/or ethanol-water mixtures recovered from industrial} \\ \text{effluents. Electrolyte: KI 0.50 M and I}_2 \ \text{25 mM.} \ \text{Adapted and reprinted from} \\ \text{ref. 138} \end{array}$

Solvent	Dye	$J_{ m sc} ({ m mA} \ { m cm}^{-2})$	$V_{\rm oc}$ (V)	FF	η (%)
H ₂ O	N3	2.70	0.37	0.60	0.6
H ₂ O	N719	2.00	0.40	0.63	0.5
$H_2O: EtOH 65: 35$	N3	4.68	0.47	0.59	1.3
$H_2O:EtOH 65:35$	N719	4.04	0.48	0.57	1.1

it seems that the adoption of a thermally stable electrolyte solvent (MPN/H₂O) and a hydrophobic dye (Z907) can lead to equally stable water-free and water-based DSSCs.

The remarkable advantage in terms of safety as well as ecocompatibility of replacing organic solvents by water has been already detailed in the introductory section. However, it should be also considered that some industrial processes (*i.e.*, biotechnology and food industry) produce water-ethanol mixtures as effluents, the separation of which is not economically advantageous. The recovery of these wastewaters surged as a hot research topic only recently, ^{135–137} after the pioneering work of Miyasaka and coworkers, who proposed their reuse in aqueous DSSCs.¹³⁸ Table 4 shows the enhancement of J_{sc} values when using EtOH-H₂O solutions, which was attributed to the activation of the mesoporous TiO₂/electrolyte (KI 0.50 M and I₂ 25 mM) interface. Pristine photoanodes were found not to be hydrophilic enough to absorb water into their interior pores, thus the addition of alcohols improved their wettability because of the higher affinity with the surface functionalities present at the inner surface of the mesopores. Moreover, the addition of ethanol reduced both the surface tension and the viscosity of the aqueous electrolyte, thus increasing the effective area of the electrochemically active TiO2-electrolyte interface with respect to the pure aqueous system, in which a hydrogen-bonded large cluster of water molecules could not effectively penetrate the whole mesoporous structure. As a result, efficiency values were doubled

The brief analysis of the recent scientific reports has evidenced a real turnaround in the DSSC research community: water is no longer intended to be a poisoner to be utterly avoided, in fact it started to be gradually introduced in progressively larger amounts. This novel trend is accompanied by electrochemical and spectroscopic investigation of the new phenomena and new interfaces, to guarantee the useful insights towards the improvement of the device characteristics. As will be explained in the following sections, each cell component is currently the subject of thorough investigation from the physical as well as engineering viewpoint in order to ensure its highest possible efficiency in an aqueous environment.

6. Additives for aqueous electrolytes

Nowadays, it is widely known that the introduction of specifically selected additives in the electrolyte may guarantee remarkable improvements in the PV performance of a DSSC.^{139–141} Indeed, for the same dye, while J_{sc} is closely related to the type and concentration of the redox mediator under use, V_{oc} can be ameliorated by introducing nitrogen-containing heterocyclic compounds, such as pyridines, aminotriazole, pyrimidine, aminothiazole, pyrazole and quinoline. Above all, in 1993 Grätzel and coworkers introduced TBP.⁸¹ Being adsorbed onto the surface of the TiO₂ photoelectrode, it prevented the leakage of the injected electrons in the electrolyte and also prevented I_3^- from contacting the surface of the photoanode. As reported in the course of this review, TBP has been already introduced in some aqueous electrolytes; however, it showed rather low solubility in water. Thus, alternative additives have been proposed for aqueous DSSCs (in this section) or their solubilisation has been achieved through the introduction of surfactants (see Section 7).

N-Alkylbenzimidazole derivatives represent another category of organic additives which were found to behave similarly to TBP. Indeed, N-methylbenzimidazole (NMBI) is often added in liquid electrolytes both for TiO2-142 and ZnO-sensitized DSSCs.143,144 Yang and coworkers proposed several bis-benzimidazole derivatives containing an ethylene glycol repeating unit (BBEG_n, see Fig. 10) for aqueous DSSCs (DMPII 0.50 M, LiI 0.10 M, I2 50 mM and TBP/BBEG_n 0.50 M in ACN/H₂O; DMPII = 1,2-dimethyl-3propylimidazolium iodide) and evaluated the cell durability.145 As listed in Table 5, V_{oc} and FF values were found to be higher in all the water-based cells. This was ascribed to the strong adsorption of water onto the TiO2 surface, which blocked the reaction between I₃⁻ ions and injected electrons; moreover, as previously stated by O'Regan's group,120 the electrochemical potential of the iodine-based redox mediator in water-based cells was positively shifted. As a drawback, the addition of water led to the detachment of the adsorbed N719 dye molecules, thus limiting the flux of the injected electrons from the excited states of the dye which adversely influenced J_{sc} values.

As shown in Table 5, the novel $BBEG_n$ additives proposed by Yang and coworkers did not guarantee improved performances if compared to TBP. Nevertheless, the resulting devices demonstrated enhanced stability upon the aging test, while TBP-based cell efficiency dropped significantly (62% in 135 h). Indeed, the chemical structure of $BBEG_n$ additives incorporated ethylene glycol linkages between two benzimidazoles, which produced H-bond bonding sites for the water molecules. In this way, ethylene glycol linkages sequestered the water molecules, thus preventing their interaction with the TiO₂-dye linkage as well as the dye –SCN functional group. Moreover, the $BBEG_n/H_2O$ phase solvated I_3^- ions, thus decreasing their concentration in the dyed TiO₂ surface and avoiding back electron transfer phenomena. Summarising, by the proper use of specifically selected and optimised additives, aqueous DSSCs may become

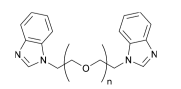


Fig. 10 Chemical structure of benzimidazole-based additives (BBEG_n) used in aqueous DSSCs, ref. 145.

Table 5PV parameters ($P_{in} = 1 \text{ sun}$) of DSSCs assembled with electrolytesbased on pure ACN or ACN-H2O mixtures. Electrolytes: DMPII 0.50 M, LiI0.10 M, I2 50 mM and TBP/BBEGn 0.50 M. Dye: N719. Adapted andreprinted from ref. 145

Additive	H ₂ O (vol%)	$J_{ m sc} ({ m mA} \ { m cm}^{-2})$	$V_{\rm oc}$ (V)	FF	η (%)
TBP	0	14.99	0.73	0.65	7.14
TBP	10	5.05	0.80	0.70	2.82
$BBEG_n, n = 1$	0	11.31	0.78	0.60	5.27
BBEG _n , $n = 1$	10	4.19	0.80	0.69	2.31
BBEG _n , $n = 3$	0	13.04	0.78	0.62	6.29
BBEG _n , $n = 3$	10	5.75	0.80	0.64	2.96

as much stable as (or eventually even more stable than) their standard aprotic counterparts. In this respect, the search for new additives for water-based liquid electrolytes is currently under intensive investigation in many research laboratories.

7. Surfactants: key ingredients for aqueous DSSCs

An electrolyte of a DSSC is a fairly complex chemical system: the solvent should be able to solubilise both inorganic salts and non-polar species (i.e., iodine and additives such as TBP and NMBI). When the aqueous electrolyte contains a mixture of water and organic solvents, homogeneity and stability of all the components in solution are fairly easy to be obtained (once optimized their concentrations), as described so far. On the other hand, if a fully aqueous DSSC has to be fabricated, a way to solubilise all of the aforementioned electrolyte components has to be found. This is a particularly complex work, which requires the modification or the functionalization of the traditionally used chemical compounds, or the introduction of a new class of additives: surfactants. The ability of surfactants in lowering the surface (or interfacial) tension between two liquids or between a liquid and a solid is well-established;¹⁴⁶ their application as detergents, wetting agents, emulsifiers, foaming agents and dispersants are vitally important in our daily life.147,148 The scientific community considers surfactants as key ingredients for drug delivery systems¹⁴⁹ as well as environmental decontamination and rehabilitation procedures.¹⁵⁰ Several applications are also envisaged in the field of energy conversion and storage¹⁵¹ due to the unique properties of surfactants particularly in the synthesis and functionalization of porous anodic^{152,153} and cathodic154,155 nanostructures.

For what concerns aqueous DSSCs, surfactants have initially been used to segregate water molecules within the micellar phase, thus promoting its solubility in the organic liquid electrolyte. The use of Triton X-100, $C_{14}H_{22}O(C_2H_4O)_n$, a nonionic surfactant having a hydrophilic polyethylene oxide chain and an aromatic hydrocarbon lipophilic or hydrophobic group, was proposed by Kim and coworkers.¹⁵⁶ They introduced Triton X-100 in the 20 mM concentration into an aqueous liquid electrolyte (LiI 0.10 M, HDMII 0.60 M, I₂ 50 mM and TBP 0.50 M in MPN/H₂O, HDMII = 1-hexyl-2,3-dimethylimidazolium iodide), and observed increased V_{oc} and FF values while increasing the amount of water (0.0–4.4 M); conversely, J_{sc} monotonically decreased. A peak efficiency value of 5.9% when $[H_2O] = 2.2$ M was obtained (Table 6).

Three hypotheses may be elaborated to justify the increased $V_{\rm oc}$ values observed in surfactant-laden liquid electrolytes:

– The reduction in the back electron transfer from the TiO_2 CB to the I_3^- ions in the electrolyte: in fact, I_3^- ions have higher solubility in the hydrophilic Triton X-100/H₂O phase than at the interface with the photoanode.

– The negative shift of the flat band potential of TiO₂: some of the protons transferred by the sensitizer (which contains –COOH groups) to the TiO₂ surface are removed by the hydrophilic Triton X-100/H₂O phase, thus leading to a shift of the flat band potential which becomes less positive than that occurring in the absence of the surfactant-added aqueous phase.^{157,158}

- The positive shift of E.

 $J_{\rm sc}$ decreased because of the reduced number of injected electrons, due to the improved $V_{\rm oc}$ caused by the increase of the barrier height for their injection.¹⁵⁹ Interestingly, the trend observed for the photocurrent differed from that expected by Kim and coworkers by means of chronoamperometry and EIS measurements. Indeed, an increase of the limiting current for the oxidation of I⁻ was detected. This accounted for the enhancement in the diffusion of I⁻ ions in the presence of Triton X-100 and water, as also indicated by the decrease in the series ($R_{\rm s}$) as well as charge transfer resistance at the counter electrode ($R_{\rm CE}$) measured for aqueous DSSCs (Table 6). Along with the observed decrease in viscosity, this should have resulted in improved $J_{\rm sc}$ and FF values.

The long-term stability of Triton X-100-added DSSCs was surprisingly high (-19% after 7 days under dark), even higher than the corresponding aprotic cell (-67%), thanks to the increased stability of $J_{\rm sc}$. In fact, Triton X-100 was able to strongly retain the molecules of solvent, thus suppressing their evaporation; its aqueous phase could even extract the water traces which were inevitably introduced during cell fabrication.¹⁶⁰ The harmful effects caused by the presence of water during irradiation (*i.e.*, hydrogen evolution and photocatalytic processes) were avoided, thus demonstrating that a surfactant-added aqueous DSSC can perform better than a standard cell. However, for the sake of the reader we must point out that the cells of this article were not sealed with a rigorous procedure; indeed, a 67% efficiency decrease in a week only is not a data comparable with the state of the art of sealed liquid DSSCs.^{161,162}

Data reported in Section 5 show that the performance of 100% water-based DSSCs is still low compared with those of

Table 6 PV ($P_{in} = 1 \text{ sun}$) and electrochemical parameters of DSSCs assembled with electrolytes containing different amounts of water. Electrolytes: Lil 0.10 M, HDMII 0.60 M, I₂ 50 mM, TBP 0.50 M in MPN/H₂O. Dye: N719. Adapted and reprinted from ref. 156

$H_2O(M)$	$J_{\rm sc}$ (mA cm ⁻²)	$V_{\rm oc}$ (V)	FF	η (%)	$R_{\rm s}\left(\Omega\right)$	$R_{\rm CE} \left(\Omega \ {\rm cm}^{-2} \right)$
0.0	13.44	0.74	0.53	5.3	20.97	2.74
1.1	12.10	0.79	0.57	5.4	20.90	2.18
2.2	11.77	0.81	0.62	5.9	20.79	2.12
3.3	11.65	0.82	0.59	5.7	20.83	1.64
4.4	10.38	0.81	0.61	5.1	20.74	1.36

traditional DSSCs, and one of the reasons is surely the incomplete wetting of the dye-coated TiO₂ interface by the aqueous electrolyte. In this context, it is well known that the addition of a surfactant increases the wettability and minimizes the separation between materials in two different phases. Yan and coworkers investigated a series of surfactants in organic solvent-free DSSCs (NaI 2.0 M, I₂ 0.20 M and GuSCN 0.10 M): hexadecyltrimethylammonium bromide (CTAB), N,N,N-trimethyl-3-(perfluorooctyl sulfonamido)propan-1-aminium iodide (FC-134), bis(2-ethylhexyl) sulfosuccinate sodium salt (AOT) and triethylammonium perfluorooctane sulphonate (FK-1).¹⁶³ As can be seen in Table 7, both anionic (AOT, FK-1) and cationic (CTAB, FC-134) surfactants dramatically improved the PV performances of N719-sensitized DSSCs. At the same time, higher charge-transfer resistance values were measured at the TiO₂/electrolyte interface, and the resulting reduced charge recombination and increased electron lifetime were attributed to the enhanced interfacial compatibility due to the addition of the surfactants.^{164,165} On the other hand, Mott-Schottky experiments and capacitance measurements suggested that the surfactant addition did not shift the TiO₂ CB edge, thus contradicting what previously reported by Kim and coworkers.¹⁵⁶ In the presence of surfactant-added aqueous electrolytes, photoanodes showed reduced contact angle (CA) values compared to the corresponding surfactant-free ones (Fig. 11), thus further demonstrating the improved wettability of the photoanode/ electrolyte interface.^{166,167}

Sunlight-to-electricity conversion of surfactant-added DSSCs was improved under low P_{in} ; as an example, the efficiency of the FC-134 (0.2 wt%)-laden cell increased from 3.96 to 4.66% when Pin decreased from 1.0 to 0.5 sun. The lower efficiency yielded under full sunlight irradiation intensity, maybe due to the inefficient charge screening of the electron transport in the mesoporous TiO_2 , ^{168,169} paved the way to the use of aqueous cells in a relatively dark environment, where DSSCs are increasingly establishing themselves as the leading device.^{170,171} As regards the long-term stability, the surfactant-free cell provided only 17% of the initial efficiency after 50 days under constant 1.0 sun illumination (at room temperature, RT, with UV-filter), where the CTAB-laden one was able to retain the 48%. Such an improvement was assigned to the surfactant molecules adsorbed at the surface of the photoanode. The resulting coating layer prevented I_3^- ions from contacting the dye-coated TiO₂ active material particles, reduced the recombination rate and avoided the sensitizer degradation by enhancing the dye-regeneration rate.

Surfactant	$J_{\rm sc}~({ m mA~cm^{-2}})$	$V_{\rm oc}$ (V)	FF	η (%)
None	7.50	0.51	0.66	2.51
CTAB (0.1 wt%)	8.94	0.51	0.69	3.14
AOT (0.1 wt%)	8.44	0.50	0.71	2.98
FK-1 (0.1 wt%)	9.99	0.51	0.69	3.55
FC-134 (0.1 wt%)	9.69	0.51	0.70	3.56
FC-134 (0.2 wt%)	10.97	0.53	0.68	3.96

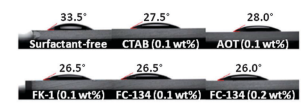


Fig. 11 CA measurements of different aqueous electrolytes at the surface of dye-coated TiO_2 films. Adapted and reprinted from ref. 163.

Due to its positive effect, AOT was recently employed by Jang and coworkers with the aim of controlling both the total and the local concentration of the redox couple (BMII 0.70 M, I2 30 mM, GuSCN 0.10 M and TBP 0.50 M in ACN/VAN; BMII = 1-butyl-3methylimidazolium iodide).¹⁷² By means of its -SO₃⁻ group,¹⁷³ AOT can be chemisorbed onto the TiO₂ nanoparticles, thus hindering the access of $\mathrm{I_3}^-$ anions and facilitating the access of I⁻ ones to enhance the dye regeneration rate. Upon concurrent addition of water, the absorbance of I_3^- at 363 nm decreased due to the triiodide hydrolysis reaction: one of the resulting products was I⁻, which further assisted the dye regeneration efficiency (from 71% to 78%). Table 8 shows that such a combined effect of water and AOT led to the improvement in both J_{sc} and V_{oc} values; the $R_{\rm CE}$ was improved as well, thus demonstrating that the triiodide reduction reaction at the counter electrode/electrolyte interface occurred much more efficiently in the AOT/water-laden devices. At present, the 10.8% efficiency value obtained by Jang and coworkers (see the insight outlined in Fig. 12) is the highest

Table 8 PV parameters ($P_{in} = 1 \text{ sun}$) of DSSCs assembled with electrolytes laden with water and/or AOT. Electrolyte: BMII 0.70 M, I₂ 30 mM, GuSCN 0.10 M, TBP 0.50 M in ACN/VAN. Dye: N719. Adapted and reprinted from ref. 172

Additives	$J_{\rm sc} ({ m mA} { m cm}^{-2})$	$V_{\rm oc}$ (V)	FF	η (%)	$R_{\mathrm{CE}}\left(\Omega\right)$
None	15.3	0.832	0.73	9.3	6.93
AOT (1.0 mM)	16.5	0.829	0.74	10.1	4.93
H_2O (10 mM)	16.9	0.850	0.71	10.2	6.89
AOT + H_2O	17.5	0.850	0.73	10.8	4.83

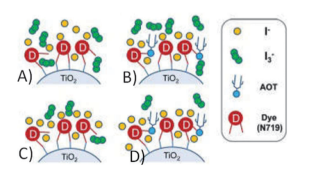


Fig. 12 Local concentration control of the I_3^{-}/I^{-} redox couple by the hydrogen bonding interaction between the AOT and the carboxyl group of the N719 dye. (A) AOT- and water-free system; (B) in the presence of AOT, I_3^{-} is hindered from approaching the TiO₂/dye surface; (C) accelerated hydrolysis of I_3^{-} by water; (D) AOT and water are simultaneously added to reduce the recombination rate and to enhance the dye regeneration yield. Adapted and reprinted from ref. 172.

value ever measured for a DSSC containing water. However, we cannot call it "aqueous DSSC", as the amount of water added to the electrolyte was minimal (10 mM).

8. Exploiting water to introduce new redox couples

As stated in Section 2, several redox mediators have been proposed during the past decades;⁵⁰ among others, the iodide/ triiodide couple⁵¹ and cobalt complexes²⁹ have proven to be the most stable and the best performing, respectively. Nevertheless, due to their ability in solubilising compounds which are insoluble in conventional aprotic organic solvents, the use of water as a solvent opened up new possibilities in the preparation and selection of the redox mediator.

The first exotic redox couple for aqueous DSSCs was a ceriumbased compound. It was proposed by Teoh and coworkers in 2008.¹⁷⁴ Ce(NO₃)₃ 0.10 M and Ce(NO₃)₄ 50 mM were dissolved in a 35:65 EtOH: H₂O mixture. This electrolyte was coupled with a variety of commercial and natural sensitizers, such as crystal violet, mercurochrome, chlorophyll, extracts of Bongainvillea brasiliensis, Garcinia suubelliptica, Ficus Reusa and Rhoeo spathacea. Indeed, due to their simple preparation techniques, wide availability and low cost, natural dyes could be the best option for an aqueous, thus truly eco-friendly solar energy conversion device. As a further insight, Teoh and coworkers used their electrolyte with a 38 nm-thick Schottky barrier¹⁷⁵ composed of Au NPs deposited onto the photoanode, in order to improve the efficiency of the electron injection.^{176,177} As a result, electrons in the LUMO of the dye passed through the Au thin layer by tunnelling the TiO_2 CB, thus avoiding the electron recombination reaction. J_{sc} and V_{oc} values measured in the presence of commercial dyes were found to be lower than those of natural dyes; the highest efficiency $(\eta = 1.49\%, J_{sc} = 10.9 \text{ mA cm}^{-2}, V_{oc} = 0.496 \text{ V}, \text{ FF} = 0.27) \text{ was}$ obtained in the presence of Rhoeo spathacea (RhS).

A cerium-based aqueous electrolyte (30:70 in EtOH:H₂O) was also coupled with natural pigments, such as the green and the red parts of *Codiaeum varie* and *Aglaonema*. Moreover, with the aim of improving the photovoltage, Su and coworkers¹⁷⁸ coupled this system with a ZrO₂ photoanode characterised by a wide band gap energy (5.8 eV) and a CB higher than that of TiO₂.^{179,180} When the green part of *Codiaeum varie* (CV-G) was used, a $V_{\rm oc}$ of 0.624 V was achieved, definitely higher than those usually measured for natural pigment-sensitized TiO₂-based DSSCs.¹⁸¹ Furthermore, the effective dielectric constant and refractive index of the ZrO₂ layer were experimentally tuned, and the light transmission of the photoanode was improved, resulting in a cell efficiency of 0.69% ($J_{\rm sc} = 0.52$ mA cm⁻², FF = 0.53).

Working in aqueous media allows the application of redox couples such as $Fe(CN)_6^{4-/3-}$, which can overcome iodate formation, light absorption and corrosiveness typical of standard iodine-based mediators. In 2012, Spiccia and coworkers introduced the ferrocyanide/ferricyanide redox couple in a truly water-based electrolyte: 0.40 M K₄Fe(CN)₆, 40 mM K₃Fe(CN)₆, 0.10 M KCl and 0.1 vol% Tween 20, dissolved in H₂O at pH 8.¹⁸² The redox

Table 9 PV parameters ($P_{in} = 1 \text{ sun}$) of DSSCs assembled with Fe(CN)₆^{4-/3-} and I⁻/I₃⁻-based electrolytes, in the presence of different solvents. Dye: MK-2. Adapted and reprinted from ref. 182

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Redox couple	Solvent	$J_{\rm sc}~({ m mA~cm^{-2}})$	$V_{\rm oc}$ (V)	FF	η (%)
${\rm Fe}({\rm CN})_{6}^{4-/3-}$	H ₂ O	7.21	0.761	0.75	4.10
${\rm I}^{-}/{\rm I}_{3}^{-}$	ACN	11.86	0.769	0.66	6.05

potentials of $Fe(CN)_6^{4-/3-}$ and I^-/I_3^- being similar,¹⁸³ a typical hydrophobic carbazole dye (MK-2) was used; besides showing an high molar extinction coefficients,¹⁸⁴ it allowed the use of relatively thin TiO₂ films, thus resulting in reduced interfacial area for recombination as well as improved charge extraction. Moreover, it showed only faint signs of desorption when stored under water for 30 days (conversely, the complete desorption of Z907 dye was observed in the same conditions). $Fe(CN)_6^{4-/3-}$ based DSSC showed an efficiency equal to 4.1% (Table 9); J_{sc} was the only value which decreased with respect to the corresponding I⁻/I₃⁻/ACN-based cell (MPII 0.60 M, LiI 0.10 M, I₂ 0.20 M and TBP 0.50 M; MPII = 1-methyl-3-propylimidazolium iodide). It was supposed that ferrocyanide, being able to sensitize TiO₂,¹⁸⁵ contributed as a co-sensitizer, but its performance was obviously lower than that of MK-2. By means of intensity modulated photovoltage and photocurrent spectroscopies, also combined with charge extraction analysis,¹⁸⁶ it was demonstrated that the $Fe(CN)_6^{4/3-}$ -based electrolyte guaranteed a three orders of magnitude faster rate of recombination if compared to the reference couple. Besides, no difference in $V_{\rm oc}$ values was observed, which was related to the negative CB shift of 150 mV induced by $Fe(CN)_6^{4-/3-}$. The only drawback of this new redox couple was the performance decay measured under unfiltered white light illumination, which was caused by the well-known photolysis and photocatalytic decomposition of the ferrocyanide/ ferricyanide redox couple.^{187,188} By introducing a 480 nm long-pass filter, no performance decay was observed. Anyway, modification of the TiO2 surface or its replacement with an alternative wide bandgap semiconductor should be considered for the fabrication of cells stable under full solar irradiation; otherwise, also the replacement of the cyanide ligands with stronger binding units may be taken into account.

Experimental investigation showed that the poor solubility of I₂ in water is the major limiting factor affecting aqueous DSSC performance. Even if this drawback can be solved by adopting suitable surfactants (see Section 7), the challenge of finding non-corrosive redox couples, which may also guarantee weak light absorption in the visible light region, has encouraged researchers to test sulphur-based systems, such as thiolate/ disulphide.^{189–192} Despite most of these species demonstrating rather low water solubility, Sun and coworkers identified a watersoluble thiolate/disulphide redox couple. It consisted of 1-ethyl-3-methylimidazolium 4-methyl-1,2,4-triazole-3-thiolate (TT⁻EMI⁺) and 3,3'-dithiobis[4-methyl-(1,2,4)-triazole] (DTT) (Fig. 13,¹⁹³). The resulting truly aqueous electrolyte (TT⁻EMI⁺ 0.20 M, DTT 0.20 M, TBP 0.50 M and Triton X-100 1 vol%) was coupled with a hydrophobic dye having a short methoxyl chain in the donor unit (D45, Table 14), which increased the probability of interaction with

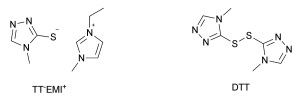


Fig. 13 Structures of the TT^-EMI^+/DTT -based redox couple. Adapted and reprinted from ref. 193.

Table 10 PV parameters ($P_{in} = 1 \text{ sun}$) of DSSCs assembled with TT⁻EMI⁺/DTT- and I⁻/I₃⁻-based electrolytes, in the presence of different dyes and solvents. Adapted and reprinted from ref. 193

Redox couple	Solvent	Dye	$J_{\rm sc} ({ m mA} { m cm}^{-2})$	$V_{\rm oc}$ (V)	FF	η (%)
TT ⁻ EMI ⁺ /DTT TT ⁻ EMI ⁺ /DTT			7.2 9.5		$0.55 \\ 0.59$	
$TT^{-}EMI^{+}/DTT$				0.570	0.53	
I^{-}/I_{3}^{-}	ACN:VAN	D45	11.7	0.790	0.61	5.6

the redox system. An efficiency equal to 2.6% was recorded (Table 10). Replacing water with ACN did not modify E, but led to the desorption of the dye from the TiO2 surface. When compared to a traditional I⁻/I₃⁻-based system (DMHII 0.60 M, LiI 60 mM, I₂ 40 mM and TBP 0.40 M in ACN/VAN; DMHII = 3-hexyl-1,2-dimethylimidazolium iodide), the aqueous electrolyte showed remarkably reduced diffusion coefficients, being $D(TT^{-})$ and D(DTT) equal to 4.01 imes 10⁻⁶ and 1.87 imes 10⁻⁶ cm² s⁻¹ respectively, while $D(I^{-})$ and $D(I_{3}^{-})$ were equal to 1.70 \times 10⁻⁵ and $1.10 \times 10^{-5} \text{ cm}^2 \text{ s}^{-1}$, respectively, thus almost one order of magnitude different. However, since PV measurements under different light intensities did not show any enhancement in the performance under weak light intensity, low electrolyte diffusion coefficients were not likely to be responsible for the low J_{sc} recorded; on the other hand, it was observed that D45 regeneration was slowed down by a factor of four in the presence of TT⁻EMI⁺/ DTT. When D45 dye was replaced by D51 (Table 14), efficiency jumped up to 3.5% (Table 10), which was attributed to the higher extinction coefficient and broader absorption spectrum of the sensitizer. During the stability test, aqueous DSSCs were able to assure only 63% of their initial efficiency after 4 h of light soaking at RT. However, despite FF values decreasing due to the poor stability of DTT in solution, Sun and coworkers highlighted that the optimization of the electrolyte composition could be considered as an effective strategy to overcome stability issues in aqueous DSSCs.

As previously stated, the recent efficiency records of DSSCs have been obtained using cobalt complexes as redox mediators.^{29,35} Being transition metal ion complexes, they are highly soluble in water which accounts for their suitable application in aqueous DSSCs. Indeed, Spiccia and coworkers easily dissolved 0.20 M of $[Co(bpy)_3]^{2+}$ and 40 mM of $[Co(bpy)_3]^{3+}$ in water, along with the addition of NMBI (0.70 M) as a V_{oc} -booster and various amounts of poly(ethylene glycol) (PEG 300) to minimize phase separation between the hydrophobic dye (MK-2) and the aqueous electrolyte.¹⁹⁴ A bell-shaped performance trend was observed as a function of the amount of PEG 300 added; the best energy conversion efficiency of 4.2% ($J_{sc} = 8.3 \text{ mA cm}^{-2}$, $V_{oc} = 0.685 \text{ V}$,

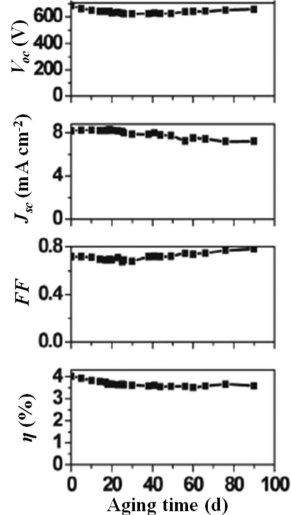


Fig. 14 Long-term stability under dark of a DSSC assembled with an aqueous electrolyte composed of $[Co(bpy)_3]^{2+}$ 0.20 M, $[Co(bpy)_3]^{3+}$ 40 mM, NMBI 0.70 M and PEG 300 1 vol%. Dye: MK-2. Adapted and reprinted from ref. 194.

FF = 0.72) was achieved upon addition of 1 wt% PEG 300. Noteworthy, such an additive was able to concurrently guarantee the highest possible electron recombination resistance and the lowest diffusion resistance of the redox species. The durability of cobalt-based electrolytes being a topic of great interest right now,¹⁹⁵ aqueous DSSCs were investigated for their stability under prolonged aging conditions by Spiccia and coworkers, demonstrating a limited 10% decrease of the initial efficiency after more than three months under dark (Fig. 14).

9. Photoanode activation: a way to improve the interfacial characteristics between electrodes and aqueous electrolytes

Nowadays, it is widely established that the interface between the photoanode and the aqueous electrolyte is a key point remarkably affecting the efficiency of water-based DSSCs. This is in fact an important issue even for standard DSSCs, where the surface modification of the photoanode is usually carried out by means of barrier layers,¹⁹⁶ N-doping,¹⁹⁷ hybrid organicinorganic linkers,¹⁹⁸ nanodecorations¹⁹⁹ and physical techniques.²⁰⁰ Furthermore, for the first time, the solid electrolyte interphase formation (SEI layer, well known and investigated in the field of Li-ion batteries,²⁰¹) has been recently evidenced even in DSSCs, which implies a novel series of challenging research goals for the scientific community.²⁰²

A simple interfacial activation for aqueous DSSCs was proposed by Miyasaka and coworkers, by means of treating the mesoporous layer with gaseous active oxygen prior to dve adsorption.²⁰³ In particular, the TiO₂ layer was concurrently etched by air rich in O_3 (300 ppm) and exposed to UV light for 60 min. Water CA values decreased from 71 to 22°, which highlighted the improved hydrophilicity of the electrode. An efficient permeation of the aqueous redox electrolyte (KI 0.50 M and I₂ 25 mM in H₂O:EtOH 65:35) through the mesoporous electrode structure was achieved; both $J_{\rm sc}$ and $V_{\rm oc}$ increased, from 4.94 to 5.80 mA cm⁻² and from 0.55 to 0.60 V, respectively. As a result, efficiency noticeably increased from 1.7% to 2.2%. Noteworthy, TBP (5 vol%), being insoluble in water, was introduced in the dye-uptake solution and not in the electrolyte. Authors hypothesized that TBP could be considered as a base and, when adsorbed onto the TiO₂ surface, could improve the rate of deprotonation of the carboxyl groups of the Ru complex to reinforce the bonding between dye and TiO₂, thus leading to improved electron injection efficiencies. In other words, TBP provided itself as a ligand for the Ru complex, forming a mixed ligand complex with tetrabutylammonium (TBA).

An alternative way to modify the photoanode could be the introduction of a Schottky barrier, which can increase the electron injection efficiency.¹⁷⁵ To this purpose, Lai and coworkers prepared Au NPs modified by tetraoctylammonium bromide, which were then anchored at the TiO2 surface by means of (3-mercaptopropyl)trimethoxysilane.²⁰⁴ A self-assembly monolayer process was performed and, by successively repeating the preparation steps, a layer-by-layer Au NPs coating was formed on the conducting substrate (Fig. 15). Being the I^{-}/I_{3}^{-} couple corrosive towards Au, a Fe²⁺/Fe³⁺ redox mediator was used, despite its high sensitivity to hydrolysis in aqueous solutions due to the formation of iron aquo complexes with water. To possibly avoid such a drawback, mixtures of H₂O/EtOH in the 65:35 ratio were selected for the preparation of the electrolyte (FeCl₂ 0.10 M, FeCl₃ 50 mM and LiNO₃ 0.10 M).²⁰⁵ In the presence of four Au NP layers, J_{sc} more than doubled (*i.e.*, from 2.76 to 5.96 mA cm⁻²); FF increased as well (*i.e.*, from 0.23 to 0.35). However, the efficiency of the aqueous DSSC remained rather low (i.e., 0.95% against 0.26% for the Au-free cell), maybe because of the redox mediator instability and/or the selection of the dye (merbromin, MB, hardly known for its high efficiency).

In liquid DSSCs, the diffusive nature of the ion flow throughout the electrolyte creates an upper limit to the current (diffusion limited current, J_{dl}) that can pass across the cell. It is controlled by the diffusion coefficient of the limiting species **Review Article**

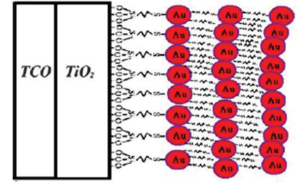


Fig. 15 Layer-by-layer Au NP assembly onto the TiO_2 electrode to act as a Schottky barrier in aqueous DSSCs. Adapted and reprinted from ref. 204.

 $(I_3^- \text{ ions})$, their concentration, the cell thickness as well as the thickness and morphology of the TiO₂ photoanode. While in organic solvent-based liquid electrolytes the J_{dl} values (~30 and \sim 100 mA cm⁻² in MPN and ACN, respectively) are larger than $J_{\rm sc}$ values, under incomplete wetting conditions the reduced cross sectional area through which ions can diffuse causes lower J_{dl} values. Since the latter situation may occur in aqueous DSSCs, O'Regan and coworkers proposed several strategies to improve the rather critical wettability of Z907-dyed TiO₂.^{206,207} Among the different surfactants being investigated, chenodeoxycholic acid (CDCA) demonstrated the highest values of $J_{\rm sc}$ (3 fold increase) and $J_{\rm dl}$ (10 fold increase): this beneficial effect was attributed both to the hydrophobic centre portion of the molecule (compatible with Z907) and the hydrophilic -OH groups. Despite the improved wetting ability, the low J_{sc} value $(\sim 2 \text{ mA cm}^{-2})$ indicated that some limitation could be derived from injection and/or collection processes. Even if the main recombination pathway has recently been proposed to be the reduction of free iodine (rather than the triiodide),²⁰⁸ it was noted that the binding coefficient (K_M) of I^- and I_2 was much weaker in water $(K_{\rm M} = 10^3)$ compared to what measured in organic solvents ($K_{\rm M} = 4 \times 10^6$ in ACN).²⁰⁹ Thus, the recombination in aqueous electrolytes should result much faster than in organic electrolytes, accompanied by a substantial decrease in the collection efficiency. To better unravel this phenomenon, I_2 concentration was reduced, resulting in increased I_{sc} (lower amount of free I2 led to slower recombination) and decreased J_{dl} (I₃⁻ is the limiting species for the diffusion current in the electrolyte). Similar experiments were performed using the D149 dye observing an analogous trend. Only higher J_{dl} values were obtained, due to the chemisorption of D149 by means of a single -COOH group, which resulted in a higher CDCA surface concentration. In general, the I2 concentration was maintained higher than 10 mM, as lower values gave unstable cells. Although the topic of dyes for aqueous cells will be comprehensively detailed in Section 10, it is appropriate to report here that this study was also carried out using an N719 sensitizer, stabilised by adding HNO₃ (pH 3) in water.

Another strategy to optimize the photoanode/electrolyte interfacial characteristics is the introduction of novel additives in the electrolyte. In this respect, O'Regan and coworkers observed a positive effect on both wetting and collection efficiency by means of guanidinium and iodide ions, respectively.²⁰⁶ In particular, by using a high concentration of guanidinium iodide (GuI 8.0 M), a remarkable increase in the J_{sc} values (from 7.89 to 10.02 mA cm^{-2}) was obtained. On the other hand, $V_{\rm oc}$ remained unchanged, due to the balance of the following two opposite phenomena: the reduction of the recombination rate due to the decrease of free jodine amount and the lowering of the redox potential of the solution due to the increased iodide concentration. Efficiencies up to 4.06% were obtained (GuI 8.0 M and I_2 20 mM in H_2O ; dye: D149/CDCA), but the long-term stability of GuI-based cells was lower than that of GuSCN/NaI-based ones, more likely because - as suggested by the authors - GuI could cause the desorption of CDCA from the surface, thus leading to the dewetting of the inner pore structure. Based on these considerations, the reader can certainly appreciate the charming and delicate balance at the base of the fabrication of an efficient DSSC.

A completely different approach for the photoanode activation has been very recently proposed in the article by Spiccia and coworkers,²¹⁰ who modified the surface properties of TiO₂ nanoparticles by means of octadecyltrichlorosilane (ODTS) in order to create an insulating layer able to reduce electron recombination by restricting the access of the cobalt redox couple at the titania surface. Condensation reaction occurred directly between Ti-OH groups and ODTS, and also by alkyl group intercalation between MK-2 dye molecules (Fig. 16A). This resulted in a strengthened insulating layer, thanks to the formation of Si-O-Si and Si-O-Ti crosslinkages induced by further condensation processes activated after the introduction of the fully aqueous electrolyte $([Co(bpy)_3]^{2+} 0.20 \text{ M}, [Co(bpy)_3]^{3+}$ 40 mM, NMBI 0.70 M and PEG 300 1 wt%). As clearly visible in Table 11, the Voc of ODTS-treated photoanodes increased with soaking time, while FF decreased due to mass transport limitations in the titania mesopores.²¹¹ ODTS-treated cells showed improved J_{sc} values than ODTS-free ones, more likely because of a negative shift in the CB of the photoanode or a suppression of the interfacial electron recombination of the injected electrons with the oxidized cobalt species (Fig. 16B¹²⁸). The first hypothesis was discarded due to the similar capacitance values provided by the different photoanodes. On the other hand, ODTS showed to be useful to improve both the electron recombination resistance and the electron lifetime by increasing soaking time values, due to a higher coverage of the exposed surface sites by the alkyl siloxane (Fig. 16C). Efficiencies up to 5.64% were obtained, which did not decrease even after 500 h of storage under dark. Moreover, molecular dynamics simulations were performed (Fig. 16D): for the untreated system, the distance $d(\text{Co-TiO}_2)$ decreased significantly from 25 to 8 Å with the dynamic simulation time, while it remained almost constant for the ODTS-treated system. This confirmed that, for the ODTS-treated cells, the $[Co(bpy)_3]^{3+}$ ions did not come in contact with the titania surface, thus lowering the chance of recombination.

Until now, a semiconductor different from TiO_2 has not been employed in aqueous DSSCs. To this purpose, a good material to be explored could be zinc oxide (ZnO), whose

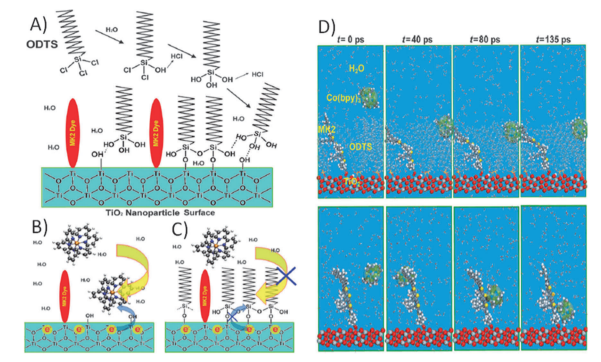


Fig. 16 (A) Proposed reaction process for alkyl siloxane anchoring to the dye-coated TiO_2 NPs treated with an ODTS solution; (B) electron recombination in non-ODTS treated aqueous DSSCs; (C) inhibition of electron recombination by aqueous DSSCs fabricated with ODTS treated photoanodes; (D) snapshot of the simulated systems of the ODTS-untreated and treated TiO_2 cluster as a function of the simulation time. In the ODTS-untreated system (bottom), the MK-2 molecule was attached at the TiO_2 cluster, whereas the ODTS-treated dyed TiO_2 photoanode (top) was obtained by attaching the crosslinked hydrolysed ODTS molecules at the vacant sites. Both systems were immersed in an explicit water environment. Adapted and reprinted from ref. 210.

Table 11PV parameters ($P_{in} = 1 \text{ sun}$) of fully aqueous DSSCs assembledwith ODTS-free or ODTS-treated photoanodes. Electrolyte: $[Co(bpy)_3]^{2+}$ 0.20 M, $[Co(bpy)_3]^{3+}$ 40 mM, NMBI 0.70 M and PEG 300 1 wt%. Dye: MK-2.Adapted and reprinted from ref. 210

$J_{\rm sc}~({ m mA~cm^{-2}})$	$V_{\rm oc}$ (V)	FF	η (%)
8.45	0.687	0.70	4.09
10.17	0.821	0.68	5.64
9.52	0.861	0.63	5.16
	8.45 10.17	8.45 0.687 10.17 0.821	8.450.6870.7010.170.8210.68

wettability and morphology can be highly tailored, as already demonstrated by research groups that usually employ this material in lithium batteries.^{212–215}

10. Dyes and stability in water: still a long road to succeed

Representing the vital component of a DSSC, the sensitizer (or dye) has been extensively investigated in the last decades.^{216–218} The main target of the scientific community efforts has been the discovery, elaboration and application of novel organometallic complexes,^{219–221} the preparation of metal-free dyes,^{222–224} the extension of the spectrum in the near IR regions^{225–227} and the invention of new synthetic methodologies being green and readily up-scalable at the industrial level.^{228–230} As regards aqueous DSSCs, the number of articles published until now that propose novel dyes *ad hoc* elaborated for the purpose has been very limited.

This is rather surprising. As reported in the previous sections of this review, several research groups more likely used common organometallic and organic dyes (maybe focusing rather on hydrophobic ones) for the fabrication of aqueous cells, the main focus being the modification of electrolyte components and the functionalization of the photoanode surface.

Bearing in mind the long-term stability as an utmost important target for a dye in aqueous DSSCs, Grätzel and coworkers initially focused their efforts on the introduction of long apolar chains able to laterally interact with Ru-based complexes; by this way, an aliphatic network was intended to form which might avoid both the TiO₂-dye debonding and the approaching of triiodide to the photoanode surface.²³¹ A long-term stability test compared the performance of N3 and [Ru(H2dcbpy)(mhdbpy)(NCS)2] dyes (where H_2 dcbpy = 4,4'-dicarboxy-2,2'-bipyridine and mhdbpy = 4-methyl-4'-hexadecyl-2,2'-bipyridine) in the presence of increasing water contents in the electrolyte (I2 10 mM in HMII/H2O; HMII = 1-hexyl-3-methylimidazolium iodide). Data depicted in Fig. 17 show that, in the presence of electrolytes encompassing 5 and 10 vol% H₂O, respectively, the performance of N3-based cells was lower with respect to that of the devices assembled with the hydrophobic dye, these latter being stable, thanks to the presence of the aliphatic chains which assured complete insolubility in water.

Thanks to their broad absorption spectra and excellent lightharvesting ability, Ru complexes would represent the sensitizer of choice for aqueous DSSCs. Nevertheless, in order to obtain cells demonstrating sufficient durability, efforts must be devoted

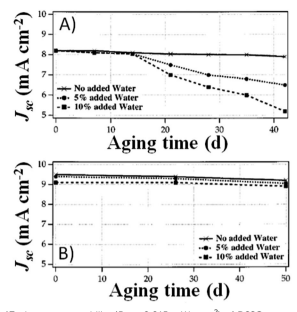


Fig. 17 Long-term stability ($P_{in} = 0.015 \text{ mW cm}^{-2}$) of DSSCs assembled with RTIL-based electrolytes (I₂ 10 mM in HMII) encompassing different H₂O contents. Dyes: (A) N3; (B) [Ru(H₂dcbpy)(mhdbpy)(NCS)₂]. Adapted and reprinted from ref. 231.

to ameliorate the bonding between dye molecules and TiO₂ NPs.²³² Miyasaka and coworkers compared two photoanodes having equal amounts of N719 and N3 sensitizer molecules adsorbed onto the surface of the TiO₂ layer.¹³⁸ In the presence of a truly aqueous electrolyte (KI 0.50 M and I₂ 25 mM), efficiencies of 0.6% and 0.5% were calculated when using N3 and N719 dyes, respectively. The cell assembled using the N719 dye showed lower J_{sc} (upon adsorption, TBA groups caused a negative Fermi level shift of the semiconductor²³³) and higher $V_{\rm oc}$ values (TBA is a base capable of negatively shifting the Fermi level as well as the TiO_2 CB) compared to the one using N3. The differences in J_{sc} were also ascribed to the adsorption strength of the dyes at the TiO₂/aqueous electrolyte interface; in fact, being more hydrophilic than N3, N719 has a higher solubility in water $(3.5 \times 10^{-3} \text{ vs. } 1.2 \times 10^{-3} \text{ M})$. The four –COOH groups of the N3 dye molecule ensure a more efficient electronic interaction at the oxide interface which, as a result, guarantees a much more stronger adsorption at the interface with the active material particles, thus improved stability under water. Moreover, the performances of all the aqueous DSSCs were found to be pH-dependent (Fig. 18A), particularly in terms of J_{sc} values which were negatively affected when a high pH (adjusted by addition of phosphate buffer solutions) was adopted; in contrast, $V_{\rm oc}$ was not affected. The $J_{\rm sc}$ decrease reflected a positive shift of the CB potential of TiO₂, as reported in Fig. 18B. However, the detachment of the dye from the electrode could also be promoted by the experimented high pH values as well.

The behaviour upon aging of Ru-based aqueous DSSCs was investigated by Miyasaka and coworkers, who observed a 50% progressive decrease of the photocurrent after 2.5 months (under dark, RT).¹³⁸ Reference cells containing organic solvent-based electrolytes showed reduced lifetime, because of solvent evaporation. After several weeks the bleaching of iodine was

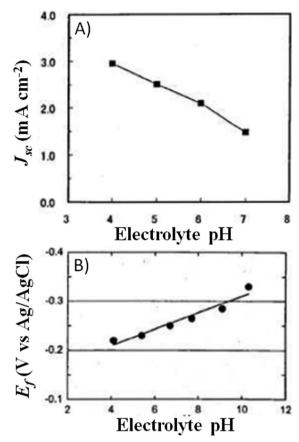


Fig. 18 (A) pH dependence of J_{sc} values ($P_{in} = 1$ sun) obtained for an aqueous DSSC sensitized with N719. Electrolyte: KI 0.50 M and I_2 25 mM; (B) dependence of the Fermi level electrochemical potential (E_f) of the mesoporous TiO₂ electrode on the pH of the above reported electrolyte. The potential data were based on the measurement of the onset potential of dark cathodic current. Adapted and reprinted from ref. 138.

detected in aqueous DSSCs. Anyway, a significant intrinsic surplus value of DSSCs is that electrolyte can be fully replaced to restore cell performance, as it was confirmed in the article by Miyasaka and coworkers where the cell efficiency was recovered by replacing the degraded aqueous electrolyte with a freshly prepared one. Of course, the procedure of electrolyte replacement is much more easy and safe when an aqueous system is envisaged than an organic volatile one that carries around severe environmental hazards.

Further surface characterization of N3-, N719- and Z907dyed electrodes was performed by Hahlin and coworkers by means of photoelectron spectroscopy (PES),²³⁴ a useful technique to get specific information on the electronic as well as the molecular surface structures of the photoanode.^{235–237} The dyesensitized samples were exposed to a mixed H₂O/EtOH solution in the 30:70 ratio for 20 min, and then assembled in organic solvent-based DSSCs (TBAI 0.60 M, LiI 0.10 M, I₂ 50 mM, GuSCN 50 mM and TBP 0.50 M in ACN; TBAI = tetrabutylammonium iodide). Water exposure led to a decrease in J_{sc} exceeding 20% for the N3- and N719-based systems, while V_{oc} decreased only for the N719-based cell; Z907-sensitized electrodes did not show any appreciable variation. Additionally, a

20 nm shift of the IPCE maximum was observed for both cells, thus clearly indicating modification in the electronic structure of the adsorbed molecules. Changes in the amount of surface adsorbed N3, N719, and Z907 dyes were finely detected by measuring the photoemission intensity from the core level Ru-3d_{5/2} relative to the Ti-2p_{3/2}: only in the case of N3 and N719 dyes, the Ru/Ti ratio decreased by about 80% and 50%, respectively. As regards the hydrophobic Z907 dye, waterinduced dye desorption was not detected, and by studying the S/Ru ratio it was concluded that no substantial permanent chemical modification, such as ligand exchange, occurred to the molecule. In agreement with the experiments of Miyasaka and coworkers,¹³⁸ it was observed that the N719 dye layer was more sensitive to water compared to the N3 one, and its S-2p and N-1s signals were mostly affected by the aqueous environment. As a result, clear shifts of the HOMO levels towards higher binding energies occurred for both the dyes (i.e., 0.17 and 0.26 eV for N3 and N719, respectively), thus leading to an increased gap between the energy levels of the HOMO and the LUMO, and subsequently a blue shift in the absorption spectrum.

As previously mentioned, very little studies have been published on the preparation of dyes having specific characteristics to be effectively used in aqueous cells. A seminal work of Ko and coworkers reported about new organic sensitizers (JK-259 and JK-262) containing 4-(2-(2-methoxyethoxy)-ethoxy)phenylamine as the electron donor molecule and cyanoacrylic acid as the electron acceptor molecule bridged by the 9,9-bis(2-(2-methoxyethoxy)-ethyl)-9H-fluorenyl linker (Table 14).238 The 2-(2-methoxyethoxy)ethyl unit was introduced to ameliorate the water compatibility, and a TBP treatment onto the TiO₂ films ensured the photoanode stability. JK-259 and JK-262 showed efficiency values of 1.16% and 2.10%, respectively, in a fully aqueous electrolyte (PMII 2.0 M, I2 50 mM, GuSCN 0.10 M, TBP 0.50 M and Triton X-100 1 wt%). However, due to the very low J_{sc} values, these efficiencies were >50% lower when compared to those measured in the corresponding organic solvent-based DSSCs. The authors justified the decrease in the photocurrent on the basis of detachment of the adsorbed dye through the hydrolysis of the TiO₂-water surface linkage, which is rather contradictory to the aim of their work which was intended for the preparation of a dye stable in water. Apart from this curious aspect, we may argue that research in this field should be much more effective because plenty of possibilities are available for performance and stability improvements.

Ko and coworkers further investigated aqueous DSSCs using three different organic sensitizers, namely D5L6,²³⁹ D21L6²⁴⁰ and JK-310; in particular, the latter two contained (hexyloxy)phenylamine and 4-(2-(2-methoxyethoxy)-ethoxy)phenylamine as electron donor units.²⁴¹ As clearly visible in Table 14, all of these dyes have a water-soluble substituent and were coupled with an aqueous electrolyte (PMII 2.0 M, I₂ 50 mM, GuSCN 0.10 M, TBP 0.50 M and Triton X-100 1 wt% in MPN:H₂O 50:50). PV measurements (Table 12) showed that V_{oc} gradually increased with the incremental addition of water (*i.e.*, from 0 to 50%); in contrast, J_{sc} sharply decreased under the same conditions. Again, dyes were found to easily detach due to their partial

Table 12PV parameters ($P_{in} = 1 \text{ sun}$) of DSSCs assembled with differentorganic dyes and liquid electrolytes. Electrolytes: PMII 2.0 M, I2 50 mM,GuSCN 0.10 M, TBP 0.50 M and Triton X-100 1 wt% in MPN or MPN/H2O.Adapted and reprinted from ref. 241

Dye	H ₂ O (vol%)	$J_{\rm sc} ({ m mA} { m cm}^{-2})$	$V_{\rm oc}$ (V)	FF	η (%)
D5L6	0	10.50	0.66	0.75	5.20
	50	6.22	0.75	0.73	3.40
D21L6	0	12.57	0.71	0.74	6.56
	50	7.46	0.77	0.77	4.41
JK-310	0	12.28	0.70	0.72	6.18
-	50	6.62	0.75	0.76	3.77

hydrophilic nature, resulting in the reduction of both photocurrent and efficiency. Such a result further confirmed that the step of tailor-making the dyes in order to get improved photoanode wettability must be carefully counterbalanced by the water solubility value of such organic molecules.

As an alternative to the synthesis of new dyes, an impressive work of modification of known sensitizers to make them efficient and stable in an aqueous environment started in 2014. Bisquert and coworkers demonstrated an efficient synthetic protocol to modify the existing cyanoacrylate moiety of organic dyes (whose carboxylate linkage is susceptible to hydrolysis²⁴²) with a highly water-stable hydroxamate anchoring group via a condensation reaction.²⁴³ The well known MK-2 dye^{184,244,245} was chosen as a target substrate, and its hydroxamate-derivative (MK-2HA, Table 14) was significantly (30-50%) less susceptible to desorption in water, as a result of the stronger interaction of the novel anchoring group with the surface of TiO₂.^{246,247} The enhanced water tolerance was also attributed to the higher pK_a of MK-2HA dye and to the less strained chelate bite angle in hydroxamic acids compared with carboxylic acid analogues. Interestingly, TAS measurements carried out on both dyes revealed that the lifetime of the charge-separated state increased with increasing immersion time in the DMF-H2O solution used to evaluate the water stability of sensitizers. Thus, such a "soaking and desorption" approach, which probably contributed to the removal of dye aggregates from the semiconductor surface,¹⁷¹ was proposed as a potentially effective treatment to improve the recombination dynamics of DSSC photoanodes. In these experiments, MK-2HA-sensitized photoanodes showed no change in the electron injection efficiency upon exposure to the desorption solution, while MK-2 ones lost nearly 65%, due to the sensitivity of the dye-TiO₂ interface toward water. Cells with MK-2 and MK-2HA dyes were fabricated and water was purposefully added to the electrolyte (MPII 2.0 M, LiI 0.10 M, I₂ 0.20 M and TBP 0.50 M in ACN/H₂O). In 200 h aging test under dark, MK-2-based DSSCs with 10 and 20 wt% water had decreased efficiency by 15% and 50% compared to the anhydrous control, mainly due to the 50% reduced J_{sc} . As regards MK-2HA-based cells, there was no significant difference in the efficiencies (slightly lower than 4%) measured with aqueous (10 wt%) and anhydrous electrolytes. Moreover, FF and $V_{\rm oc}$ of water-based DSSCs increased with time: authors hypothesized that the water content in the electrolyte could serve to accelerate the desorption of aggregates, leading to more favourable interfacial electron transfer dynamics.



Fig. 19 Crossed sections of working electrodes sensitized with D35 (left) and V35 (right) with a drop of deionized water positioned on the top. CA values are also reported. Adapted and reprinted from ref. 249.

D35 is another well known dye in the DSSC field,²⁴⁸ and Boschloo and coworkers synthesized its hydrophilic analogous, by substituting the hydrophobic alkyl chains with glycolic ones.²⁴⁹ The resulting hydrophilic dye (V35, Table 14) showed an extraordinary interaction with water, and the authors claimed that the use of wettability-improver surfactants was not necessary (as can be seen in Fig. 19), thus reducing cell cost and avoiding possible mass transport problems. The new dye was initially coupled with a fully aqueous electrolyte containing only NaI and I2, and then gradually modified as reported in Table 13. Interestingly, efficiency increased when KI replaced NaI, probably due to the ability of glycolic chains (like crown ethers) to coordinate small cations (such as Na⁺).^{250,251} Performance was further improved by doubling the concentration of KI, thus lowering the amount of free iodine. The addition of GuSCN had a negative effect, while CDCA improved efficiency up to 2.20%. Moreover, V35 dye was further investigated in aqueous DSSCs employing alternative cathodic materials at different pH values, as will be described in Section 11.

Pigments extracted from natural matrices may result in a valuable alternative to synthetic dyes.^{252,253} Despite the efficiencies obtained are still lower compared to the reference Ru-based dyes, thanks to their low cost and ready availability, these pigments are nowadays very much considered by those research groups who are focused on the development of the so-called biophotovoltaic devices.181,254,255 In Section 8, the very recent work of Su and Lai on natural pigments coupled with alternative redox mediators has been mentioned,¹⁷⁸ but the very first demonstration of the good compatibility between pigments and water was provided by Rabani and coworkers.²⁵⁶ In 2001, they exploited the promising prospects as a sensitizer of the seed coats of the pomegranate (PG) fruit, which is a rich source of anthocyanins.²⁵⁷ By operating under strong acidic conditions (pH 1) and after having ascertained that Na⁺ was the most effective counter ion in photocurrent generation, the resulting aqueous DSSC (NaI 1.0 M and I2 0.10 M) demonstrated a Jsc value of

Table 13PV parameters ($P_{in} = 1 \text{ sun}$) of V35-sensitized DSSCs assembledwith different fully aqueous electrolytes. Adapted and reprinted fromref. 249

Electrolyte	$J_{\rm sc}({\rm mA}{\rm cm}^{-2})$	$V_{\rm oc}\left({\rm V}\right)$	FF	η (%)
NaI 2.0 M and I ₂ 20 mM	2.30	0.500	0.67	0.80
KI 2.0 M and I ₂ 20 mM	4.07	0.550	0.70	1.55
KI 4.0 M and I ₂ 20 mM	4.78	0.570	0.65	1.76
KI 4.0 M, I2 20 mM and GuSCN 0.50 M	3.33	0.555	0.67	1.25
KI 4.0 M, I ₂ 20 mM and CDCA sat.	4.86	0.600	0.76	2.20

2.2 mA cm⁻² and a V_{oc} of 0.44 V (FF was not reported). Stability tests showed no appreciable decrease of the photocurrent after 24 h illumination. It is important to recall here that the size of the photoanode NPs is a key factor affecting the performances of natural pigment-based cells. In their investigation, Rabani and coworkers used 5 nm diameter NPs which guaranteed small pores, thus preventing or minimizing the adsorption of undesired molecules (*i.e.*, foreign impurity phases deriving from the original matrix) which could result in reduced photocurrent by means of visible light absorption competition.

An overview of the dyes used so far for aqueous DSSCs is given in Table 14, together with the redox mediators with which they have been tested. Very recently, novel metal-free dyes containing an anthracene/phenothiazine unit in the spacer have been synthesized and proposed for aqueous DSSCs, and a cell performance equal to 4.96% was achieved.²⁵⁸

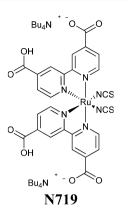
11. Low-cost and water-compatible cathodes

Being the most popular choice for high-efficiency DSSCs, platinum has been widely recognized as the benchmark material at the cathode side, due to its high catalytic activity and excellent conductivity.²⁵⁹ As a result, for many years, research on novel cathodic materials has been considered rather a secondary issue by the scientific community, especially if compared to the impressive efforts addressed to the improvement of the other cell components. Nevertheless, Pt presents a series of drawbacks, the most significant being the high cost and the scarce availability, which have restricted so far the mass production of DSSCs,²⁶⁰ as already occurred to fuel cells and related systems using Pt as a catalyst.²⁶¹ Moreover, Pt may readily undergo severe dissolution in electrolytes containing the iodine-based redox couple, resulting in the formation of PtI4 and reduced cell performances;²⁶² Pt is not compatible with the recently proposed sulphur-based redox shuttles as well.¹⁸⁹ For all these reasons, we may reasonably argue that the breakthrough in the widespread diffusion of DSSCs may be realised only when counter electrodes at sustainable costs and stable long-term performance are developed.²⁶³ In this respect, transition metal compounds (*i.e.*, carbides, nitrides) have drawn considerable attention as alternatives to Pt cathodes, due to their excellent catalytic activity towards electrolytes based on both sulphur- and cobalt-based redox couples.^{264,265} Organic polymers (i.e., PEDOT, PPy, PANI, etc.) demonstrated promising prospects as flexible and transparent cathodes.^{266,267} In the last two years, low-dimensional carbonaceous materials (i.e., CNTs arrays and graphene films) started to be widely studied and applied, especially in iodine-free DSSCs.^{268,269}

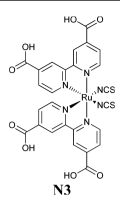
As previously reported in the course of this review (Section 8), redox couples to be used in aqueous DSSCs are often different from the standard ones conceived for aprotic media. The use of water as a solvent (sometimes mixed with ACN or MPN) makes the electrolyte/cathode compatibility a new topic to be deeply investigated. Obviously, aqueous electrolytes being a recent

COOH

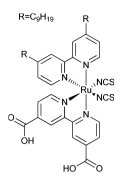
Table 14 Structural formulas of sensitizers usually employed in aqueous DSSCs, together with the redox pairs with which were coupled. Further details about the other cell components and the PV performance are reported in the manuscript and in Table 20



Always with I^{-}/I_{3}^{-} with increasing $H_{2}O$ content: 0.018 wt%,¹⁷² 4 wt%,¹⁵⁶ 10 wt%,¹⁴⁵ 65 wt%,¹³⁸ and in 100 wt% $H_{2}O^{138,163,206}$

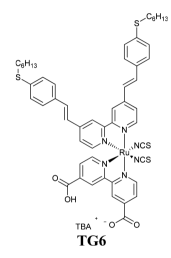


Always with I^{-}/I_{3}^{-} with increasing $H_{2}O$ content: 3 wt%,¹²⁴ 4 wt%,⁴³ 5 wt%,³⁰³ 10 wt%,²⁹⁴ 65 wt%,^{138,203} and in 100 wt% $H_{2}O^{138,328}$

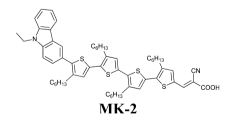


Z907

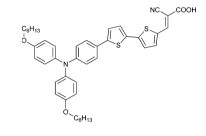
Always with I^-/I_3^- with increasing H_2O content: 10 wt%, 128 50 wt%, 310 and in 100 wt% H_2O^{206}

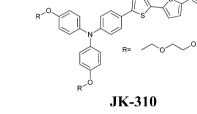


Always with I^-/I_3^- with increasing H_2O content: 20 wt%, $^{120}_{120}$ 50 wt%, $^{318}_{318}$ and in 100 wt% $H_2O^{120,206}$

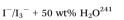


 $\begin{array}{l} \label{eq:constraint} Fe(CN)_6{}^{4-}/Fe(CN)_6{}^{3-} \mbox{ in 100 wt\% } H_2O^{182} \\ [Co(bpy)_3]^{2+}/[Co(bpy)_3]^{3+} \mbox{ in 100 wt\% } H_2O^{194,210} \end{array}$





 I^-/I_3^- + 50 wt% H_2O^{241}

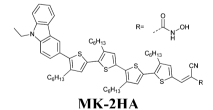


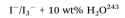
D21L6

D5L6

 $I^-\!/{I_3}^- + 50 \ wt\% \ H_2O^{241}$

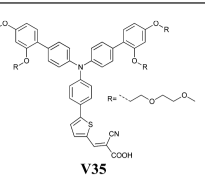
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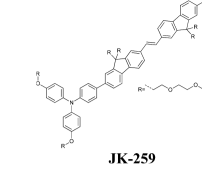


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Table 14 (continued)



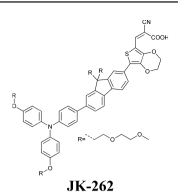
 $I^-\!/{I_3}^-$ in 100 wt% H_2O^{249}



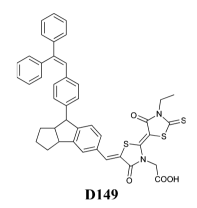
Соон

 $I^-/{I_3}^-$ in 100 wt% H_2O^{238}

C₆H₁₃ C₆H₁₃



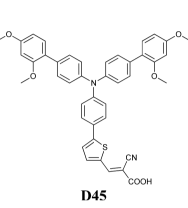
 $I^-/{I_3}^-$ in 100 wt% H_2O^{238}



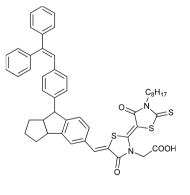
 $I^-/{I_3}^-$ in 100 wt% H_2O^{206}



 Fe^{2+}/Fe^{3+} + 65 wt% H_2O^{204}

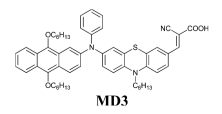


 $TT^-EMI^+\!/DTT$ in 100 wt% H_2O^{193}

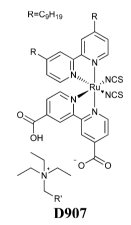


D205

TEMPOL/TEMPOL⁺ in 100 wt% H_2O^{274}



 $TEMPO/I^-\!/{I_3}^-$ in 100 wt% H_2O^{258}



D51

 $TT^-EMI^+\!/DTT$ in 100 wt% H_2O^{193}

 $I^{-}\!/{I_{3}}^{-} + 65 \ wt\% \ H_{2}O^{314}$

Natural dyes/pigments CV-G with Ce^{3+}/Ce^{4+} + 30 wt% H_2O^{178} RhS with Ce^{3+}/Ce^{4+} + 65 wt% H_2O^{174} PG with I^-/I_3^- in 100 wt% H_2O^{256}

research interest, very few literature reports useful to identify an optimum cathodic material have been available so far. Notwithstanding, some interesting studies have been already proposed, which will be hereby briefly summarised.

When using Co complexes as aqueous redox shuttles, Spiccia and coworkers observed a high charge transport resistance between the water-based electrolyte and the Pt counter electrode.¹⁹⁴ Thus, they screen-printed a mesoporous ITO film onto the FTO substrate, and then added a thermally-deposited Pt layer. The resulting cathode (13.5 µm-thick) led to an efficiency of 5.0% (4.2% with Pt only), which increased by another 0.1% after 48 h. The ITO/Pt cathode showed a R_{CE} equal to 1.6 Ω , around five times lower than that of pure Pt, along with an almost doubled exchange current (investigated by means of cyclic voltammetry), thus indicating improved electron transfer to the oxidized redox couple. The reason for this behaviour lied in the high surface area of the ITO mesoporous film, which enabled a higher Pt loading and facilitated the reduction of the oxidized species of the aqueous electrolyte at the counter electrode surface.

Poly(3,4-ethylenedioxythiophene) (PEDOT) is currently one of the most considered materials for the preparation of both rigid and flexible cathodes to be used in DSSCs based on alternative redox couples. Sun and coworkers tried to improve the FF of DSSCs assembled with a thiolate/disulphide redox couple by using PEDOT as a counter electrode.¹⁹³ However, they failed because of the poor stability of PEDOT in the aqueous electrolyte, which also resulted in its easy detachment from the FTO glass. Quite surprisingly, the opposite behavior was reported by Boschloo and coworkers with fully aqueous cells containing V35 dye and an iodide/triiodide redox couple.249 Indeed, Table 15 shows that PEDOT counter electrodes outperformed Pt ones due to increased J_{sc} and V_{oc} values. Such a positive effect was attributed to the leaf-like PEDOT structure, which conferred a high-surface area to the cathode, thus increasing its catalytic activity. Moreover, a reduced tendency of the dye to desorb in the presence of PEDOT was observed: authors speculated that PEDOT partially trapped in its matrix the ions capable of being absorbed onto the surface of TiO_2 ,²⁷⁰ preventing in this way the desorption of the dye. V35 was also tested with the best electrolyte reported by O'Regan and coworkers (see Section 9,²⁰⁶); however, the elevated concentration of GuI decreased significantly $V_{\rm oc}$ values (Table 15) due to the absorption of guanidinium cations on the TiO₂ surface.²⁷¹ Further optimization

Table 15PV parameters ($P_{in} = 1 \text{ sun}$) of V35-sensitized DSSCs assembledwith different electrodes and electrolytes. Adapted and reprinted fromref. 249

Electrolyte	Cathode	$J_{ m sc}$ (mA cm ⁻²)	$V_{\rm oc}$ (V)	FF	η (%)
KI 4.0 M, I ₂ 20 mM and CDCA sat.	Pt	4.86	0.600	0.76	2.20
KI 4.0 M, I ₂ 20 mM and CDCA sat.	PEDOT	5.26	0.625	0.74	2.45
KI 2.0 M, I ₂ 10 mM and CDCA sat.	PEDOT	6.85	0.650	0.67	3.01
GuI 8.0 M, I_2 20 mM and CDCA sat.	PEDOT	5.76	0.550	0.62	1.97



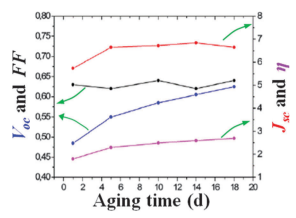


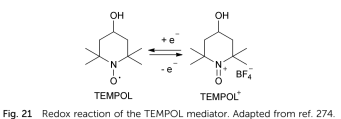
Fig. 20 Long-term stability under dark of DSSCs assembled with a fully aqueous electrolyte (KI 4.0 M and I₂ 20 mM at pH 8.0) and a PEDOT counter electrolyte. Dye: V35. PV parameters were measured under $P_{in} = 1$ sun. Adapted and reprinted from ref. 249.

of iodine and CDCA concentrations yielded the highest efficiency recorded with V35 dye, namely 3.01% (4% under 0.5 sun). Noteworthy, all the electrolytes proposed by Boschloo and coworkers contained CDCA as a $V_{\rm oc}$ -improver agent: it did not shift significantly the TiO₂ CB,²⁷² but reduced the recombination processes.

To further demonstrate the long-term stability of the PEDOT-based aqueous cells, Boschloo and coworkers set up several aging tests, also studying the effect of the pH of the electrolyte on the PV parameters.²⁴⁹ As shown in Fig. 20, both V_{oc} and FF were higher in the case of electrolytes (KI 4.0 M and I₂ 20 mM) at pH 9.0; however, a lower photocurrent and an overall efficiency were obtained with respect to cells at pH 8.0. One very important aspect was given by fluctuations over time of the PV parameters (Fig. 20), and the better efficiencies were collected several days later device assembly. This is a typical characteristic of aqueous cells, thus scientific groups are suggested to monitor their devices over time rather than immediately abandoning their new materials due to low performance measured immediately after DSSC assembly.

Also materials other than PEDOT have surprisingly demonstrated increased performance and stability as cathodes in aqueous electrolyte-based DSSCs. In their pioneering work, Kim and coworkers revealed for the first time that the corrosion potential of silver metal increased in aqueous electrolytes containing Triton X-100.¹⁵⁶ The surfactant may in fact create a sort of protective layer on the Ag film surface which reduced the direct interaction with I_3^- ions, which in turn greatly diminished the dissolution of the metal. In the presence of the surfactant, an increased overpotential for the electron transfer from the Ag layer to I_3^- was additionally recorded. Such encouraging results unveiled the possibility of using Ag instead of the more expensive Pt in futuristic commercial DSSCs.

Very recently, the coating of the cathode surface with a polymeric film has been proposed to improve the electrode/ electrolyte interfacial properties.²⁷³ In this respect, Nishide and coworkers used 4-hydroxy-2,2,6,6-tetramethylpiperidinoxyl (TEMPOL, Fig. 21) as a redox mediator in an aqueous DSSC.²⁷⁴



TEMPOL is a hydrophilic TEMPO derivative, belonging to the class of nitroxide radical molecules, which have attracted remarkable attention as organic-based redox-active materials due to their rapid and reversible one-electron-transfer capability to form the corresponding oxoammonium cations.275-277 This alternative redox mediator (to be added to the list shown in Section 8) was immobilized in a Nafion[®]:Pt two-layers cathode. Nafion[®] is typically used to prepare polymer electrolyte membranes,²⁷⁸ but also as a trapping agent for electroactive cations in sensing and photoelectrochemical devices.^{279,280} Exploiting its properties, a significant increase of both J_{sc} and FF values was evidenced with respect to the Pt-based cell, which was ascribed to the lower cell resistance due to the high efficiency in the regeneration reaction of the redox mediator as well as to the steep concentration gradient of the TEMPOL cation promoted by the Nafion[®] layer. In the presence of the Nafion[®]:Pt two-layers counter electrode and using the D205 dye, the overall cell efficiency increased from 0.11% ($J_{sc} = 1.1 \text{ mA cm}^{-2}$, $V_{oc} = 0.76 \text{ V}$, FF = 0.14) to 2.1% ($J_{sc} =$ 4.5 mA cm⁻², $V_{oc} = 0.69$ V, FF = 0.64).

12. Cell sealing in the presence of water: a new intriguing challenge

Being the long-term stability a key objective of the present scientific research activity in the DSSC community, amounts of materials have been explored as sealants, such as thermoplastic hot-melt foils (*i.e.*, Surlyn[®],²⁸¹ Bynel[®],²⁸² *etc.*), UV curable glues³⁷ and glass frit (GF).^{283,284} GF is believed to be one of the strongest candidate as a sealant material in DSSC glass modules, as it possesses the same characteristics of the substrate, thus optimum compatibility; moreover, it is non-permeable, stable under UV light and at elevated temperatures.^{285–287} On the other hand, it has been quite difficult to find lead-free GF,²⁸⁸ demonstrating at the same time low melting temperature and high chemical stability towards the redox mediator. Moreover, GF must not leach elements into the electrolyte, which could cause the depletion of the I₃⁻ ions.^{115,289}

The use of an aqueous electrolyte opens up an amount of novel opportunities in the development of specific sealants for water-based DSSCs. A seminal work was proposed by Hinsch and coworkers, who fabricated Bi_2O_3 - SiO_2 - B_2O_3 and ZnO- SiO_2 - Al_2O_3 GFs, with characteristic melting temperatures in the range of 400–520 °C.²⁹⁰ When Bi_2O_3 - SiO_2 - B_2O_3 was coupled with an aqueous electrolyte (I₂ 0.10 M, GuSCN 0.10 M and NBBI 0.50 M in MPII/ACN/H₂O), UV-Vis measurements revealed a significant decrease in the I₃⁻ absorbance with time (Fig. 22A),

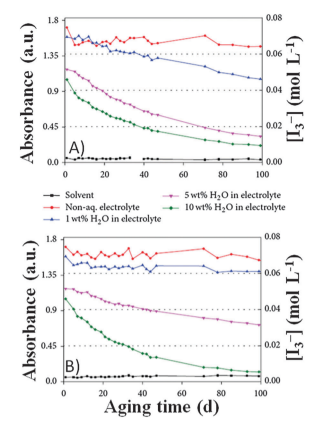


Fig. 22 Absorbance at 430 nm as a function of the aging time for DSSCs filled with solvent, non-aqueous electrolyte and redox electrolyte encompassing 1, 5 and 10 wt% H₂O, in the presence of different GFs as sealants: (A) $Bi_2O_3-SiO_2-B_2O_3$; (B) ZnO-SiO_2-Al_2O_3. Adapted and reprinted from ref. 290.

while energy dispersive X-ray (EDX) analysis revealed a precipitate of bismuth and iodine. Indeed, BiI_4^- formation was observed, and its absorption band at 460 nm competed with the one characteristic for the sensitizer.²⁹¹ As regards the ZnO–SiO₂–Al₂O₃ GF, the electrolyte bleached because of I_3^- depletion (Fig. 22B); EDX revealed the leakage of zinc from the GF. In turn, the metal formed a thermally stable compound with iodine,²⁹² being detrimental on the DSSC performance. Moreover, for both the GFs, the I_3^- depletion was enhanced while increasing the content of water in the electrolyte.

The GFs proposed by Hinsch and coworkers represent only the first example of the sealing material for aqueous DSSCs, thus other metal oxide formulations must be elaborated to assure the development of truly stable cells. However, in the case of aqueous DSSCs, also the polymeric sealants would return in vogue, since the water permeability would no longer be a problem for cell stability.

13. Quasi-solid electrolytes: the step forward

In Section 2, the main drawbacks of liquid electrolytes have been thoroughly discussed, volatility and difficulty of hermetic sealing being the most significant. We have already shown that volatility may be remarkably reduced by introducing water as solvent of the redox mediator in place of ACN; nevertheless, the use of specifically developed quasi-solid electrolytes would clearly facilitate the operations of cell assembly and sealing, also guaranteeing a prolonged stability to the device. As a result, many research groups working on aqueous electrolytes have directed their efforts towards the realisation of water-based DSSCs in the quasi-solid state.

13.1 RTIL-water mixtures

The quasi-solidification of a liquid electrolyte is usually carried out through the introduction of RTILs or polymeric materials.^{18,48,68} The pioneering work on the development of RTIL-laden polymer electrolytes was published by Mikoshiba and coworkers, who used water to face the slow I_3^- diffusion in the highly viscous RTILs.²⁹³ By coupling MPII with different components (*e.g.*, LiI/TBP/I₂, TBP/I₂, LiI/I₂, I₂), it was observed that the addition of water improved the efficiency of N3-based cells, the best results being obtained using 5 wt% H₂O in the presence of LiI/I₂, while 10 wt% in the case of the LiI/TBP/I₂ system. The improvement was attributed to the reduced electrolyte viscosity and to the increased V_{oc} : a remarkable efficiency of 4.2% was obtained.

A more detailed analysis regarding the effects of water on the RTIL-based electrolytes was conducted by Hayase and coworkers.²⁹⁴ As shown in Table 16, several additives were introduced to increase the conductivity of MPII: among all, water was demonstrated to be the best one, besides being the one having the lowest environmental impact. Table 17 shows that increasing the water content led to reduced viscosity (μ) and increased ionic conductivity (σ , also reflected in the improved diffusion coefficient for I₃⁻). Other positive aspects were the R_{CE} decrease and *E* increase while increasing the

Table 16 μ and σ values of MPII-based electrolytes added with differentadditives in the 10 wt% ratio. Adapted and reprinted from ref. 294

Additive	μ (mPa s)	$\sigma ~({ m mS~cm^{-1}})$
Formamide	108	1.8
NMP	238	1.2
GBL	153	1.9
EC	104	1.5
PC	220	1.1
Diethoxyethane	139	2.1
EtOH	57	3.6
H ₂ O	18	15
Ethylene glycol	104	2.6

amount of water. As regards the TiO₂ surface, when 8% of water was added in the electrolyte, it was observed that the flat band potential shifted positively from -1.11 to -1.01 eV, due to the adsorption of H₂O molecules onto the semiconductor.^{82,83} This modified the characteristics of the electron diffusion in the nanostructured photoanode: electron lifetime (τ), electron diffusion constant (*D*) and electron diffusion length (*L*) decreased upon the introduction of water. This was due to the replacement of the Li⁺ ions adsorbed onto the TiO₂ surface by water molecules, resulting in a less effective ambipolar diffusion of the electrons which, in turn, led to an increased electron recombination probability.

Imidazolium-based RTILs are very popular due to their peculiar characteristics of being solvents and salts at the same time. However, the synergistic effect of using mixtures of different ionic liquids has been also proposed to possibly improve the PV performance. Among all, bistriflimide (TFSI⁻) gave excellent performance when combined with imidazolium-based RTILs.²⁹⁵ Even if TFSI⁻-based RTILs are generally considered as hydrophobic,^{296,297} coulometric Karl Fischer titration enlighten the presence of significant amounts of water (*i.e.*, around 7 mol%) most likely absorbed from the ambient air.²⁹⁸ To better unravel this phenomenon, Kim and coworkers theoretically investigated the effects of water on the microstructure as well as the transport properties of a mixture of 1-ethyl-3-methylimidazolium (EMIm⁺), TFSI⁻, I⁻ and I₃⁻.²⁹⁹ By considering a 0–25% mole fraction (χ_w) of water, three regimes were obtained (Fig. 23A):

– ice-like regime ($\chi_w < 7.69\%$, Fig. 23B), with water molecules mostly isolated and intercalated between two EMIm⁺ and two TFSI⁻ units, resulting in an ice-like entropy.

– Nano-cluster regime ($\chi_w = 7.69-14.29\%$, Fig. 23C), with nanosized water clusters bridged between two EMIm⁺ and two TFSI⁻ units.

– Liquid-like regime (χ_w = 14.29–25.00%, Fig. 23D), that is clusters composed of several water molecules located between EMIm⁺ and TFSI[–] ions.

In order to examine the effect of water absorption on the transport properties of the electrolyte, the self-diffusion coefficients were measured based on the Green–Kubo relationship,³⁰⁰ and it was found that the addition of water up to 10 mol% enhanced the diffusivity of both EMIm⁺ and TFSI⁻, with marginal variations in the diffusivity of I⁻ and I₃⁻. More in detail, the increase of entropy and mobility of EMIm⁺ caused local fluctuations in I⁻ and I₃⁻, thus enhancing the rate of Grotthuss-like electron transfer³⁰¹ as well as the rate of oxidized

Table 17 μ , σ , apparent diffusion constant for I_3^- [$D(I_3^-)$], R_{CE} , E, τ , D and L values for MPII-based electrolytes with different water contents. Electrolytes tested were $I_2 0.50$ M in MPII: H_2O for μ , σ and $D(I_3^-)$; $I_2 0.30$ M in MPII: H_2O for R_{CE} ; MPII 60 mM and $I_2 10$ mM in ACN/ H_2O for E; Lil 0.50 M, $I_2 0.30$ M, TBP 0.58 M in MPII/ H_2O for τ , D and L. $P_{in} = 1$ sun. Dye: N719. Note: we think that a factor of $10^{-5/-7}$ is missing in the reported $D(I_3^-)$ values. Adapted and reprinted from ref. 294

H_2O (wt%)	μ (mPa s)	$\sigma \ ({ m mS \ cm^{-1}})$	$D({\rm I_3}^-)({\rm cm}^2~{\rm s}^{-1})$	$R_{\rm CE} \left(\Omega \ {\rm cm}^{-2} \right)$	$E(\mathbf{mV})$	τ (s)	$D (\mathrm{cm}^2 \mathrm{s}^{-1})$	L (cm)
0	564	0.7	6.5	3.04	_	0.32	1.36	0.66
1	285	1.35	9.5	1.99	0.09		—	
5	52.8	5.81	23	0.24	0.17	0.12	1.23	0.38
10	19.2	14.12	34	0	0.24	0.12	1.23	0.38

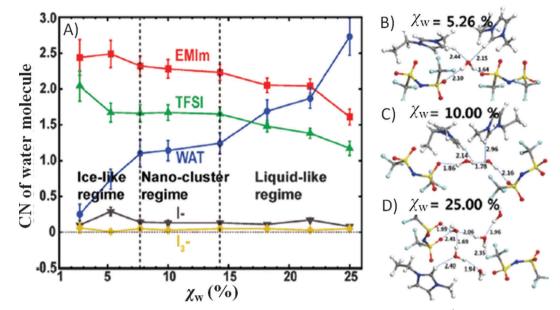


Fig. 23 (A) Dependence of water coordination (CN) on χ_w in the presence of the following mole fractions: EMIm⁺ 0.50, TFSI⁻ 0.36, I⁻ 0.13 and I₃⁻ 0.01. Representative water structure mediating EMIm⁺-TFSI⁻ interactions after 30 ns molecular dynamics for: (B) ice regime, (C) nanocluster regime, and (D) liquid-like regime. Adapted and reprinted from ref. 299.

dye reduction. On the other hand, when using high amounts of water, the large water clusters allowed water molecules to access the TiO_2 nanostructure, thus weakening the Lewis acid–base interaction of the carboxylic group of dye molecules with the anode particles. Based on all of these simulations, it was clear that the H₂O–RTIL system provided the best performance in the nanocluster regime, because of allowing spontaneous water absorption into the RTIL, retention in the RTIL phase by binding as nanoclusters to EMIm⁺ or TFSI⁻ ions and lubricating fluctuations in the RTIL to enhance Grotthuss electron transport.

13.2 Gel-polymer aqueous electrolytes

The second category of quasi-solid electrolytes is based on a polymeric matrix, capable of gelifying the redox mediator solution and trapping it within the macromolecular network.³⁰² In 2004, Hayase and coworkers firstly proposed an aqueous gelpolymer electrolyte obtained by reacting poly(vinylpyridine) (PVP, $M_{\rm p}$ = 80 000) with 1,2,4,5-tetra(bromomethyl)benzene to make chemically crosslinked ionomer structures.³⁰³ The resulting polymer³⁰⁴ trapped 0.30 M I₂ in MPII-H₂O mixtures, where water was useful to decrease the gel viscosity, thus guaranteeing a good physical contact between the gel and the mesopores of the photoanode. Moreover, $D(I_3^{-})$ was improved upon water addition, because of the phase separation between PVP and MPII, which in turn retarded the interaction between the cations fixed on the polymer matrix and I_3^- ions.³⁰⁵ Another advantage of water addition was the decrease in $R_{\rm CE}$. As a further interesting aspect of their work, Hayase and coworkers carried out the treatment of the N719sensitized photoanode with acetic acid, which bonded on the free TiO_2 sites. As a consequence, the electron recombination with I_3^- , which is the most relevant drawback of RTIL-based electrolytes,

where high I₂ amounts are added to increase the limiting currents, was inhibited. An efficiency value of 2.4% (J_{sc} = 10.9 mA cm⁻², V_{oc} = 0.52 V, FF = 0.42) was obtained in the presence of 5 wt% H₂O.

In the field of gel-polymer electrolytes, PEG is one of the most successful material due to its ability to play the role as a host for several metal-salt systems, being at the same time a good binder for other phases.^{306–308} Moreover, to achieve high ionic conductivity values, polymer blending with PEG is a valuable strategy.³⁰⁹ With the aim of confirming the practical application of such an approach in an aqueous environment, Nateghi and coworkers prepared a gel-polymer electrolyte consisting of poly(vinylpyrrolidone) (PVPi, $M_n = 40\,000, 2 \text{ wt\%}$), PEG 400 4 wt%, KI 0.20 M and I₂ 40 mM in ACN/H₂O.³¹⁰ The effect of water concentration on the polyiodide formation was investigated by analyzing the intensity ratio between the polyiodide (I_n^-) and the I_3^- species: a decrease of the I_n^-/I_3^- ratio occurred upon water addition. Theoretical calculations indicated that I_n^- species were more effective electron acceptors than I_3^- due to more delocalized charges. This facilitated the reaction with electrons due to reduced repulsion;³¹¹ moreover, the decrease of the I_n^{-}/I_3^{-} ratio contributed to reduce the charge recombination losses. As a consequence, $V_{\rm oc}$ values of the aqueous quasi-solid DSSCs increased because of water addition; this effect was also due to water adsorption onto the TiO₂ surface. On the other hand, the presence of water negatively affected the Jsc values, due to the lower concentration of I_3^- in the aqueous electrolyte and the detachment of the adsorbed Z907 dye molecules. Overall, cells with 50 wt% H₂O showed an efficiency equal to 2.31% (J_{sc} = 7.37 mA cm⁻², V_{oc} = 0.63 V, FF = 0.50), while in the absence of water the average value was 2.88% (J_{sc} = 13.7 mA cm⁻², V_{oc} = 0.58 V, FF = 0.35). When tested for their stability (under dark, 65 °C, 85% RH),

aqueous cells were able to retain the 63% of their initial efficiency after 168 h of operation, remarkably better than the 12% retention for the ACN-based quasi-solid DSSCs, thus further confirming the positive aspects of the lower volatility of aqueous electrolytes.

Amphiphilic block copolymers (ABCs) represent an emerging class of macromolecular matrices, because of their ability to selfassemble into liquid crystalline phases such as cubic, hexagonal and/or lamellar; in these configurations, polymer gels exhibit enhanced σ values with respect to other crystalline phases.^{312,313} Being the behaviour of ABCs more relevant in water-based solvents, Soni and coworkers prepared gel-polymer electrolytes with Pluronic F77 [HO(CH₂CH₂O)₅₂(CH₂CH(CH₃)O)₃₅(CH₂CH₂O)₅₂H] and Pluronic P-123 [HO(CH₂CH₂O)₂₀(CH₂CH(CH₃)O)₇₀(CH₂CH₂O)₂₀H].³¹⁴ These are triblock copolymers (widely known as poloxamers) composed of a central hydrophobic chain of poly(propylene oxide) (PPO) flanked by two hydrophilic chains of poly(ethylene oxide) (PEO). F77 ($M_n = 6600$), P-123 ($M_n = 5750$) or pure PEG ($M_n = 6000$) were introduced in the 35 wt% ratio in aqueous electrolytes containing BMII 2.0 M, I₂ 50 mM and TBP 0.50 M: under these conditions, the cubic phase was obtained, which is the crystalline phase of ABCs having the highest σ , due to the presence of interconnected bi-continuous channels. The conductivity trend showed in Table 18 followed the sequence P-123 > F77 > PEG: in fact, due to the lower amount of PEO units in the P-123 gel-polymer electrolyte, the degree of hydration per PEO chain was found to be higher when compared to that of F77, which promoted ion mobility. Moreover, the size of the F77 micellar core (Fig. 24A) was smaller than that of P-123. As regards PEG, it did not form a microcrystalline ordered gel, thus I_3^- ions could not diffuse so faster as in P-123- and

Table 18 Effect of the ABCs: H_2O 35: 65 matrices on both electrolyte (BMII 2.0 M, I_2 50 mM and TBP 0.50 M) and DSSC PV parameters ($P_{in} = 1$ sun). An ACN-based cell is also reported for comparison purposes. Dye: D907. Adapted and reprinted from ref. 314

F77-based electrolytes. However, as shown in Table 18, despite the lower σ , when introduced in quasi-solid DSSCs F77-based systems demonstrated higher J_{sc} and V_{oc} values than P-123-based ones. Indeed, while P-123 formed a gel with very high viscosity, F77 showed remarkable structural polymorphism (Fig. 24B): by lowering its temperature, a conversion into the simple micellar solution having reduced μ occurred. Thus, F77 gels could diffuse into the photoanode nanopores at low temperature (ensuring the efficient regeneration of the oxidized dye in the whole electrode), while maintaining a gel-like consistence above 37 °C. Moreover, the higher PPO content in the P-123 gel hindered the accessibility of TBP, thus lowering the cell photovoltage with respect to that of the F77-based cell. Stability tests were carried out over 500 h and only around 3% reduction in the overall performance of DSSCs based on ABCs occurred; in contrast, dye desorption was observed in the case of PEG-based gels, due to the highly hydrated polymer chains.

Another elegant strategy to prepare electrolytes being effectively water as well as solvent leakage resistant is the use of thixotropic three-dimensional (3D) networks,³¹⁵ the latter obtained by a chemical, photochemical or thermal route.^{316,317} To this purpose, Ko and coworkers used xanthan gum ($M_n = 2 \times 10^6$), a water soluble and environmentally friendly polysaccharide rich in hydroxyl groups ready to form a 3D network, the viscosity of which reversibly decreases upon application of an external stress.³¹⁸ After a thorough optimization of the stability of all the ingredients in the reactive formulation – a key aspect in this field, where apolar, polar and ionic substances are concurrently used – a 1:1 mixture of xanthan gum (3 wt% in H₂O) and liquid electrolyte (PMII 4.0 M, I₂ 0.30 M, TBP 1.5 M

Table 19 Effect of liquid (PMII 4.0 M, I₂ 0.30 M, TBP 1.5 M and GuSCN 0.20 M in MPN or MPN : H₂O 1:1) and quasi-solid (1:1 in xanthan gum 3 wt% in H₂O and PMII 4.0 M, I₂ 0.30 M, TBP 1.5 M and GuSCN 0.20 M in MPN) electrolytes on different electrochemical and PV parameters ($P_{in} = 1$ sun). Dye: TG6. Adapted and reprinted from ref. 318

Matrix	$\sigma (mS cm^{-2})$	^I) $D(I_3^-)$ (cm ² s	$^{-1}$) $J_{\rm sc}$ (mA cm	n^{-2}) $V_{\rm oc}$ (V)	FF	η (%)	Electrolyte	$D(I_3^{-})(cm^2 s^{-})$	¹) $J_{\rm sc}$ (mA cm	$^{-2}) V_{\rm oc} (V)$) FF	η (%)
ACN P-123 F77 PEG		$\begin{array}{c} - \\ 4.3 \times 10^{-7} \\ 3.4 \times 10^{-7} \\ 2.6 \times 10^{-7} \end{array}$	16.8 4.6 6.2 2.1	0.676 0.491 0.595 0.600	0.57 0.53	1.3 2.1	MPN MPN:H ₂ O MPN:H ₂ O:xantan gum	$\begin{array}{c} 1.15 \times 10^{-5} \\ 5.60 \times 10^{-6} \\ 3.78 \times 10^{-6} \end{array}$	12.86 9.69 9.49		0.61 0.76 0.77	4.99

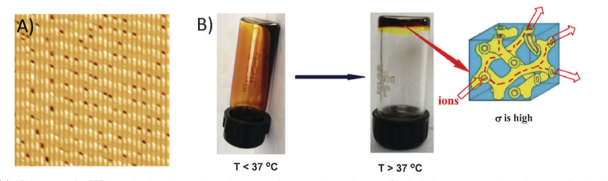


Fig. 24 (A) AFM image of a F77-based gel-polymer electrolyte coated onto a glass substrate. Black dark spots prove the existence of a circular area ranging from 3 to 5 nm, which might be the core of F77 micelles consisting of hydrophobic PPO moieties. The contrast of this area to the rest of the film could be due to the penetration of relatively hydrophobic I_3^- into the PPO core. (B) Photos and schematic illustration of the structural polymorphism of a F77-based gel-polymer electrolyte. Adapted and reprinted from ref. 314.

$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Anode ^a	Dye^{b}	Electrolyte	$H_2O(\%)$	$Cathode^c$	$\int_{\rm sc} J_{\rm sc} ({ m mA} { m cm}^{-2})$	$V_{\rm oc}$ (V)	FF	μ (%)	$\eta_{\mathrm{loss}}{}^d$	Ref.
N3 III 0.100 M and b.100 M/L in MNN 3 14.1 0.54 N3 110 0.100 M, HDIM MNN 3 - 14.1 0.54 N31 10.000 M, HDIM MNN 3 - 14.1 0.56 N31 10.000 M, HDIM MNN 5 Pt 10.9 0.23 N719 DMPH 0.50 M, LIO M, J.5 00 M and BBEG, 10 Pt 4.119 0.6 N719 DMPH 0.50 M, LIO JM, J.5 00 M and BBEG, 10 Pt 4.11 0.69 N719 DMPH 0.50 M, LIO JM, J.5 00 M and BBEG, 10 Pt 8.1 0.69 N719 DMPH 0.50 M, LIO JM, J.5 00 M and BBEG, 10 Pt 8.1 0.69 N711 DM MJ, 20 M, LIO JM, J.5 00 M, ABEG, 10 Pt 8.1 0.69 N71 DM MJ, 20 M, J.6 00 M, 10 AM and TBP 0.50 M in MPH 10 Pt 8.3 0.67 N811 DM MJ, 20 MM, 20 MM AND Pt Pt 11.55 0.68 0.55 N811 DM MJ, 20 MM, 20 MM AND Pt Pt 8.3 0.67 N811	$TiO_2 (11.5 \ \mu m \ T + 14.5 \ m c1) + m c1$		BMII 0.70 M, I ₂ 30 mM, GuSCN 0.10 M, TBP	0.018	Pt	17.5	0.85	0.73	10.8	2% (1000 h, 1 sun)	172
N3 Lif 0.10 M and L 10 mi MNN 4 Pt foil 1.05 0.541 N719 L010.00 M and L 10 mi MNN 4 Pt 11.17 0.61 N719 L010.00 M HID 0.60 ML 3 S0 mM APR 4 Pt 11.17 0.63 N719 DMPI 0.50 ML 10.10 ML 4.50 mM and BBEG, 10 Pt 4.19 0.8 N719 DMPI 0.50 ML 10.10 ML 4.50 mM and BBEG, 10 Pt 4.19 0.8 N719 DMPI 2.00 ML 4.5.00 ML 4.5.0 mM and BBEG, 10 Pt 8.1 0.69 N719 DMPI 2.00 ML 4.0.10 ML 4.5.00 ML 4.6.0 MI 10 Pt 8.1 0.69 N719 DMMI 1.00 ML 4.0.10 ML 4.0.10 M m ACN 10 Pt 8.3 0.67 N82 DMMI 1.00 ML 4.0.10 M m ACN MPP 1.00 M m ACN 10.10 ML 19.0.10 M m ACN 10.66 0.71 N82 DMMI 1.00 ML 4.0.10 M m ACN MPP 1.00 M m ACN 10.66 0.71 N82 DMMI 1.00 ML 4.0.10 M m ACN 10.67 0.82 0.66 N82 DMMI 1.00 M M MNN MCACN 20	µш эь) т ны₄ TiO ₂	N3	Lil 0.50 M and 50 mM I_2 in MPN	3	I	14.1	0.54	0.59	4.5	I	124
NTJ9 Lin OLO M, HDMI LOGO M, JS OTIMA JR MAPI A Pr Lin Z 0.63 NT19 US 0.500 M, EVR 2%, and TBME 2%, in MPI 5 P 10.9 0.52 NT19 D500 M in ACN 5 P 10.9 0.52 NT19 D500 M in ACN 5 P 1.0 0.53 NT19 D500 M in ACN 10.20 M, L9 0.20 M in MPI 10 P 4.19 0.83 N3 Lin 0.50 M, LP 0.20 M and TBP 0.50 M in MPI 10 P 8.1 0.69 N3 Lin 0.50 M, LP 0.28 M and L 0.50 M in MPI 10 P 8.1 0.69 N3 Lin 0.50 M, L9 0.30 M and TBP 0.53 M in MPI 10 P 8.3 0.67 N3 Lin 0.50 M, L9 0.01 M and LPP 2.00 M in PDH P 8.3 0.65 N3 Lin 0.50 M, L9 0.01 M and TBP 0.50 M in MPI 10 P 8.3 0.65 N3 Lin 0.50 M, L9 0.01 M and CBSC 0.10 M and TBP 2.0 P P 2.56 0.58 N3 Lin 0.50 M, L9 0.01 M and CBSC 0.10 M. TBP 2.0	$TiO_2 (4.5 \mu m)$	N3	Lif 0.10 M and I ₂ 10 mM in MPN	4	Pt foil	1.96	0.541	0.67	4.8		43
N3 I_0.30 M, PVP 2% and TBMB 2% in MFII 5 Pt 109 0.32 N719 DMPII 0.50 M, II.0.10 M, I_2 30 MM and BBEG, if = 10.50 M, II.0.10 M, I_2 30 MM and BBEG, if = 10.50 M, II.0.10 M, I_2 30 M, TPP 0.30 M, IM PII 5 7 0.39 0.38 SJ III DMPII 0.50 M, II.0.10 M, I_2 30 M, TPP 0.30 M, IM MDI 10 Pt 8.1 0.69 Jim + X83 DMI PLO SM, II.0.10 M, I_2 0.30 M, TPP 0.30 M, IM MDI 10 Pt 8.1 0.69 Jim + X83 DII 0.50 M, IJ.0.20 M, TPP 0.30 M, IM MDI 10 Pt 8.3 0.67 Jim + X83 DII 0.50 M, I_3 0.50 M, M MDI 10 Pt 8.3 0.67 Jim + X83 DII 0.50 M, I_3 0.50 M, M MDI 10 Pt 8.3 0.67 Jim + Z07 DII 0.50 M, II DRD DM ANDI 10 Pt 5.5 0.8 Jim + Z05 DM MPP DM ANDI DM ANDI </td <td>BL + TiO₂ (10 μm)</td> <td>N719</td> <td>Lil 0.10 M, HDMII 0.60 M, l₂ 50 mM, TBP 0.50 M and Triton X-100 20 mM in MPN</td> <td>4</td> <td>Pt</td> <td>11.77</td> <td>0.81</td> <td>0.62</td> <td>5.9</td> <td>29% (7 d, dark)</td> <td>156</td>	BL + TiO ₂ (10 μ m)	N719	Lil 0.10 M, HDMII 0.60 M, l ₂ 50 mM, TBP 0.50 M and Triton X-100 20 mM in MPN	4	Pt	11.77	0.81	0.62	5.9	29% (7 d, dark)	156
NT9 DMPT 0.50 M, Lif 0.10 M, L ₃ 50 mM and BBEG ₆ 10 Pt 4.19 0.8 $35 \ Jml$ WR2HA MET 0.00 M, L ₃ 20.00 M in ACN Pt 5.75 0.8 $55 \ Jml$ MK2HA MPT 2.0M, Li 0.10 M, L ₃ 20.00 M in ACN Pt 5.75 0.8 719 MPT 2.0M, Li 0.10 M, L ₃ 20.00 M in ACN Pt 5.75 0.8 710 MPT 2.0M, Li 0.10 M, L ₃ 20.30 M in MPT 10 Pt 8.3 0.67 710 MPT 2.0M, Li 0.10 M, L ₃ 0.25 M in MPT 10 Pt 8.3 0.67 710 TG6 PMIL 2.0M, L ₃ 0.10 M and TPP 0.36 M in MPT 10 Pt 8.3 0.67 710 CV-G C6(NO ₃)s 0.10 M and TPP 0.36 M in MPT 10 Pt 7.368 0.62 710 CV-G C6(NO ₃)s 0.10 M and C6(NO ₃ M, TPP 0.36 M in MPT 10 Pt 7.368 0.65 710 CV-G C6(NO ₃)s 0.10 M and TPR 0.36 M in TPP 2.46% 50 Pt 7.368 0.65 710 CV-G C6(NO ₃)s 0.10 M and TPR 0.36 M in TPP 2.4%	$TiO_2 (9 \mu m) +$	N3	I_2 0.30 M, PVP 2% and TBMB 2% in MPII	IJ	Pt	10.9	0.52	0.42	2.4		303
NT9 NT19 PRT 200M, J10,010M, J, 50,00M, and BBBG, PF 5.75 0.8 55 Jm) + MK2HA MK2140, M, J10,010M, J, 50,00M, TBP 0.50 M in MPT 10 PF 8.1 0.69 75 Jm) + MK2HA MFT 2.00 M, J10,010M, J, 50,00M, TBP 0.50 M in MPT 10 PF 8.1 0.69 710 MFT 2.00 M, J10,010M, and TBP 0.50 M in MPT 10 PF 8.3 0.67 710 TGL TGG PMT 10, 01, M, J10,010M, and TBP 0.50 M in MPT 10 PF 8.3 0.67 710.1 TGG PMT 10, M, J10,010M, and TBP 0.50 M in MPT 10 PF 8.5 0.84 711 TGG PMT 10, M, J10,010M, and Ce(NO)J, 50 mM, TBP 20 PF 0.52 0.62 711 TCGG PMT 10, MJ, J10,010M, TBP 50 PF 5.2 0.53 711 CV+G Ce(NO,J1,010 M and Ce(NO,J1,50 mM, TBP 50 PF 5.2 0.62 711 CV+G CoJM, J13,010 M and Ce(NO,J1,50 m M, TBP 50 PF 5.2 0.62 <	TiO_2 (12 µm)	N719	Х,	10	Pt	4.19	0.8	0.69	2.31	7% (135 h)	145
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	TiO ₂ (12 μ m)	N719	М,	10	Pt	5.75	0.8	0.64	2.96	40% (135 h)	145
N3 III 0.50 M, TBP 0.58 M and I_2 0.30 M in MPI 10 Pt 8.8 0.67 III X03 TiCla, TG Nin Nin Nin 0.50 M, I, 0.50 M, I, 0.53 M, 0.51 M, MAPI 10 Pt 8.8 0.67 TiCla, TG PG PMII J. 0.0, 1, 0.50 M, I, 0.51 M, 0.53 M, 0.51 M, MAPI 10 Pt 8.8 0.67 TiCla, TG PMI J. 0.0, 1, 0.1 M and CelNO ₁ J, 80 mM in EOH 20 Pt 8.5 0.84 D516 PMII 2.0 M, I, 40 mM, PGG 4 w% and PY2 wt% 5 0 Pt 7.368 0.653 D516 PMII 2.0 M, I, 30 mM, GUSCN 0.10 M, TBP 50 Pt 5.22 0.75 D5116 PMII 2.0 M, I, 30 mM, GUSCN 0.10 M, TBP 50 Pt 7.46 0.77 V+7 Jim TG6 PMII 2.0 M, I, 30 mM, GUSCN 0.10 M, TBP 50 Pt 5.65 0.65 0.50 M and Triton X-100 1% in MPN TG6 PMII 2.0 M, I, 30 mM, GUSCN 0.10 M Pt 7.46 0.77 0.51 M III 2.0 M, I, 30 mM GUSCN 0.10 M, TBP 50 Pt 2.22 </td <td>$TiCl_4 + TiO_2 (25 \ \mu m) + TiCl_4 + TiO_2 (25 \ \mu m) + TiCl_4 + TiO_2 (25 \ \mu m) + TiCl_4$</td> <td></td> <td>MPII 2.0 M, Lil 0.10 M, I₂ 0.20 M, TBP 0.50 M in</td> <td>10</td> <td>Pt</td> <td>8.1</td> <td>0.69</td> <td>0.72</td> <td>4.02</td> <td>2.5% (200 h, dark)</td> <td>243</td>	$TiCl_4 + TiO_2 (25 \ \mu m) + TiCl_4 + TiO_2 (25 \ \mu m) + TiCl_4 + TiO_2 (25 \ \mu m) + TiCl_4 $		MPII 2.0 M, Lil 0.10 M, I ₂ 0.20 M, TBP 0.50 M in	10	Pt	8.1	0.69	0.72	4.02	2.5% (200 h, dark)	243
N3 Lil 0.50 M, 1_9 0.30 M and TBP 0.58 M in MPH 10 Pt 8.8 0.67 $TIGI_4$ TG6 PMI 1.00 M, 1_9 0.15 M, GuSCN 0.10 M and TBP 20 Pt 15.67 0.68 $TIGI_4$ TG6 PMI 1.00 M, 1_9 0.15 M, GuSCN 0.10 M and TBP 20 Pt 15.57 0.68 $110T_+$ CGCN0,3_10 0.10 M and CG(NO,3, 50 mM in BEOH 30 Pt 5.52 0.623 $110T_+$ CGCN0,3_10 0.10 M and CG(NO,3, 50 mM, GuSCN 0.10 M, TBP 50 Pt 7.368 0.623 $110T_+$ Z007 KG10A, 1_3 0 mM, GuSCN 0.10 M, TBP 50 Pt 7.368 0.653 $110T_+$ Z007 KG10A, 1_3 0 mM, GuSCN 0.10 M, TBP 50 Pt 7.46 0.77 1756 PMII 2.0 M, 1_3 0 mM, GuSCN 0.10 M, TBP 50 Pt 7.46 0.77 $1+7$ µm TG6 PMII 2.0 M, 1_3 0 mM/ TBP 50 Pt 7.46 0.77 $100T$ PMII 2.0 M, 1_3 0 mM/ TBP 50 Pt 7.46 0.77 $100T$ D00T PMII 2.0	TiO,	N3	Lil 0.50 M, TBP 0.58 M and I ₂ 0.30 M in MPII	10	Pt	8.8	0.67	0.72	4.2^{f}	I	293
Implicit Z00. + CDAA PMII 2.00 M, 15 so mM, GuSCN 0.10 M and TBP 10 Pt 15.51 0.68 TICI, TG6 PMII 2.00 M, 15 so mM, GuSCN 0.10 M and TBP 20 Pt 5.52 0.684 $t_{IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII$	TiO_2	N3		10	Pt	8.8	0.67	0.72	4.2 ^f		294
$ \begin{array}{ccccccc} \mbox{TG}(4 & \mbox{TG}) & \mbox{TG}) & \mbox{TG}(4 & \mbox{TG}) & \mbox{TG}(4 & \mbox{TG}) & \mbox{TG}) & \mbox{TG}(4 & \mbox{TG}) & \mb$	BL + 11O ₂ (10 μm) + TiCl ₄	Z90/ + UDCA		10	Ρτ	/0.cl	0.68	0./1	7.6./	7% (1000 n, 1 sun)	128
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	TiO_2^4 (7 μ m) + $TiCl_4$	TG6	PMII 2.0 M, I_2 50 mM, GuSCN 0.10 M and TBP 0.50 M in MDN	20	Pt	8.5	0.84	0.73	5.2	Gain 2% (750 h, 1	120
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$ZrO_2 (17.3 \mu m)$		$Ce(NO_3)_3$ 0.10 M and $Ce(NO_3)_4$ 50 mM in EtOH	30	Pt	0.52	0.624	0.53	0.688		178
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	TiCl ₄ + TiO ₂ (9 μ m T + 5 μ m SL)		KI 0.20 M, I ₂ 40 mM, PEG 4 wt% and PVP 2 wt% in ACN/VAN	50	Pt	7.368	0.625	0.5	2.306	37% (168 h, 1 sun)	310
JK-310 PMII 2.0 M, 12, 50 mM, 100, 170, 170, 17BP 50 Pt 6.62 0.75 $+7$ Jµm TG6 PMII 2.0 M, 15, 50 mM, GUSCN 0.10 M, TBP 50 Pt 7.46 0.77 0.50 M and Triton X-100 1% in MPN 0.50 M and Triton X-100 1% in MPN 50 Pt 7.46 0.77 0.50 M and Triton X-100 1% in MPN 50 Pt 7.46 0.77 0.50 M and Triton X-100 1% in MPN 50 Pt 9.5 0.65 $+7$ Jµm TG6 PMII 4.0 M, 12 0.30 M, TBP 0.75 M and GUSCN 0.20 M 50 Pt 9.7 0.68 μ D907 BMII 2.0 M, 12 50 mM and TBP 0.50 M in PEG 65 Pt 2.1 0.6 μ D907 BMII 2.0 M, 12 50 mM and LiNO3,0.10 M in 65 Ag 5.96 0.46 μ D907 BMII 2.0 M, 12 50 mM and LiNO3,0.10 M in 77 65 Pt 4.4 0.61 μ D907 BMII 2.0 M, 12 50 mM and LiNO3,0.10 M in 77 65 Pt 4.4 0.40 μ D907 BMII 2.0 M, 12 50 mM and 12 0.50 M in PTO	$TIO_2 (T + SL)$	D5L6	$PMH 2.0 \text{ M}, I_2 50 \text{ mM}, GuSCN 0.10 \text{ M}, TBP 0.000 \text{ M}, 0.10 \text{ M}, TBP 0.000 \text{ M}, 0.100 \text{ M}, 0.000 \text{ M}, 0.0000 \text{ M},$	50	Pt	6.22	0.75	0.73	3.4		241
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$TiO_2 (T + SL)$	JK-310	PMII 2.0 M, 12.50 mM, GuSCN 0.10 M, TBP	50	Pt	6.62	0.75	0.76	3.77		241
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$TiO_2 (T + SL)$	D21L6	0.50 M and 111001 X-100 1% IN MPN PMII 2.0 M, 12 50 MM, GUSCN 0.10 M, TBP	50	Pt	7.46	0.77	0.77	4.41		241
$+7 \ \mu m$ TG6 PMII 1.96 M, I_2 0.15 M, TBP 0.75 M and GuSCN 50 Pt 9.7 0.68 μm) + D907 BMII 2.0 M, I_2 50 mM and TBP 0.50 M in PEG 65 Pt 2.1 0.6 μm) + D907 BMII 2.0 M, I_2 50 mM and TBP 0.50 M in PEG 65 Pt 2.1 0.6 μm) + D907 BMII 2.0 M, I_2 50 mM and LiNO ₃ 0.10 M in 65 Pt 2.1 0.6 μm) + D907 BMII 2.0 M, I_2 50 mM and LiNO ₃ 0.10 M in F123 65 Pt 4.6 0.496 μm) + D907 BMII 2.0 M, I_2 50 mM and TBP 0.50 M in P123 65 Pt 4.24 0.51 μm) + D907 BMII 2.0 M, I_2 50 mM and 50 mM Ce(NO ₃) ₄ in EtOH 65 Pt 4.94 0.55 μm) + D907 BMII 2.0 M, I_2 50 mM and TBP 0.50 M in F77 65 Pt 4.94 0.55 μm) + D907 BMII 2.0 M, I_2 50 mM and TBP 0.50 M in F77 65 Pt 4.94 0.55 μm) + D907 BMII 2.0 M, I_2 50 mM and I_2 25 mM in EtOH 65 Pt 4.94 0.55 μm) + D907	TiO, (12 um T + 7 um	TG6	0.50 M and Triton X-100 1% in MPN PMII 4.0 M. I. 0.30 M. TBP 1.5 M. GuSCN 0.20 M	50	Pt	9.5	0.65	0.77	4.78	7% (288 h. dark.	318
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	SL + TiCl ₄		and xanthan gum 3 wt% in MPN	2	2	2				65 °C, 85% RH)) +
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	TiO ₂ (12 μ m T + 7 μ m SI.) + TiCl.	1G6	PMII 1.96 M, I ₂ 0.15 M, TBP 0.75 M and GuSCN 0.10 M in MPN	50	Pt	9.7	0.68	0.76	4.99	48% (288 h, dark, 65 °C. 85% RH)	318
$ \begin{array}{lcccccccccccccccccccccccccccccccccccc$	$BL + TiO_2 (12 \ \mu m) + mc H$	D907	BMII 2.0 M, I ₂ 50 mM and TBP 0.50 M in PEG	65	Pt	2.1	0.6	0.44	0.6		314
$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	$TiO_2 + Au NPS$	MB	0.10 M, FeCl ₃ 50 mM	65	Ag	5.96	0.46	0.65	0.95	I	204
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	BL + TiO ₂ (12 μ m) +	D907	BMII 2.0 M, I ₂ 50 mM and TBP 0.50 M in P123	65	Pt	4.6	0.491	0.57	1.3		314
RhS Ce(NO ₃) ₃ 0.10 M and 50 mM Ce(NO ₃) ₄ in EtOH 65 Ag 10.9 0.496 μm) + D907 BMII 2.0 M, I_2 50 mM and TBP 0.50 M in F77 65 Pt 4.94 0.55 μm) + D907 BMII 2.0 M, I_2 50 mM and TBP 0.50 M in F77 65 Pt 4.94 0.55 $+ UV/O_3$ N3 + TBP KI 0.50 M and I_2 25 mM in EtOH 65 Pt 5.8 0.6 PG Nal 1.0 M and I_2 0.10 M 100 Pt 2.2 th 0.44 th N3 LiI 0.10 M, I_2 10 mM, 0.5 wt% k-carrageenan 100 Pt 2.69 0.42 N3 LiI 0.50 M and I_2 25 mM 0.5 wt% k-carrageenan 100 Pt 2.14 0.44 N3 KI 0.50 M and I_2 25 mM 100 Pt 2.14 0.47	$TO_2 (15 \mu m)$	N719	KI 0.50 M and I_2 25 mM in EtOH	65	Pt	4.24	0.51	0.60	1.3	Ι	138
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	TiO ₂ (1.8 μ m) TiO ₂ (15 μ m)	RhS N3	$Ce(NO_3)_3$ 0.10 M and 50 mM $Ce(NO_3)_4$ in EtOH K1 0.50 M and T. 25 mM in FtOH	65 65	Ag Df	10.9 4 94	0.496	0.27	1.49		174 138
+ UV/O3 N3 + TBP KI 0.50 M and I2 25 mM in EtOH 65 Pt 5.8 0.6 PG NaI 1.0 M and I2 0.10 M 10 0 M 100 Pt 2.2 ^h 0.44 ^h N3 Li1 0.10 M, I2 10 mM, 0.5 wt% k-carrageenan 100 Pt plate 2.69 0.42 N719 KI 0.50 M and I2 25 mM 100 Pt plate 2.69 0.44 N3 KI 0.50 M and I2 25 mM 100 Pt 2.14 0.44 N3 KI 0.50 M and I2 25 mM 100 Pt 2.14 0.44	$BL + TiO_2 (12 \ \mu m) + TiO_2 $	D907	BMII 2.0 M, I ₂ 50 mM and TBP 0.50 M in F77	65	Pt	6.2	0.595	0.53	2.1	3% (500 h)	314
N3 Li1 0.10 M, I ₂ 10 mM, 0.5 wt% k-carrageenan 100 Pt plate 2.69 0.442 and HNO ₃ (pH 2) 7719 K1 0.50 M and I ₂ 25 mM 100 Pt 2.14 0.44 N3 K1 0.50 M and I ₂ 25 mM 100 Pt 3.61 0.47 N3 K1 0.50 M and I ₂ 25 mM	TiO ₂ (20 μm) + UV/O ₃ TiO ₂ (3.3 μm)	N3 + TBP PG		65 100	Pt Pt	$5.8 \\ 2.2^{h}$	$\begin{array}{c} 0.6 \\ 0.44^{h} \end{array}$	0.63	2.2	— 0% (24 h, under	203 256
and HNO ₃ (PH 2) N719 KI 0.50 M and I ₂ 25 mM 100 Pt 2.14 0.44 N3 KI 0.50 M and I ₂ 25 mM 100 Pt 3.61 0.47	${ m TiO_2}$ (10 ${ m \mu m}$)	N3	i wt% к-carrageenan	100	Pt plate	2.69	0.442	0.48	0.586	lıght") —	328
	TiO ₂ (15 μm) TiO ₂ (15 μm)	N719 N3		100 100	Pt Pt	$2.14 \\ 3.61$	$0.44 \\ 0.47$	$0.64 \\ 0.65$	0.6 1.1	— 138 50% (75 d, dark, RT) 138	$138 \\ 138$

Review Article

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Anode ^a Dye ^b	Electrolyte	H_2O (%)	$Cathode^{c}$	$\int_{ m sc} J_{ m sc} (m mA~ m cm^{-2})$	$V_{\rm oc}$ (V)	FF	(%) <i>u</i>	$\eta_{\mathrm{loss}}^{~~d}$	Ref.
$\operatorname{TiCl}_{4} + \operatorname{TiO}_{2} (20 \ \mu \text{m T} +]\text{K-259}$	PMII 2.0 M, I ₂ 50 mM, GuSCN 0.10 M, TBP	100	Pt	2.28	0.66	0.79	1.16		238
$\frac{4}{10}$ m SL J^{+} 1104 TiCl ₄ + TiO ₂ (3 µm T + 5 V35 m cr) + m:Cl	0.50 M and 11.00 A-100 1% KI 4.0 M and 1 ₂ 20 mM, pH 8	100	PEDOT	5.60	0.485	0.63	1.71	Gain 58% (18 d,	249
TiCl ₄	Z907 + CDCA Nal 2.0 M, I ₂ 20 mM, GuSCN 0.50 M and CDCA 100	100	Pt	4.91	0.62	0.64	1.88	uaik) —	206
TiO ₂ cis-	Ru(ii) $L_2(H_2O)_2^{2-i}$	KI 0.10 M and I ₃ ⁻ 1.0 mM	100	Pt flag	0.38	0.52	0.7	2.0 [′]	0% (4 d, 0.07
118 TiO ₂ (2.8 μ m T + 3.1 μ m D205	TEMPOL 1.0 M and NaBF $_4$ 1.0 M	100	Pt:Teflon	4.5	0.69	0.64	2.1	I	274
TiCl ₄ + TiO ₂ (20 μ m T + JK-262	PMII 2.0 M, I ₂ 50 mM, GuSCN 0.10 M, TBP	100	Pt	3.78	0.68	0.82	2.1		238
$4 \ \mu m \ sL) + 1104 \ TiO_2 (3 \ \mu m \ T + 5 \ V35 \ m m \ cr) + m \ cr)$	V_{120} M and 111001 A-100 1% KI 4.0 M, I_2 20 mM and CDCA until saturation	100	Pt	4.86	0.600	0.76	2.20		249
$\lim_{T \to 0} \sum_{1} + 1101_{4}$ TiO ₂ (7 µm) + TiCl ₄ TG6	PMII 2.0 M, I ₂ 50 mM, GuSCN 0.10 M, TBP 0 50 M and Triton X-100 1%	100	Pt	4.7	0.74	0.69	2.4	I	120
TiCl ₄ + TiO ₂ (3 μ m T + 2 D45 \dots cr) + TiCl ₄	$TT = EMI^+ 0.20 M$, DTT 0.20 M, TBP 0.50 M and 100	100	Pt	7.2	0.65	0.55	2.6	37% (4 h, 1 sun, RT) 193) 193
TiO ₂ (8.9 μ m) + TiCl ₄ TG6	1.% 111001 A-100 Nai 2.0 M, I ₂ 20 mM, GuSCN 0.50 M and CDCA 100	100	Pt	7.34	0.59	0.63	2.64		206
TiCl ₄ + TiO ₂ (3 μ m T + 5 V35	nM and CDCA until saturation	100	PEDOT	6.85	0.650	0.67	3.01		249
$TiO_2 (4.4 \ \mu m) + TiCl_4$ N719	Nal 2.0 M, I ₂ 20 mM, GuSCN 1.0 M, HNO ₃ (pH	100	Pt	8.5	0.59	0.63	3.08		206
TiCl ₄ + TiO ₂ (3 μ m T + 2 D51 μ m Sr) + π iCl	3) and CDCA until saturation TTT-Emf 0.20 M, DTT 0.60 M, TBP 0.50 M and 100 106. Triton V-100	100	Pt	9.5	0.61	0.59	3.5		193
TiO ₂ (13.3 μ m) N719 + CDCA		100	Pt	10.97	0.53	0.68	3.96	63% (50 d, 1 sun, RT_11V filter)	163
TiO ₂ (4.5 μ m) + TiCl ₄ D149 BL + TiO ₂ (1.3 μ m T + 5 MK-2 μ m SL) + TiCl ₄	Gul 8.00 M, I ₂ 20 mM and CDCA until saturation 100 $K_4Fe(CN)_6$ 0.40 M, $K_3Fe(CN)_6$ 40 mM, KCl 0.10 100 M, Trizma-HCl buffer 50 mM (pH 8) and Tween 2000 100	100	Pt Pt mirror	10.02 7.2	0.61 0.76	0.67 0.75	4.06 4.1		206 182
TiO ₂ (1 μ m T + 3 μ m SL) MK-2	$[Co(bpy)_3]^{2+}$ 0.20 M, $[Co(bpy)_3]^{3+}$ 40 mM, NMBI 100 0.70 M NMMI 3nd bEC 300 100	100	Pt	8.3	0.68	0.72	4.2	10% (90 d, dark)	194
TiO ₂ (9 μ m T + 3 μ m SL) MD3	TEMPO 0.40 M, NOBF ₄ 0.40 M, Lil 0.10 M, I ₂ 50 100 mM, DMPII 0.60 M, GuSCN 0.10 M and Tween 2000 10.60 M	100	Pt	9.56	0.77	0.67	4.96	I	258
TiO ₂ (1 μ m T + 3 μ m SL) MK-2	[Co(br)3] ²⁺ 0.20 M, [Co(bpy) ₃] ³⁺ 40 mM, NMBI 0 70 M NMBI and PEG 300 1%.	100	Pt:ITO	9.8	69.0	0.74	5	Gain 2% (2 d, dark)	194
TiO ₂ (1 μ m T + 3 μ m SL) MK-2 + TiCl.	$[Co(bpy)_3](NO_3)_2 0.20 M, [Co(bpy)_3](NO_3)_3 40 mM NMBI 0.70 M and DEC 200 1 meV.$	100	Pt	10.17	0.821	0.68	5.64	0% (500 h, dark)	210

^{*a*} T means transparent layer, SL scattering layer, BL blocking layer. When "TiCl₄" is written before "TiO₂", it means that the treatment has been carried out to fabricate a BL. When "TiCl₄" is reported after "TiO₂", it means that further data were not reported. ^{*b*} If a coadsorbent was employed, it has been reported in the table. ^{*c*} When "Pt" alone is reported, it means that Pt has been sputtered or thermally deposited following the traditional procedures. ^{*d*} If available, aging test conditions have been reported in the table. ^{*c*} Photoanodes were dipped in 1% accetic acid solution in toluene for 5 min, followed by rinsing with toluene. ^{*f*} Same PV data were reported by the same group in two different papers. ^{*g*} The cell was stored for 7 months in the dark before starting the aging test. ^{*h*} *P*_{in} not reported. ^{*i*} It is the 2,2'-bipyridyl-4,4'-dicarboxylate ligand. ^{*j*} *P*_{in} = 0.07 sun.

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and GuSCN 0.20 M in MPN) was selected. As reported in Table 19, resulting aqueous DSSCs, both liquid and quasi-solid, showed higher $V_{\rm oc}$ and FF, but lower $J_{\rm sc}$ values with respect to the MPNbased liquid counterpart. The decrease in J_{sc} was related to the desorption of the dye from the TiO₂ electrode surface. Actually, even if the hydrophobic TG6 dye was used, it was not possible to fully prevent the dye molecules from being desorbed; moreover, the I_3^- diffusion coefficient was decreased when water was introduced. As regards $V_{\rm oc}$, both E of the electrolyte and the CB of the TiO₂ positively shifted in the presence of aqueous electrolytes; however, the shift was found to be more significant for E, thus resulting in enhanced photovoltage values. Noteworthy, the quasi-solid aqueous cell exhibited a performance in terms of overall efficiency comparable with that of the liquid one (i.e., 4.78% vs. 4.99%): this was ascribed to the thixotropic nature of the gel-polymer matrix, which could infiltrate as a liquid into the photoanode, in spite of the scarce wetting ability typically observed when solid polymer electrolytes are used, 319,320 being their macromolecular chain radius of gyration larger than the pore size of TiO₂.^{69,321,322} During extreme 288 h aging tests (65 °C and 85% RH), the water-based liquid DSSC retained around 50% of its initial efficiency, while the quasi-solid one remarkably provided the 93% of its initial value. Such results demonstrated that the xanthan gum was able not only to prevent the leakage of the solvent from the DSSC, but also to reduce the number of dye molecules desorbed from the photoanode at high temperature.

As evidenced in the present section, the overall performances of quasi-solid water-based DSSCs are definitely interesting, particularly because of the excellent long-term stability demonstrated by these devices. Nevertheless, there is still room for further improvements. As an example, in terms of materials selection, a highly ambitious challenge would be the replacement of the oil-derived polymer matrixes by means of biosourced ones.^{57,323-326} In the field of water-based DSSCs, a first attempt has been proposed by Kaneko and coworkers, who reported that polysaccharides such as agarose and $\kappa\text{-carrageenan}^{327}$ can form a tight and elastic solid able to retain the aqueous electrolyte (LiI 0.10 M and I₂ 10 mM in H₂O, pH 2).³²⁸ Detachment of the N3 dye from the TiO₂ electrode was not observed even after prolonged irradiation times; moreover EIS analysis showed that the electron donation from the I^- to the oxidized sensitizer took place in a similar way both in the quasi-solid and in the liquid state. Unfortunately, the authors published only a preliminary efficiency value of 0.586% (J_{sc} = 2.69 mA cm⁻², V_{oc} = 0.442 V, FF = 0.48) in the presence of 0.5 wt% k-carrageenan, without any optimization of the electrolyte composition which, in our opinion, could have led to definitely higher performance.

14. Conclusions

DSSC represents the most widely studied device for the conversion of solar energy into electricity, particularly when low costs and good performance are envisaged. In this review article, the key role of water, being one of the most promising research After an exhaustive analysis of the interactions between water and the other cell components – previously considered exclusively harmful – we have shown how the solvent-of-life may adequately replace the traditionally used organic solvents. This would make DSSCs cheaper as well as less dangerous in terms of flammability and toxicity, thus even more environmentally friendly. In order to fabricate efficient aqueous DSSCs, several research groups worldwide are involved in an impressive scientific work, proposing new dyes, photoelectrodes, counter electrodes, redox couples, additives and sealants. As summarized in Table 20, the results obtained so far in terms of efficiency and durability are promising, sometimes outstanding, and fully justify the recent boom in the research and development of aqueous DSSCs.

Summarising, it seems that the main factor that negatively affects the efficiency of aqueous DSSC if compared to those based on organic solvents is the quality of the photoanode/ electrolyte interface. Indeed, excessive hydrophilicity of the dyesensitized surface favours sensitizer molecule desorption, thus decreasing the photocurrent and the stability over time; on the other hand, highly hydrophobic dyes do not allow the complete wettability of the electrode which, in turn, results in a less effective regeneration process for aqueous electrolytes. An optimum between these two extremes has not been achieved so far, and the creation of an intimate photoanode/electrolyte interface will be one of the main research goals in the coming years. At present, the most effective strategies towards realistic aqueous DSSCs include the use of cobalt complexes (preferably in the gel form) as redox mediators, hydrophobic dyes combined/functionalized with surfactants or weakly hydrophilic dyes obtained by the modification of already available sensitizers, and water-tolerant counter electrodes with a wide surface area. However, the study of aqueous DSSCs is at an early stage and comparative studies that could greatly improve the knowledge of these systems are still lacking.

The sum of the above reported intriguing features accounts for the promising prospects of the novel materials described here to be effectively implemented in the emerging business of aqueous DSSC manufacturing. The use of water in solar energy conversion devices was intended to be a big bet by a great part of the scientific community but, based on the results discussed in this review article, it may very soon become the key for success of this green technology to finally enter into the mass production stage.

Abbreviations

ABC	Amphiphilic block copolymer
ACN	Acetonitrile
AOT	Bis(2-ethylhexyl) sulfosuccinate sodium salt
BBEG _n	Bis-benzimidazole derivatives containing an
	ethylene glycol repeating unit

BMII	1-Butyl-3-methylimidazolium iodide	PVP	Poly(vinylpyridine)
CA	Contact angle	PVPi	Poly(vinylpyrrolidone)
CB	Conduction band	$R_{\rm CE}$	Charge transfer resistance at the counter
CDCA	Chenodeoxycholic acid		electrode
CPMD	Car–Parrinello molecular dynamics	RH	Relative humidity
CTAB	Hexadecyltrimethylammonium bromide	RhS	Rhoeo spathacea
CV-G	Green part of Codiaeum varie	$R_{\rm s}$	Series resistance
D	Electron diffusion constant	RT	Room temperature
D(ion)	Apparent diffusion constant for the given ion	RTIL	Room temperature ionic liquid
DMHII	3-Hexyl-1,2-dimethylimidazolium iodide	TAS	Transient adsorption spectroscopy
DMPII	1,2-Dimethyl-3-propylimidazolium iodide	TBA	Tetrabutylammonium
DRIFT	Diffuse-reflectance infrared Fourier transform	TBAI	Tetrabutylammonium iodide
DSSC	Dye-sensitized solar cell	TBP	4- <i>tert</i> -Butylpyridine
DTT	3,3'-Dithiobis[4-methyl-(1,2,4)-triazole]	TCO	Transparent conductive oxide
E	Redox mediator potential	TDDFT	Time-dependent density functional theory
E_{c}	CB energy	TEMPOL	4-Hydroxy-2,2,6,6-tetramethylpiperidinoxyl
E_{f}	Fermi level electrochemical potential	$TFSI^{-}$	Bistriflimide
EC	Ethylene carbonate	$TT^{-}EMI^{+}$	1-Ethyl-3-methylimidazolium 4-methyl-1,2,4-
EDX	Energy dispersive X-ray analysis		triazole-3-thiolate
EIS	Electrochemical impedance spectroscopy	UV-Vis	Ultraviolet-visible
\mathbf{EMIm}^+	1-Ethyl-3-methylimidazolium	VAN	Valeronitrile
FC-134	N,N,N-Trimethyl-3-(perfluorooctyl sulfonamido)-	V _{oc}	Open-circuit voltage
	propan-1-aminium iodide	3D	Three-dimensional
FF	Fill factor	η	Cell efficiency
FK-1	Triethylammonium perfluorooctane sulphonate	$\lambda_{\rm max}$	Absorption maximum
FTIR	Fourier transform infrared	μ	Viscosity
GBL	γ-Butyrolactone	σ	Ionic conductivity
GF	Glass frit	τ	Electron lifetime
GuI	Guanidinium iodide	χw	Mole fraction of water
GuSCN	Guanidinium thiocyanate		
HDMII	1-Hexyl-2,3-dimethylimidazolium iodide		
HMII	1-Hexyl-3-methylimidazolium iodide	Acknow	vledgements
IPCE	Incident photon to current efficiency		
$I_{\rm sc}$	Short-circuit current		ly acknowledges financial support by DSSCX pro-
$J_{\rm d1}$	Diffusion limited current		010-2011, 20104XET32) from Ministero dell'Istru-
$J_{\rm sc}$	Short-circuit current density	zione, dell'U	niversità e della Ricerca.
L	Electron diffusion length		
MAN	Methoxyacetonitrile	Referen	CA5
MPII	1-Methyl-3-propylimidazolium iodide	Referen	CES
MPN	3-Methoxypropionitrile	1 R. A. M	ulhall and J. R. Bryson, Energy price risk and the
NBBI	<i>N</i> -Butylbenzimidazole		bility of demand side supply chains, <i>Appl. Energy</i> ,
NMBI	<i>N</i> -Methylbenzimidazole		23 , 327–334.
NMP	<i>N</i> -Methylpyrrolidone		, T. Soares, Z. Vale and H. Morais, Distributed
NPs	Nanoparticles		on and demand response dispatch for a virtual
ODTS	Octadecyltrichlorosilane	-	player energy and reserve provision, <i>Renewable</i>
PC	Propylene carbonate		2014, 66 , 686–695.
PEDOT	Poly(3,4-ethylenedioxythiophene)		ndpal and L. Broman, Renewable energy education:
PEG	Poly(ethylene glycol)		status review, Renewable Sustainable Energy Rev.,
PEO	Poly(ethylene oxide)	-	I, 300–324.
PES	Photoelectron spectroscopy		s, C. A. Silva and S. Connors, Design and imple-
PE5 PG			on of hybrid renewable energy systems on micro-
	Pomegranate Irradiation intensity		
P _{in} PMII	Irradiation intensity		nities: a review on case studies, <i>Renewable Sustainable</i>
	1-Propyl-3-methylimidazolium iodide		ev., 2014, 31 , 935–946.
	Doly(propylene ovide)		
PPO PSC	Poly(propylene oxide)		Jebaselvi and S. Paramasivam, Analysis on renewable
PPO PSC PV	Poly(propylene oxide) Perovskite solar cell Photovoltaic		ystems, Renewable Sustainable Energy Rev., 2013, 28,

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