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Usage of Ionic Liquid Electrolyte in Tin and Zinc Oxide Composite Dye-sensitized Solar Cells

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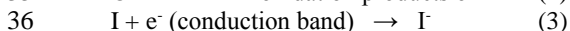
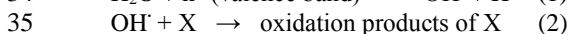
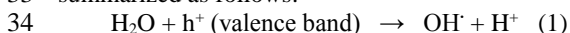
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1 Though Dye-sensitized solar cells (DSCs) made from
2 less photo-catalytically active tin and zinc oxides are
3 inefficient, composite tin-zinc oxide films yield efficiencies
4 comparable to those fabricated out of titanium dioxide films.
5 Studies conducted reveal that DSCs based tin-zinc oxide
6 films are highly stable when high boiling point solvents are
7 incorporated to solubilize the conventional iodide/tri-iodide
8 redox system.

9 **Keywords:** Dye-sensitized solar cell, Tin oxide, Zinc
10 oxide
11

12 Researchers worldwide have engaged in development
13 of dye-sensitized solar cells (DSCs) for nearly three decades
14 [1–7]. Despite this effort, the device continues to be plagued
15 with issues of long term stability preventing large scale
16 practical application. Instability originates from photo-
17 degradation of the dye and/or the electrolyte, evaporation or
18 leakage of the electrolyte owing to imperfect sealing [8,9].
19 The latter cause is largely eliminated by the use of
20 electrolytes based on high boiling point ionic liquids instead
21 of volatiles such as acetonitrile [10]. The photo-degradation
22 of the dye and the electrolyte happens mainly due to
23 photocatalytic reactions at the TiO₂ surface, induced by
24 ultraviolet (UV) component of solar radiation in presence of
25 water. The moisture contamination of the electrolyte is
26 practically unavoidable, even with the use of hydrophobic
27 electrolytes [11]. Holes generated via direct absorption of UV
28 photons by TiO₂, produce highly oxidative hydroxyl free
29 radicals via reaction with water molecules [12]. The
30 concomitant electron, reduces iodine discoloring the
31 electrolyte, an effect almost always seen in prolonged
32 illumination of a DSC. The reactions involved can be
33 summarized as follows:



37 Where X denote the dye or the electrolyte.

38 The above problem could be solved by use of a less
39 photo-catalytically active larger band-gap (compared to ~ 3.2
40 eV for TiO₂) oxide semiconductor having appropriate
41 positioning of the conduction band edge [2,5]. Tin (IV) oxide
42 (band gap ~ 3.8 eV) satisfies the above requirement, however,
43 solar cells based on this material are inefficient and energy
44 conversion efficiency values attained are order of 1% or less.
45 The reason attributed is faster recombination of dye cation
46 with conduction band electrons, probably due to the low
47 effective mass of conduction band electrons in SnO₂.
48 Imposition of ultra-thin barriers of higher band gap oxides

49 over SnO₂ crystallites, relieve this problem greatly,
50 suppressing recombination and efficiencies comparable to
51 that of the TiO₂ system has been achieved with SnO₂ based
52 DSCs, when the crystallite are coated ZnO or MgO [5]. This
53 work examines the stability of a DSC made of SnO₂/ZnO
54 (ZnO coated SnO₂) using an ionic liquid based electrolyte
55 (0.4 M I₂ in 1-methyl-3-propylimidazolium iodide with ~ 1 %
56 by vol 4-tert butyl pyridine) and compares it with a TiO₂
57 system based on the same electrolyte.

58 Before making the samples, fluorine-doped tin oxide
59 (FTO) plates (1 cm × 2 cm) were cleaned by ultra-sonication
60 in a detergent for 5 min and thoroughly washed with distilled
61 water. Then the cleaned glass plates were rinsed with
62 isopropyl alcohol and purged with air to dry. Commercially
63 available chemicals of purity 98% or more were used for
64 sample preparation without further purifications. Fabrication
65 method of each film and the construction of solar cell are
66 described below.

67 First, SnO₂ films were fabricated as follows. Colloidal
68 tin oxide (3 ml, 15% aqueous dispersion Alfa Aesar) and
69 glacial acetic acid (6 drops) were grounded in a mortar. Triton
70 X-100 (5 drops) and ethanol (20 ml) was added to the mixture
71 and the suspension sonicated for 15 minutes were sprayed
72 onto cleaned FTO substrates heated to 150 °C. Finally, SnO₂
73 films were sintered at 500 °C for 30 minutes.

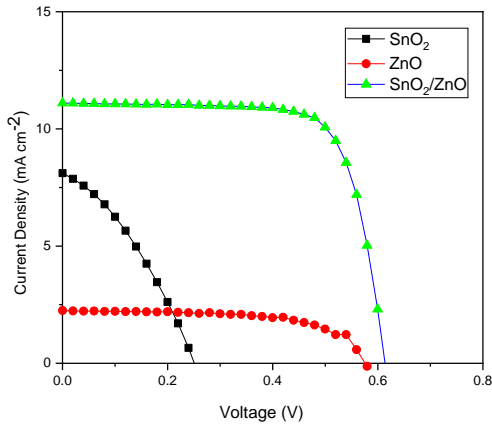
74 Zinc oxide powder (0.06 g Aldrich), glacial acetic acid
75 (6 drops) and Triton X-100 (5 drops) were mixed with ethanol
76 (20 ml) for the fabrication of ZnO films. The suspension was
77 sonicated for 15 minutes and sprayed onto cleaned FTO
78 substrates heated to 150 °C. ZnO films were sintered at
79 500 °C for 30 minutes.

80 SnO₂/ZnO composite films were fabricated grounding
81 colloidal tin oxide (3 ml, 15% aqueous dispersion Alfa Aesar),
82 glacial acetic acid (6 drops), ZnO (0.06 g Aldrich) in a mortar
83 and then adding triton X-100 (5 drops), ethanol (20 ml) to the
84 mixture and the suspension sonicated for 15 minutes was
85 sprayed onto cleaned FTO substrates heated to 150°C. Lastly,
86 composite films were sintered at 500 °C for 30 minutes.

87 Plates coated with oxides (SnO₂, ZnO and SnO₂/ZnO)
88 were heated to 80 °C and kept soaked overnight in a 3x10⁻⁴
89 M solution of N719 dye in 1:1 v/v ratio acetonitrile and tert-
90 butyl alcohol. The Pt counter electrode was clipped firmly to
91 the dye-anchored surface and the space between the two
92 electrodes was impregnated with the ionic liquid based
93 electrolyte (0.4 M I₂ in 1-methyl-3-propylimidazolium iodide
94 with ~ 1 % vol 4-tert butyl pyridine). A mask with a window
95 of 0.25 cm² was pasted the working electrode and solar cell
96 characteristics were ascertained using a simulated sunlight

1 using SPD SS-25 LED Solar Simulator and VK-PA-300K PV
2 Power Analyzer (AM 1.5 at 100 mW cm^{-2}).

3 The IV characteristics of DSCs made of SnO_2 , ZnO and
4 SnO_2/ZnO films are shown in Figure 1 and the IV parameters,
5 short-circuit photo-current (J_{sc}), open-circuit voltage (V_{oc}),
6 fill factor (FF) and efficiency (η) are summarized in Table 1.
7 The efficiencies of cells made of SnO_2 and ZnO separately
8 are 0.69% and 0.83% respectively, whereas that of SnO_2/ZnO
9 is 5.04%. Clearly conspicuous improvements in J_{sc} , V_{oc} and
10 FF are also seen in the SnO_2/ZnO based solar cell.

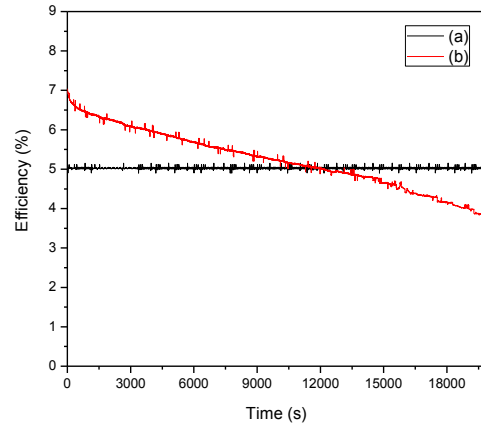


11
12 **Figure 1.** IV characteristics of the DSCs based on SnO_2 , ZnO and
13 SnO_2/ZnO films using the ionic liquid electrolyte constituted of
14 0.4 M I_2 in 1-methyl-3-propylimidazolium iodide with $\sim 1\%$ by
15 vol 4-tert butyl pyridine.

16 **Table 1.** IV parameters of DSCs based on SnO_2 , ZnO and
17 SnO_2/ZnO films using the ionic electrolyte constituted of 0.4 M I_2
18 in 1-methyl-3-propylimidazolium iodide with $\sim 1\%$ by vol 4-tert
19 butyl pyridine.

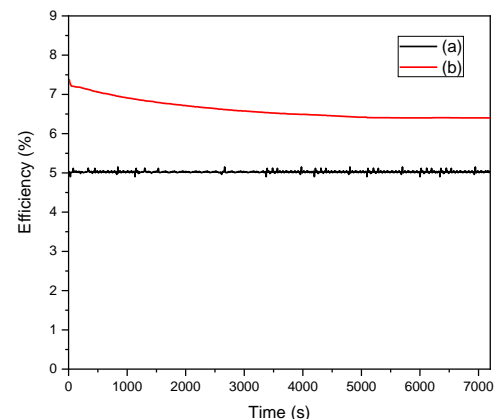
Working Electrode	J_{sc} (mA cm^{-2})	V_{oc} (V)	FF (%)	η (%)
SnO_2	8.11	0.251	0.34	0.69
ZnO	2.25	0.577	0.64	0.83
SnO_2/ZnO	11.10	0.614	0.74	5.04

20
21 Figure 2 compares the time variation of the efficiencies of
22 SnO_2/ZnO cell that utilizes the high boiling point ionic liquid
23 and volatile electrolytes. The less viscous volatile electrolyte
24 yields higher initial efficiency, but the electrolyte loss causes
25 rapid decline of the efficiency. A detectable decrease in the
26 efficiency of ionic liquid system was not observed during the
27 period of illumination.
28
29



30
31 **Figure 2.** Comparison of the time variation of the efficiencies of
32 SnO_2/ZnO DSCs based (a) on ionic liquid electrolyte (0.4 M I_2 in
33 1-methyl-3-propylimidazolium iodide with $\sim 1\%$ by vol 4-tert
34 butyl pyridine) and (b) volatile electrolyte ($0.4 \text{ M LiI}/10^{-2} \text{ I}_2$ in
35 methoxide acetonitrile containing $\sim 1\%$ by vol 4-tert butyl
36 pyridine).

37 When SnO_2/ZnO and TiO_2 based cells with ionic
38 electrolyte were compared, TiO_2 system displayed clear
39 evidence for gradual decrease of efficiency (Figure 3). It is
40 also evident that rate of decrease of the efficiency of the TiO_2
41 cell is initially faster and the rate gradually decrease, possibly
42 because, the degradation reactions (1)- (3) eliminate moisture.
43 Figure 3 compares the time variation of the efficiencies of
44 SnO_2/ZnO and TiO_2 cells using the ionic liquid electrolyte
45 (0.4 M I_2 in 1-methyl-3-propylimidazolium iodide with $\sim 1\%$
46 by vol 4-tert butyl pyridine). TiO_2 cell with ionic electrolyte
47 yields higher efficiency, but the efficiency reduces by about
48 10% over 2 hour period. However in the SnO_2/ZnO system
49 detectable decline in efficiency was not observed. A 10%
50 reduction in efficiency of TiO_2 based solar cell could be
51 attributed to photocatalytic effect of TiO_2 induced by
52 ultraviolet component of the solar simulator radiation.
53



54
55 **Figure 3.** Comparison of the time variation of the efficiencies of the (a)

1 SnO₂/ZnO and (b) TiO₂ based DSCs using the ionic liquid electrolyte
2 (0.4 M I₂ in 1-methyl-3-propylimidazolium iodide with ~ 1 % by vol 4-
3 tert butyl pyridine).

4 The work reported above has conducted a comparative
5 study of DSCs made of SnO₂, ZnO, SnO₂/ZnO and TiO₂ films
6 sensitized with the N719 dye, using an I⁻/I₃⁻ redox shuttle in
7 a high boiling point ionic liquid electrolyte. Results indicate
8 that the SnO₂/ZnO cell, yields nearly an order of magnitude
9 higher efficiency compared to the efficiencies of systems
10 based on individual oxides. More importantly findings
11 demonstrate that the DSCs made of SnO₂/ZnO films and the
12 ionic liquid electrolyte are highly stable and resistant to
13 degradation during prolonged illumination.

14
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18 References and Notes

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