Review Article

A Review on Current Status of Stability and Knowledge on Liquid Electrolyte-Based Dye-Sensitized Solar Cells

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The purpose of this review is to gather the current background in materials development and provide the reader with an accurate image of today’s knowledge regarding the stability of dye-sensitized solar cells. This contribution highlights the literature from the 1970s to the present day on nanostructured TiO2, dye, Pt counter electrode, and liquid electrolyte for which this review is focused on.

1. Introduction

The photovoltaic effect was discovered by the French scientist Antoine Cesar Becquerel in 1839. His observations were presented two subsequent times at the Académie des Sciences in Paris by his son Alexandre Edmond [1]. In their experiences, they shared the observations of an intriguing phenomenon related to the generation of current flowing when setting two platinum plates in contact with an aqueous acidic galvanic cell under illumination. This generation of a galvanic current under light occurs when the surface of one of the two electrodes is modified by halide vapor (iodide, chloride, and bromide). Depending on the experimental procedure for this deposition, they noticed a current flowing in one or in the other way when placed under illumination, likely owing to a halide film exhibiting n or p characteristics. For a long time, this starting point of the photovoltaic effect was considered as a scientific curiosity. A mechanistic explanation to account for this effect was drawn by A. Einstein in 1905, who obtained his first Nobel Prize in Physics for his theory describing the origin of the photoelectric effect in 1921. At that time nobody envisioned that the photonic field would become a new area of strong fundamental and applied researches from which the underlying technologies would be a stake for human beings. The starting point for PV applications goes back to the Cold War and the two superpowers’ irrational objective to become the first nation to invade the Moon. Thanks to a worldwide support for billions of dollars, Chapin Pearson and Fuller from Bell Laboratories pioneered the silicon p-n technology, paving the way to a substantial enhancement in the light-to-electricity power conversion efficiency (PCE) compared to the selenium-based electrodes exhibiting at best ca. 1%. The principal objective of this new p-n Si technology was to supply the spatial vehicles in energy by making use of extraterrestrial sun power. With its 1.1 eV indirect bandgap affording panchromatic light absorption, they first demonstrated 4.5% power conversion efficiency in 1954 and 6% a few months later and even 8% was reported in the pioneering patent published in 1957 [2, 3]. Solely one year was required for technological transfer to Western Electric. They were the first to commercialize the product to a “large” public with small systems of 14 mW exhibiting 2% conversion efficiency for 25 $ (1500 $ per W).

Photovoltaic research intended public democratization after the two petrol crises and the ensuing financial crises in 1973 and 1979. Because of the soaring price for crude petrol and the public awareness of its limitations in terms of price stability and availability on Earth, research on alternative low-cost energy production and storage has been prompted. This has largely contributed to the development of PV but also
of electrochemical batteries through the discovery of electrochemical insertion compounds which are currently used in lithium and lithium-ion technology. The most prospective low-cost PV technology is the third generation developed at the end of the 1980s. Organic PV (OPV) and hybrid organic/inorganic dye-sensitized solar cells are fulfilling not only these low-cost expectations and low environmental footprint requirements but also the easy processing, promising a fast technological transfer. What the history tends to forget is that forward-thinking Elliot Bermann envisioned already organic PV back in the late 1960s. He was supported by Exxon Company which integrated prospection for PV cutting-edge research relevant for 30 years ahead in their development projects. They rapidly cost-cut the conversion price by 5 times for the first two years’ activity (from 100 to 20 $/W_p$).

This review is dedicated to the second member of the third generation, namely, dye-sensitized solar cells (DSCs) [4]. Its working principle lies in complete disruption with the other technologies: polycrystalline materials of large bandgap are used, high purity of the materials is not necessarily required, nanocrystalline semiconductors contain large amount of point defects, the working principle relies on electrochemical processes, exciton separation and transport are detached processes, and finally no internal electric fields onset the charge separation (Figure 1). The original idea of providing light sensitiveness to larger bandgap semiconductors has more than one hundred years’ history. It has been prompted by Professor Vogel who pioneered a dye-sensitized silver halide emulsion for practical use in argentic photographic films [5]. This principle was turned away from its primary scope to energy related applications in particular by Gerischer and Tributsch in 1968 and 1972 who used ZnO single crystals sensitized by chlorophyll [6–8]. New impetus was finally given by O’Regan and Graetzel who published on the sensitization of mesoscopic film of nanocrystalline anatase TiO$_2$ particles enabling, in association with red heteroleptic ruthenium (+II) polypyridyl complexes, a remarkable enhancement in the light harvesting properties of the photoanode [4].

High photoelectrochemical stability of DSC was rapidly supported by the fact that Ru(II) polypyridyl complexes in association with TiO$_2$ or ZnO can endure over $10^7$–$10^8$ turnovers of electron injection from dye excited state to electron acceptor levels in the semiconductor which in turn would guarantee more than 10 years’ lifetime of the panel [9]. Rapidly passing the threshold of 10% power conversion efficiency (PCE) in 1993 [10], despite endless efforts on material development, the efficiency had stagnated for a long time in the range of 11.0–11.5% [11–16] before scoring three new subsequent records, first 12.3% and 13.0% under A.M. 1.5 G conditions (100 mW/cm$^2$) using organic dye molecules in association with a stronger oxidant redox active cobalt polypyridyl complex [17, 18]. More recently, certified 14.1% and even certified 17.9% PCE have been achieved by...
replacing the dye with a hybrid organic/inorganic lead halide perovskite absorber and the liquid electrolyte by a solid hole transporting material (HTM) [19, 20] giving birth to a new technology called perovskite solar cells, for which progresses are remarkably fast.

Beside these excellent performances of light-to-electricity power conversion efficiency which clearly meets the standard requirement for market introduction, the race for highest PCE outshined the requirement for high stability and ageing predictability for mass industrialization in many aspects. The much lower number of publications in this area witnesses this deficiency. Matsumura et al. in 1980 already suspected issues of dye stability in devices artificially mimicking the photosynthetic processes as dye-sensitized solar cells [21].

The scope of this review is to provide the reader with an overview about the current material progress which has been realized and the knowledge acquired on the stability issues of dye-sensitized solar cell technology. In this review, we will outline the recent development made in dyes and in electrolyte formulation. We will also gather the actual knowledge regarding the degradation mechanisms which are the source of efficiency decreases under accelerated ageing protocol (IEC61646). The review is then separated into three independent sections: a rapid reminder about the principle of dye-sensitized solar cells including research progress in the different cell components, an overview about the development of new electrolytes based on solvent and solvent-free formulation, and a description of the current understanding on the degradation mechanisms in dye-sensitized solar cells.

2. Dye-Sensitized Solar Cells Principle: Role and Progress in the Different Components

A schematic presentation of the operating principles of the DSC is given in Figure 1. At the heart of the electrochemical system, a mesoscopic layer of anatase TiO$_2$ is sintered together to provide efficient electronic transport. The layer is typically composed of well-dispersed 20–30 nm size particles of anatase TiO$_2$ sheltered by a scattering layer constituted of ca. 400 nm particles whose role is to scatter unabsorbed photons (Figure 2). Attached to the surface of the nanocrystalline film is a monolayer of the light sensitive dye. Photoexcitation of the latter induces charge separation, ultrafast femto/picosecond electron injection into the conduction band of the oxide, and hole capture by the reductive redox species composing the electrolyte. This regeneration reaction enables recovering the original oxidation state of the dye. For a sufficient concentration of reductive species, this reaction is fast enough to prevent the recapture of the conduction band electron by the oxidized dye and to hamper dye degradation. The role of the dye is then pivotal for the cell operation as it will govern light absorption ability of the device. The main family of redox couples used in DSC is
the two-electron system I$_3^−$/I$^−$. Subsequently for dye regeneration, tri-iodide diffuses towards the counter electrode and collects two electrons from the external circuit to form back iodide. The counter electrode is typically supported by electrocatalytic platinum nanoparticles reducing the charge transfer resistance for I$^−$/I bond breaking.

Alternatives to anatase TiO$_2$ nanoparticles have been investigated. The most evident approach was to explore the two other main polymorphs of TiO$_2$: brookite [22] and rutile, where the latter can enhance the scattering characteristics of the reflecting layer owing to its higher refractive index [23]. Alternatively, ZnO [24–43], SnO$_2$ [44–48], Nb$_2$O$_5$ [49–55], and In$_2$O$_3$ [56] were also proposed along three other ternary metal oxides SrTiO$_3$ [57–59] and BaSnO$_3$ perovskites [60] or the spinel ZnSn$_2$O$_4$ [61, 62]. So far, the anatase TiO$_2$ remains the leading contender photoanode as it affords the best performances of charge collection. The optimization of this charge collection efficiency has been the stake in research activities in the mesostructuration of the photoanode or by introducing additional point defects through aliovalent doping. The reduction of particles dimensionality offers faster electron transport when the photoanode is composed of vertically aligned nanowires or nanotubes [63–71]. The synthesis of TiO$_2$ beads leads to the most spectacular improvements resulting from the excellent particles interconnections, shortened mass transport pathways into the mesopores, and combination of light scattering ability and high surface area offering improved light confining properties of the photoanode [16, 72–74]. Remarkable efficiencies exceeding 11% (A.M. 1.5 G) were obtained on optimized beads in combination with high molar extinction coefficient C101 or C106 ruthenium dyes.

The strategy to introduce point defects relies on the high electronic sensitiveness to doping of the 3d$^n$ electronic configuration adopted by the Ti$^{4+}$. The literature in this topic for DSC extends rapidly, even though this approach was proscribed for a long time as it could lead to a harmful increase in the density of free carriers. However, close to thermodynamic synthetic methods are employed which maintain TiO$_2$ a n-type semiconductor regardless of the dopant. It also allows charge compensation mechanism to maintain a low concentration carriers [75]. We can list a series of hypervalent cations which have been successfully incorporated into the anatase crystal structure: W$^{6+}$ [76], Nb$^{5+}$ [77–80], and Ta$^{5+}$ [81], subvalent cations: Zn$^{2+}$ [82], Cr$^{3+}$ [83], Fe$^{2+}$ [84], Sc$^{3+}$ [85], Y$^{3+}$ [86], and Ga$^{3+}$ [87], or isovalent cations such as Sn$^{4+}$ [88] and Ce$^{4+}$ [89]. The common observation is that hypervalent doping affects the energy of acceptor trap states. A careful control of dopant concentration can therefore adjust the trap energetics and thus is favorable in some cases to accelerate the charge injection. By contrast, subvalent dopants influence neither the energy nor the distribution of traps (Figure 3). With the appropriate type of dopant and concentration, improvement of charge collection efficiency can be achieved, thus contributing to improve the power conversion efficiency, even though the efficiencies reported so far are still lying below 10%.

Efforts to replace iodine/iodide redox couple have been motivated to reduce the nearby 600 mV energy loss from improper energy alignment between the dye HOMO level and the thermodynamic redox potential of iodine/iodide. It is secondarily motivated by the strong corrosive character of iodine and its deep orange coloration penalizing the blue light conversion of the solar spectrum. A significant reduction of this loss-in potential can bring the DSC to pass the threshold of 20% power conversion efficiency [90]. Figure 4 gathers the actual redox molecules and hole transporting materials proposed so far, listed in energy scale. The closely related two electrons Br$_3^−$/Br$^−$ have been suggested despite their high corrosive character [91–93]. In combination with carbazole-based sensitizers, Li et al. achieved DSC cells going beyond the 1 V photovoltage threshold. The sulfide/polysulfide redox couple is typically used for quantum-dot solar cells [94] and the alternate disulfide/thiolate redox couple showed appealing features in combination with ruthenium and organic dyes [95–97]. Another one-electron redox system was investigated, namely, the cobalt (+III+/II) polypyrrolid complexes [98–103], copper [104], iron [105], or nickel [106]. Fast redox couple based on nitroxide was proposed by Zhang et al. [107].

The output voltage is limited in principle by the dye absorption bandgap value (1.5 eV < $E_g$ < 2.5 eV). The integration of cobalt complexes has been beneficial to boost the photovoltage of champion devices. It is interesting to comment that the braking efficiency record in DSC by means of photovoltage improvement does not differ from the other single junction technology for which new records were systematically reached from a photovoltage improvement.

The central role of the sensitizing dye molecule generates demands in terms of the triptych chemical, electrochemical, and photoelectrochemical properties. The dye should of course combine chemical and photoelectrochemical stability but should also possess a high molar extinction coefficient with panchromatic light absorption and adequate HOMO/LUMO energy alignments. The pioneering work at EPFL presented a series of ruthenium polypyrrolid complexes among the ubiquitous N719 and Z907 dyes which are benchmark sensitizers still today [108]. Molecular ligand engineering for ruthenium dyes was judiciously scrutinized by different groups to tune the spectral response of the sensitizers either by introducing a ligand with a low-lying $π^*$ molecular orbital and by destabilization of the metal $t_{2g}$ orbitals through the introduction of a strong donor unit (Figure 5). N719 and related sensitizers are typically anchored onto TiO$_2$ according to a bidentate mode [109]. It signifies that two of the four carboxylic acid functions remain unbounded, thus not participating efficiently in the photon-induced charge releasing into TiO$_2$. To address this issue, one approach lies in the replacement of one of the two dicarboxybipyridyl units by an ancillary ligand displaying a slight donating character at the excited state while including a long alkyl chain to improve the dye solubility, to provide a hydrophobic character for longer term stability [110, 111], and to reduce iodine access to the surface for hampering recombination processes. It was also found that enhancement of
the π-conjugation in the ancillary ligand not only exerts an energetic stabilization of the LUMO levels, thus red-shifting the dye absorption, but also favorably affects the molar extinction coefficient of the dye reaching more than 20000 M\(^{-1}\)⋅cm\(^{-1}\) [12, 15, 112–115]. The integration of this second generation of dyes paved the way to improved performances in terms of light harvesting and power conversion efficiency. Well above 11% power conversion efficiency was achieved and excellent stability data reported at 60°C/100 mW⋅cm\(^{-2}\) conditions. The extension of conjugation in the acceptor ligand by going from bipyridine, terpyridine (e.g., N749) to quaterpyridine (e.g., N1044) contributes to a broadening in the absorption of the dye affording visible light panchromatic response [116–118], unfortunately at the expense of the molar extinction coefficient.

The second option to tune the complex absorption towards the red consists in destabilizing the t\(_{2g}\) orbitals of the ruthenium by the introduction of a stronger donor unit than thiocyanate. Many attempts to replace the thiocyanate donor ligands have been made and motivated also by the fact that the monodentate NCS is believed to be the weakest part of the complex from a chemical point of view. For long time, these efforts had yielded limited success as the conversion efficiency remained well below few percent [119]. A paradigm arose from the YE series developed by Bessho et al. who achieved 10.1% power conversion efficiency under standard illumination conditions (A.M. 1.5 G) [120]. The lowest energy MLCT band is red-shifted by 25 nm and contains a new absorption band at 485 nm with a remarkable molar extinction coefficient. This new band is characterized by an electronic transition from HOMO, which has a sizable π-orbital contribution from the cyclometalated ligand, and the LUMO formed by a set of π* orbitals localized on the bipyridines ligands evidencing strong electronic coupling with acceptor states in TiO\(_2\). These features explain the high photocurrent in the range of 17 mA/cm\(^2\) and the photovoltage

**Figure 3:** Effect of doping TiO\(_2\) by Nb\(^{5+}\), Ga\(^{3+}\), and Y\(^{3+}\) on energy and distribution of trap states and on charge collection efficiency.
Figure 4: Scale of chemical potential for the main (a) redox complexes and (b) redox molecules used in dye-sensitized solar cells.
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C12H25
C8H17
C6H13
C6H13
C8H17
C8H17
C8H17... fast electron transfer. A very recent in-depth review dedicated to metal-free sensitizers can be found in [131].

[121]. The removal of NCS ligands altogether in tridentate cyclometalated ligands also provides an enhancement in the absorption ability of the complex. This was demonstrated in the series of TF-dyes compared to the thiocyanate-related N749 (so-called black dye) [122]. With TF-dyes, Chou et al. reached close to 10% efficiency and even stability. They exhibit significantly higher molar extinction coefficient than ruthenium dyes, lying in the range of 20000 to 400000 M\(^{-1}\) cm\(^{-1}\). Four main families have been reported: coumarins which approached \(\eta = 7.7\%\) [123], indoles, and natural pigments among which are anthocyanin, flavonoid, carotenoid, and chlorophyll [124].

D-\(\pi\)-A structures actually stand out from the others as they combine high power conversion efficiencies beyond 10% [125, 126] and stability standard passing even the accelerated ageing test of 85\(\degree\) C/dark (Y123 dye). One key unit in the design of D-\(\pi\)-A originated from the introduction of cyanoacrylic function as an acceptor group which enabled extending light absorption ability of the dye compared to the related coumarin D-\(\pi\)-A [127]. It provides around 25% more short-circuit current density and exhibits a molar extinction coefficient close to 100000 M\(^{-1}\) cm\(^{-1}\) at 552 nm. Sensitizers having an intense absorption band in the near-infrared region can be typically acquired through phthalocyanines which are reaching absorption maxima to 700 nm. Squaraines dyes can even get close to 800 nm [128, 129]. Despite these interesting absorption profiles, NIR dyes have typically showed unimpressive power conversion efficiencies due to either too strong aggregation and lack of directionality in the excited state for the phthalocyanines [130] or issues for fast electron injection in the case of the NIR squaraines. This is explained by a too strong energetic stabilization of the \(\pi^*\) orbital delocalized throughout the dye structure. For these reasons, this call for the development of new semiconductor having lower lying energy conduction band edge for endorsing fast electron transfer. A very recent in-depth review dedicated to metal-free sensitizers can be found in [131].

![Figure 5: Main families of different organic dyes developed for dye-sensitized solar cells.](image)
3. Liquid Electrolyte Development Based on Solvent and Solvent-Free Formulation for Stable Devices

Champion efficiencies reported above are systematically obtained in conjunction with an electrolyte based on a volatile solvent. Pure acetonitrile or acetonitrile/valeronitrile solvent mixtures are often preferred. Their intrinsic characteristics, mostly in terms of volatility, cannot guarantee enough stability to pass the accelerated IEC61646 protocol. The excessive vapor pressure exerted shortcomings to the cell sealing even at temperatures well below 60°C, besides additional chemical/electrochemical reactions in cell which may occur. This volatility issue was circumvented by replacing acetonitrile with a different class of lower volatile solvents. γ-Butyrolactone (GBL), propylene carbonate (PC), propionitrile (PN), sulfolane, butyronitrile (BN), or 3-methoxyproponitrile (MPN) had been proposed. Their integration led to substantial device lifetime prolonging. It seems that the last two actually show the best compromise between high efficiencies and stability. The sulfolane-based electrolyte is promising for extreme conditions even though its high viscosity impedes high photocurrent production owing to mass transport limitations [132].

The electrolyte is not solely composed of iodide and iodine. It also contains two main additives. The beneficial impact of these additives on the cell performance is undeniable. However, the exact role/action in the complete cell remains speculative in some extent. Table 1 gathers the benchmark composition of nitrile-based electrolyte which is used to obtain high efficiencies or alternatively good stability for low-volatile MPN or BN solvent. Note that solvent-free ionic liquids are composed of binary or ternary eutectics melts including different components and composition than the reported in Table 1. A review was dedicated to this topic by Zakeeruddin and Grätzel [133]. The two additives typically incorporated in electrolyte are guanidinium thiocyanate and a Brönsted base, namely, terbutylpyridine for high efficiency electrolyte or replaced by the N-butyl benzimidazole or the benzimidazole in some cases for stable electrolytes. The utilization of a Brönsted base affords the deprotonation of the sensitized TiO₂ surface. It contributes to up-shift the quasi-Fermi level in TiO₂ and therefore the open circuit voltage of the cell. N-Butyl benzimidazole in stable electrolyte composition is preferred over terbutylpyridine, regardless of whether it is in MPN or BN (Figure 6). The comparison of the two molecules suggests that they have similar strength for deprotonation since the distribution of the subconduction band energetic levels is not modified. The charge collection efficiency is similar or slightly better in the case of the terbutylpyridine for which a marginally higher electron lifetime is compensated by a lower electron transport (Figure 6). The guanidinium thiocyanate has a more subtle function. The literature often refers to the guanidinium cation going to the surface of TiO₂ between the dye molecules in order to explain the improvement of charge collection efficiency. However, it is also straightforward that this additive is beneficial for the cell stability, in particular to maintain a high fill factor.

An impedance study carried out on TCO-Pt/electrolyte/Pt-TCO symmetric cells, performed at the laboratory, has highlighted that the charge transfer resistance evolution becomes strongly dependent on whether the electrolyte contains or not this guanidinium thiocyanate additive. Indeed, whereas the benchmark electrolyte composition containing 0.1 M of guanidinium thiocyanate shows a slight decrease in $R_\text{ct}$ to ca. 1Ω·cm²², such evolution appears in complete contrast with an electrolyte composition free of guanidinium thiocyanate (Figure 7). Beside its aforementioned function, we concluded that the guanidinium thiocyanate molecule also plays the role of a protective agent by molecular self-assembling onto Pt to prevent its dissolution through soluble PtI₆²⁻ complexes. This is further supported by visual observation of the symmetric cell during ageing which tends to bleach at the specific area in contact with the electrolyte.

In terms of stability performances, 10% efficiency on lab cells (<1cm²) with PCE retention over 95% after 1000 hours under 60°C/100 mW/cm² light soaking experiments currently stands for the best values reported so far on lab devices. This was achieved by using either BN [134] or alternatively gelified MPN-based electrolyte [135]. Dyesol recently reported its HSS electrolyte passing even 95°C dark accelerating test for 1000 hours in combination with N719 dye with efficiencies in the range of 5% under standard illumination conditions (A.M. 1.5 G) [132].

The standard accelerated ageing tests protocol used for evaluating the stability of dye-sensitized solar cells is by default the protocol established for terrestrial thin film PV (IEC61646):

(i) 1000 hours at 85°C (±2°C) in dark under 85% (±5%) humidity;
(ii) 1000 hours at 60°C under light illumination (800–1000 W/m²);
(iii) 300 thermal cycles between −40°C and +85°C.

Stable performances were reported over 2000 hours at 60°C in dark maintaining 5.5% PCE [136], 2.5 years in outdoor conditions without specification about module efficiency [137], or 2280 hours at room temperature under 80 mW/cm² light soaking [138], or more recently Dyesol Ltd. achieved as long as 25600 hours at 55°C–60°C under continuous light soaking with preserving 4% efficiency [139]. Since the seminal publication by Graetzel, only very few articles have reported stability at higher temperatures, requiring, as it seems, more viscous liquids to alleviate sealing issues, as, for instance, gelified electrolyte or solvent-free ionic liquids [140–143]. Apart from the very few reports announcing this achievement, still no consensual publications are actually passing the challenging threshold of 1000 hours at 85°C in dark condition with efficiencies above 5% [132, 144, 145]. Nevertheless, this assessment emphasizes on the one hand the high credibility of DSC for larger scale applications, but on the other hand, it also point out the important gap to fill in between the champion efficiencies and the stable devices for which this gap closing will require more focus on the understanding of the ageing mechanisms leading to the cell failure, and, in a next step to design specifically more
Table 1: Composition and structure of main liquid electrolyte components.

<table>
<thead>
<tr>
<th>Name</th>
<th>Acetonitrile</th>
<th>3-Methoxypropionitrile</th>
<th>1,3-Dialkylimidazolium iodide</th>
<th>Iodine</th>
<th>Ter-butylpyridine</th>
<th>N-Butyl benzimidazole</th>
<th>Guanidinium thiocyanate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Structure</td>
<td>H₃C—C≡N</td>
<td>H₃C—O—CH₂—C≡N</td>
<td>R=N=N—R'</td>
<td>I</td>
<td>phenyl</td>
<td>N=CH₂—N=CH₂</td>
<td>H₂N—N=CH₂</td>
</tr>
<tr>
<td>High efficiency (concentration)</td>
<td>Solvent</td>
<td>—</td>
<td>1 M</td>
<td>0.03 M</td>
<td>0.5 M</td>
<td>—</td>
<td>0.1 M</td>
</tr>
<tr>
<td>Stable efficiency (concentration)</td>
<td>—</td>
<td>Solvent</td>
<td>1 M</td>
<td>0.15 M</td>
<td>—</td>
<td>0.5 M</td>
<td>0.1 M</td>
</tr>
</tbody>
</table>
Figure 6: Comparison of cell characteristics during ageing at 60°C/100 mW·cm$^{-2}$ using an electrolyte constituted of TBP or NBB (composition: 1 M DMII, 0.1 M GuNCS, 0.15 M I$_2$, and 0.5 M NBB/TBP) in (a) 3-MPN solvent and in (b) BN solvent. A comparison of the trap state distribution and energy and a comparison of electron lifetime and transport time as a function of charge density are also included.
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0.1
1
0 1 02 03 04 05 0
Ageing days
No GuNCS
BN
MPN
MPN
BN
103
102
101
0.1M...an example of withdrawn aged electrolyte analyzed by LC/MS after 500 hours' ageing at 60°C/100 mW·cm⁻² measured in TCO-Pt/electrolyte/Pt-TCO symmetric cell configuration comparing BN and 3-MPN-based electrolyte, with 0.1M GuNCS and without GuNCS.

robust and better performing molecules (i.e., ligand, solvent, additive(s) in electrolyte, etc.).

4. Current Understanding of Chemical/Photoelectrochemical Degradation Paths

Whereas the durability of single-crystalline and amorphous silicon modules is established to a lifetime attaining 20 years without question, the stability and ageing predictability of emerging PV technologies are still under close observation to predict at least 10 years' lifetime under working environment. Mastering the dynamics of charge transport and charge recombination that take place in dye-sensitized solar cells is a crucial issue for reaching high performances. Nevertheless the stability of each cell component taken separately and the understanding of all the chemical, electrochemical, and photochemical reactions interplay at the different material interfaces or components are a key to bring further development on DSC, not only to improve the device stability under severe ageing conditions but also to close the gap between champion and stable efficiencies. Such complete understanding of the ageing mechanisms will pave the way to the specific design of more robust components. Integrated in an advanced generation, these new materials will give impetus to DSC commercialization and broader integration into the PV panorama. Only a very few groups are actively working in this more fundamental but crucial domain. Beside the intrinsic sealing and permeability issues of the device which have been revealed to influence the device stability as will be discussed below, six distinct intrinsic features contributing to the cell degradation have been spotted in the literature (Figure 8).

(i) Likely one of the most preeminent degradation path stems from the well-known iodine consumption in electrolyte: this reaction takes place during ageing. It translates into a well-visible bleaching in coloration of the electrolyte. It is crucial to overcome this side reaction as it is one important source of failure since iodine concentration controls the kinetic for tri-iodide mass transport. Excessive depletion will turn to short-circuit current density limitation of the device.

(2) Dye desorption at elevated temperatures (>60°C): this possible event takes its origin from the rivalry between dye solubility in the electrolyte and the binding strength of the anchoring group upon TiO₂. It can also be promoted by the ruthenium hexacoordination rupture as it will be discussed below. Water intake in the electrolyte has also been proposed to assist this dye desorption mechanism. The development of hydrophobic dyes was anticipated to go against this reaction and explained the better long term stability of Z907 versus N719.

(3) There is lack of chemical and photochemical stability of the monodentate thiocyanate ligand in heteroleptic ruthenium (+II) complexes which tends to undergo substitution reactions with different external components.

(4) There is platinum dissolution for electrolytes free of thiocyanate (cf. Figure 7).

(5) Formation of a polymeric solid electrolyte interphase (SEI) forming on TiO₂ and sensitized-TiO₂ (see Figure 10 inset).

(6) UV irradiation causes direct bandgap excitation of TiO₂ leading to conceivable dye or electrolyte component oxidation [137, 146]. Addition of MgI₂ or CaI₂ in the electrolyte was proposed to improve the electrolyte tolerance to UV irradiation [135]. It is also typical to cover the glass photoanode by an antireflecting polymer coating which also plays the role of UV filters [146].

In the following we will review in more details the two most severe issues to overcome, namely, the iodine depletion in electrolyte and the dye chemical/photoelectrochemical stability.

4.1. Iodine Consumption. Iodine consumption has been spotted by numerous groups who gave different attributions to explain this depletion. A total of eight hypotheses/observations, sometimes controversial, can be listed.

(i) Formation of IO₃⁻ induced by water traces [147–150]: in this case it is postulated that I₂ reacts with ingress water or residual water contained in the electrolyte. It leads to the formation of the iodate anion (IO₃⁻). As commented also by some of these authors, the formation of IO₃⁻ has, however, never been really detected so far either by spectroscopic methods (UV-Vis, Raman, etc.) or by chromatography (LC/MS). Figure 9 reports an example of withdrawn aged electrolyte analyzed by LC/MS after 500 hours' ageing at...
Figure 8: Schematic gathering the known stability issues in dye-sensitized solar cells.

Figure 9: LC/MS chromatogram of aged MPN-based electrolyte in contact with TiO$_2$ during 500 hours at 85°C/dark. The chromatogram for $m/z$ value of 175 is reported for comparison.

85°C in dark and compared to the specific signal of $m/z$ value of IO$_3^-$ [151].

(ii) $I_2$ reacts with glass frit used in some technologies as a sealant [152]. The experimental evidences reported in the reference are conclusive. However, this depletion is experienced regardless of the type of sealant utilized (i.e., Bynel or Surlyn polymers, glass frit, etc.). This is therefore not the original path for this reaction even though it will contribute to this consumption.

(iii) Sublimation of molecular $I_2$ has been proposed by different groups [145, 153, 154]. This hypothesis has been controverted by ex situ and in situ spectroscopic methods such as Raman on aged electrolyte and GC/MS or TGA/MS performed also on aged electrolyte or on electrolyte heated at 60°C or 85°C with gas output analysis. The results were suggesting the absence of any spectroscopic band or mass which could be attributed to iodine or iodide-based compounds even in form of traces [151].

(iv) Bandgap excitation of TiO$_2$ leads to the photoelectrochemical reduction of tri-iodide [155]. This reaction would be particularly unexpected knowing that TiO$_2$ has a strong n-type semiconducting character with direct bandgap excitation. This reaction takes place faster without than with dye and therefore could not be attributed to a recombination procedure between iodine and electron in conduction band. Last but not least [152], also pointed out that iodine consumption proceeds in dark and is only thermally activated as discussed below.
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Figure 10: Evolution of UV-Vis absorption spectrum for reference MPN-based electrolyte (in black), electrolyte aged alone during 500 hours at 85°C/dark (in blue), and electrolyte aged in contact with TiO₂ (in red). In inset are reported high resolution transmission electron micrographs comparing the TiO₂ nanoparticles before and after ageing showing the formation of SEI layer.

(v) The electrolyte bleaching is solvent dependent. This has been clearly highlighted from the work led by Dyesol by Jiang et al. [132, 156]. The authors compared γ-butyrolactone, 3-methoxypropionitrile, and the so-called nitrile-free solvent HSS. They have concluded that iodine depletion is significantly hampered when going from γ-butyrolactone to HSS. They have also reported slower iodine consumption using tetruglyme compared to 3-MPN [156].

(vi) Iodine consumption is not an intrinsic reaction taking place in the electrolyte alone. It is triggered by the surface of TiO₂ which plays a catalytic role in the electrolyte degradation [151]. The authors concluded that iodine depletion is exclusively activated by temperature while light has no action on this reaction, at least for 3-MPN-based electrolyte (Figure 10).

(vii) Iodine reacts with the electrolyte additives such as the 4-tert-butylpyrididine (TBP) leading to an idopyridinate complex and with thiocyanate ligand of the dye to form I⁻NCS⁻ species in the particular case the electrolyte is exempt of any TBP [157, 158].

(viii) Formation of a solid electrolyte interphase (SEI) nucleating on the surface of TiO₂ in the dye monolayer pinholes (Figure 10 inset) [151]: the authors highlighted that SEI formation traps or solvates iodine (and other electrolyte components) explaining its depletion from bulk electrolyte. It is supported by UV-Vis and XPS spectroscopies combined to ToF-SIMS which revealed on the one hand its very complex composition and on the other hand its high concentration of iodine/iodide-based species along other degraded components of the electrolyte. To the current understanding, this SEI layer is originating from the polymerization between acrylonitrile radicals leading to the formation of a very cohesive polyacrylonitrile polymer. GC/MS experiments showed that in presence of TiO₂ the 3-MPN thermally disrupts to form two highly volatile compounds: acrylonitrile (bp. = 77°C) and methanol (bp. = 65°C) [159].

The free radical polymerization of acrylonitrile is well-established to take place spontaneously by mild thermal activation such as temperatures in a range of 50°C. The polymerization rate can get substantially faster in the presence of metal halide catalyst [160]. The formation of these two degradation compounds can also give a more rational explanation to the sudden electrolyte evaporation pointed out by many authors in the field, which operates at a more or less long term in the ageing, an event typically the culprit of poor cell sealing.

The literature often refers to iodine consumption in electrolyte as it is the chromatic component of the electrolyte with its characteristic strong purple color. At HOPV2014 conference our group presented that many other electrolyte components are in reality consumed. We determined by cyclic voltamperometry using a platinum microelectrode that about 50% of iodide is depleted after 500 hours’ ageing at 85°C (Figure II(a)), FT-IR carried out on aged electrolyte indicates that thiocyanate from guanidinium thiocyanate is almost completely depleted (Figure II(b)), and last but not least N-butyl benzimidazole is also quantitatively consumed ca. 50%) by LC/MS (Figure II(c)) [159]. These experiments were realized looking at the interface between naked TiO₂ and electrolyte. When sensitized, these reactions are still occurring but at slower rate. This was similarly experienced in the case of iodine consumption for which Asghar et al. have made careful comparison between unsensitized TiO₂ and sensitized TiO₂, highlighting that dye monolayer only slows down this reaction [161]. MPN-based electrolyte stability is strongly affected by the surface of TiO₂ which tends to destabilize significantly the thermal stability of electrolyte components by about 50°C (Figure 12). The surface of TiO₂ also collaterally induces gas formation as aforementioned in this review [151, 159].

4.2. Chemical/Photochemical Stability of Benchmark Ruthenium Sensitizers. This part is more focused on ruthenium polypyridyl complexes. Much less is known about the core stability of organic dyes. Tanaka et al. reported that major degradation in indoline-based D-π-A dye, namely, the yellow DI31 chromophore, was decarboxylation of the cyanoacrylic acid anchoring group [162]. This decarboxylation reaction is contributing to dye desorption, lowering significantly the device performance whether the ageing is at 60°C/1sun or 85°C/dark. This reaction has been attributed to harmful action of iodine or amine components to the dye. This reaction can likely be extended to some degree to the family of “push-pull” D-π-A dye, even though some of other dyes in this family have been reported to be very stable upon 60°C/light and 85°C/dark ageing conditions. Most of organic dyes are known to cause stability issues owing to their empathy to undergo fast photooxidation when they are
Figure 11: (a) Comparison of cyclic voltamperometry recorded on Pt microelectrode for MPN-based electrolyte aged in contact with TiO$_2$ during 100, 200, and 500 hours at 85°C/dark. (b) ATR-FT-IR spectra comparison of MPN-based reference electrolyte before and after 1000 hours ageing at 85°C/dark in contact with TiO$_2$. (c) Quantification by LC/MS of NBB concentration as a function of ageing time at 85°C/dark in contact with TiO$_2$.

anchored onto TiO$_2$. Following this hypothesis, increased stability will thus require a fast recovery of the reduced form to prevent this side reaction to occur.

The chemical/photo-(electro-)chemical durability of ruthenium polypyridyl-based sensitizers in contact with TiO$_2$ had been questioned well before the seminal paper from Anderson et al. in 1979 [163]. The underlying effect of the nanostructuration of TiO$_2$ on the ruthenium-complex stability was discussed [4]. The authors presented photocurrent stability over 2 months subjected to continuous visible light illumination stress with less than 10% degradation corresponding to stability over $5 \times 10^6$ turnovers. Similar arguments were reported on monomeric dye without noticeable degradation [164].

These intriguing results are in relative contradiction with other reports on cis-Ru(bpy)$_2$(SCN)$_2$ (bpy = 2,2’ bipyridine) but also on trimeric ruthenium dyes [165]. The lack of structural robustness of N719 and Z907 had been reported in particular by Tributsch, Hagfeldt, and Lund. These authors became consensually alarmed on the vulnerability of the strong electron donor thiocyanate ligand which can easily exchange with electrolyte components as a result of the antibonding character of these orbitals. This ligand exchange reaction is triggered not only by light action (photolysis) but also can be activated by temperatures above 80°C. The monodentate thiocyanate ligand can then exchange with acetonitrile or 3-methoxypropionitrile when used as a solvent (in integral or fragmented part such as C≡N), with 4-tert-butylpyridine, water residues, or even iodide [165–173]. The different exchange reactions reported are summarized in Figure 14. This reactivity of the monodentate thiocyanate ligand was explained by Grünwald and Tributsch on the basis of an incomplete reduction of the photooxidized dye by I$_3$”/I$^-$ redox couple in combination with a certain lack of stability.
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... oxide (ITO) layer can irreversibly lose its conductivity when voltages above 1.5 V are applied, need to remind that

of the oxidized form of the complex [165]. This explanation was further confirmed by Kohle et al. who showed in solution that dye instability can also get prompted by a too low concentration of iodide, in other words when decreasing the rate constant for the dye regeneration [174]. The results provided by Lund et al. suggest again that TiO₂ plays a significant catalytic activity towards this ligand exchange reaction which is accelerated by a factor of 2.

It is also suggested that some degradation products of N719 and Z907 dyes are in equilibrium together [171] (Figure 13). It was for instance evoked in three interrelated equilibrium reactions:

\[
[\text{RuLL}^+(\text{NCS})_2]^+ + 3 \text{MPN} \rightleftharpoons [\text{RuLL}^+(\text{NCS})(3\text{-MPN})]^+ + \text{NCS}^- \quad (1)
\]

\[
[\text{RuLL}^+(\text{NCS})(3\text{-MPN})]^+ + 4 \text{TBP} \rightleftharpoons [\text{RuLL}^+(\text{NCS})(4\text{-TBP})]^+ + 3 \text{MPN} \quad (2)
\]

\[
[\text{RuLL}^+(\text{NCS})_2]^+ + 4 \text{TBP} \rightleftharpoons [\text{RuLL}^+(\text{NCS})(4\text{-TBP})]^+ + \text{NCS}^- \quad (3)
\]

The rate for this ligand exchange reaction can be lowered by a factor of 2 when including a buffer concentration of thiocyanate anions in the electrolyte (e.g., guanidinium thiocyanate) [169, 175]. The degradation products based on 4-TBP can be prevented by replacing this latter with N-butyl benzimidazole or the closely related benzimidazole which enhances the device stability. This can explain in part or in whole the better stability achieved with NBB (Figure 6(a)). Although such ligand exchange reactions will affect the optical MLCT (metal-to-ligand charge transfer) contribution, entailing either slight bathochromic or hypsochromic shift of absorption [132], the real implication of this in situ/in operando dye structure modification on the practical cell characteristics remains unclear. Finally, again in the work published by Grünwald and Tributsch, the authors exposed the possibility that the ruthenium hexacoordination can get fragmented, leading to the desorption of the metal core unit and the retaining of the anchoring bipyridine group attached upon TiO₂ [165].

Besides these intrinsic degradation mechanisms highlighted to date, more external contributions to the cell degradation have also been exposed, namely, sealing conditions, substrate corrosion, and sealing issues. Assessment in these technical aspects is relatively complex to detail since they are rarely published. An interesting work reported by Fredin et al. aims at alarming the importance to consider the temperature of sealing as it will determine the device efficiency [176]. This work was examined by comparing N719 with the organic D5 sensitizer. The authors found that N719 appears more sensitive to thermal degradation than electrodes sensitized with the organic D5. Under temperatures lying between 120°C and 250°C for 5 minutes, the cell efficiency can vary from a maximum of ca. 4% to almost nothing when cells were sealed with an excessive sealing temperature (> 200°C). By using IMPS/IMVS spectroscopies, they also highlighted that the electron diffusion length, referring to the charge collection efficiency, can get as low as 10%.

The encapsulation of the cells should prevent the exchange of material between the inner part of the cell and the ambient environment. It is crucial for the cell stability that the encapsulation creates a barrier against water and oxygen ingress which can be assimilated by the electrolyte and the mesoporous sensitized TiO₂ layer. Conversely water has also been reported to enhance the power conversion efficiency performances of DSC based on ionic liquids [177]. The proportion of water in the cell can drastically modify the proton concentration in the electrolyte. Note that proton is a potential determining cation; in other words it tends to form specific adsorption on the surface of TiO₂ leading to downward shift in energy of the Fermi level and conduction band edge [178]. A critical threshold value of water concentration to maintain high stability and improve the cell efficiency is difficult to evaluate although it is an important factor governing the device stability. One difficulty to assess this threshold value results from the inappropriateness of DSC’s electrolyte towards Karl-Fischer titration because of iodine. On the other hand, this threshold value is expected to strongly depend not only on the type of dye used but also on the type of electrolyte.

Finally, the general photocatalytic properties of TiO₂ should be mentioned since it could trigger the formation of hydrogen peroxide from oxygen and water, which itself can destructively oxidize organic compounds [179, 180]. The production of H₂O₂ is initiated by the reduction of O₂ from conduction band electrons in TiO₂ leading to superoxide anion O₂⁻ which in turn will react with water. It should also be stressed that because of the instability in nature of the superoxide anion radical, it can also directly participate in internal chemical reactions to other organic cell components [181].

Related to the sealing of the cell is the conductive substrate (transparent conducting oxide, TCO) itself which in some specific conditions can sustain a number of events leading to its conduction loss properties. The indium-doped tin oxide (ITO) layer can irreversibly lose its conductivity when voltages above 1.5 V are applied, needless to remind that
such voltages can be reached in series modules [182]. This may bring issues, for instance, for flexible devices for which the substrate is typically made of ITO deposited on polyethylene naphthalate (PEN) or polyethylene terephthalate (PET).

5. Conclusive Remarks

After being strongly questioned subsequently to the seminal publication of Graetzel et al. describing nanocrystalline dye-sensitized solar cells, the macroscopic stability of DSCs is now reported in hundreds of publications. To date IEC61646 accelerated protocol is passed by a few companies although the power conversion efficiency performances are still lagging well behind the champion numbers reported by the public research laboratories. The development of stable DSCs was principally relying on beliefs and excellent intuitions from the community gathered after more than 20 years of research. We can only regret that yet very little is known about the interrelated chemical reactions responsible for the device ageing and the stress factors responsible for this degradation. Which is the most critical parameter: UV, visible light, or temperature? The experiments gathered in our group on electrolyte stability pointed out the temperature to be the most critical parameter.

In practice, stability investigation papers mainly report the evolution of the cell characteristics as a function of time. Less often, IMVS-IMPS, EIS, and transient photovoltage/photocurrent decay were techniques used to monitor the influence of ageing on the charge transfer kinetics and distribution and on the energy of the surface trap states in the nanocrystalline TiO$_2$. For further improvement, not only for stable power conversion efficiency but also for prolonging significantly the lifetime of DSC to get competitive with silicon technology, there is an urgent need to establish a set of experiments and to develop techniques for careful analyses and characterization of the degradation products which are formed upon device ageing. In this area, only a very few groups are actively contributing to puzzling the complexity of dye-sensitized solar cells.

In this review, we spotted that TiO$_2$ plays a major role in the device stability. This goes particularly against the preconceived idea that the inorganic part of the cell is the most robust compared to organic and organometallic compounds. In fact TiO$_2$ displays a catalytic role not only for the dye degradation as it had been clearly demonstrated by the pioneering work of Lund’s group but also for the electrolyte stability, in particular those based on alkyl and alkoxy nitrite solvents which we are currently scrutinizing. Understanding the multifaceted and interplay chemistry of dye-sensitized solar cells and developing new materials based on such knowledge will indubitably open up significant stability breakthrough in this mystic artificial photosynthetic solar cell (Figure 14).
Conflict of Interests

The author declares that there is no conflict of interests regarding the publication of this paper.

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