Application of Polyvinyl Acetate (PVAc) in Dye Sensitized Solar Cells (DSSC) as Electrolyte in The Presence of Acetonitrile as Highly Soluble Organic Solvent and KI as Inorganic Redox Salt

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Abstract—The dye sensitized solar cells technology is one of the highly efficient, low cost and easily fabricated renewable energy technology. The highly soluble PVAc was prepared by solution polymerization. The PVAc was employed as an electrolyte in third generation, Dye sensitized solar cell (DSSC). The PVAc was characterized using SEM, XRD and conductivity measuring techniques. The prepared electrolyte behaved as gel electrolyte. SEM images shows the excellent dispersion PVAc in liquid electrolyte. The conversion efficiency achieved using gel electrolyte was 4.18%, which was comparable to liquid electrolyte with a value of 4.57%. This new technique reduces the degradation of DSCs rising from volatilization and leakage of the liquid electrolyte and endorses the commercialization process of DSSC.

Keywords—Polyvinyl acetate, Acetonitrile, Gel electrolyte, Dye sensitized solar cell.

I. INTRODUCTION

Solar cells are the renewable energy resources that pick up their energy from sun and supply this endless energy for human beings. It is one of the most hopeful energy resources due to its profusion, purity, safety and higher economic values that causes energy creation in distant rural areas [1]. The Solar energy can be used for three purposes: Photovoltaic energy i.e. Solar PV, solar thermal energy and passive solar energy [2]. Technology of producing electricity directly from the solar energy was discovered by Alexander Edmond Becquerel, who was a French physicist at 1839 and that was the start of solar energy [3]. A solar cell is an electric device that directly convert incident light into electricity by photovoltaic effect [4].

The solar cell technology is grouped into three different generations, as shown in Fig.1. Illustration of solar cell technology, it is grouped into first, second, and third generations and These generations are additionally arranged into various kinds [5].

In 1991 the seminal work was initiated on DSSCs by O’Regan and Grätzel [6]. The DSSC is an emerging technology which is one of the third generation PV technology.

An extensive research has been made in this technology on its applications during the last two decades because of its cheaper and easier fabrication process [7]. This technology mimics the natural photosynthesis in the way of light absorption [8]. DSSCs work better than other PV devices in dark condition that is in cloudy climate or in sunset and in diffused light radiations which makes it outstanding choice for indoor applications like sunroof and windows [9]. So far, the efficiency achieved by DSSC is from ∼7% to ∼14% [10]. A DSSC basically comprises a semiconductor (e.g. ZnO, SnO2 and TiO2 etc.) of few microns thick film as working electrode, a dye i.e. Organic dye or inorganic metal complex dye, a counter electrode (e.g. carbon materials and Pt etc.) also coated on conductive glass substrate and an electrolyte (i.e. Co2+/Co3+ and I3−/I− etc. as a redox couples) inserted between counter electrode and dye, as shown in Fig.2 [11]. The light is absorbed
by the dye molecules and electrons are injected into the conduction band of semiconductor. The electrolyte in contact with dye, provide electrons to the oxidized dye and restores it to its initial state. Then the oxidized electrolyte moves toward counter electrode and reduced to its neutral state [13].

The organic solvents have the volatility problem of its solvent caused by thermal stress and as a result cell deterioration is caused over long use. Consequently, ionic liquid solvents have been the promising electrolyte because of its good thermal and chemical stability, tuneable viscosity, high ionic conductivity and low volatility. Researchers are in interest to discover new liquid electrolyte, to increase the efficiency of DSSCs, such as 1,3-dialkylimidazolium etc. [14,15]. Low-viscosity ionic liquid is mixed with imidazolium iodide, to higher the efficiency [15].

Polymers has a unique network structure in gel electrolyte and shows, long term stability, higher electrical conductivity and good interfacial contact as compare to liquid electrolyte [16]. Polyvinyl acetate was discovered by Germany scientist Fritz Klatte in 1912. The manufactured polymer has the formula (C4H6O2)n and the general formula with the polyvinyl esters family is - [RCOOCHCH2]. It is a kind of thermoplastic materials and its IUPAC name is poly (1-acetylxiethene). Polyvinyl acetate is profoundly soluble polymer in organic solvents [17, 18].

This study, therefore demonstrate the application of polymer in DSSC’s electrolyte in the presence of acetonitrile as solvent and KI (potassium iodide) as redox ion. The prepared Polymer was characterized through FTIR, XRD and the IV characteristics was done by solar simulator.

II. EXPERIMENTAL

A. Materials:

Vinyl acetate, sodium dodecyl sulphate, acetone and ethanol were purchased fromuni-chem Lab Ltd (India) whereas KI, iodine and acetonitrile from Schalab SL (Spain). The FTO glasses and N3 dye was obtained from Solaronics (Switzerland). All the chemicals purchased were in pure form.

B. Synthesis of PVAc

The Polyvinyl acetate was synthesized with the help of free radical solution polymerization technique. Simply, 30 mL of monomer vinyl acetate was taken in flask and in a certain amount of water and 5 mL of methanol and 0.1 G of ammonium persulfate (NH4)2S2O8 as an initiator were added. The reaction was refluxed at 70 °C for 2 hours. Once the mixture turned gelatinous 40 mL of methanol was mixed to the system and then stopped heating. The prepared polymer was dried in vacuum chamber for 24 hours at 40 °C. Then it is characterized for further applications.

C. Electrolyte preparation:

The prepared liquid electrolyte consisted of 0.4 M 1-methyl-3-propyl imidazolium iodide, 0.2 M KI, 0.3 M pyridine in acetonitrile as solvent. A 0.4 g of the PVAc was mixed with 1.5 g of liquid electrolyte and then mixer was stirred till complete dissolution of the polymer and then the electrolyte converted into quasi solid electrolyte and then employed both the electrolytes in DSSC separately. The prepared gel and Liquid electrolytes and DSSC module are shown in Fig.3.

III. RESULTS AND DISCUSSION

A. SEM Micrographs:

The SEM micrograph of PVAc is shown in fig 4. The fig 4. (a) shows that PVAc has uniform, non-porous and amorphous structure while fig 4. (b) shows the scanning electron micrograph of the gel electrolyte on the TiO2 working electrode.
As shown in the figure, the TiO2 surfaces are completely covered by the gel electrolyte. The big holes show the spongy network of PVAc through which the liquid electrolyte can pass through it and retains the volatility of the organic solvent by making bond with acetonitrile as a result stability of DSSC is improved.

B. XRD Analysis:
As shown in Fig 5, native PVAc is amorphous as expected. The peaks at 15°, 22° represent PVAc. As the figure shows the almost zero peaks which shows that it has non-crystalline structure [20].

C. Electrical Conductivity:
The fig 6 showed the conductivity of 2.3×10^-3 S/cm at high frequency of PVAc. The lower conductivity of the polymer is because of low level protonation of PVAc chain. The increase of the polymer conductivity with frequency is the radiation which cause splitting of the polymer chain. The conductivity is based upon electronic jumping which cause high energy free electrons, ions [19-22]. The fig 7 shows the conductivity of the gel electrolyte. This increase of the electrolyte conductivity is due to the free iodide ions present in the liquid electrolyte.

D. Performance Parameters:
The photovoltaic performance was done under illumination (AM 1.5G, 100 W cm⁻²) on DSSC employing gel electrolyte as shown in fig 7. The Voc, Jsc and FF of DSSC based on liquid and gel electrolyte are 0.712 V, 12.78 mA cm⁻² and 57 %, respectively, yielding conversion efficiency of 4.18%. The values (Voc, Jsc, FF, and η) for the liquid electrolyte are 0.686 V, 13.88 mA cm⁻², 59%, and 4.57%, respectively. By comparing the data of the two electrolytes show that the increase of the Voc of liquid electrolyte is compensated by the slight decrease of FF of the gel electrolyte.

CONCLUSION
In summary, the highly soluble PVAc was prepared by solution polymerization. The PVAc was employed as an electrolyte in DSSC. The PVAc was characterized using SEM, XRD and conductivity measuring techniques. The prepared electrolyte behaved as gel electrolyte. SEM images shows the excellent dispersion PVAc in liquid electrolyte. The conversion efficiency achieved using gel electrolyte was 4.18%, which was comparable to liquid electrolyte with a value of 4.57%. the gel electrolyte has the advantage of high efficiency, cheap, easy fabrication, environmentally friendly and stable. As a result, the stability and efficiency of DSSC can be improved by
improving the conductivity of the gel electrolyte in case of polymer nanocomposite.

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