**In situ** synthesized SrF$_2$/polyvinylidene fluoride nanocomposite film based photo-power cell with imperious performance and stability

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**ABSTRACT**

A self-charged photo-power cell, named as light sensitive energy storage device has been developed in a simplified way for application as a portable photo-charged power bank. The Photo-Power Cell is fabricated by assembling a photoelectrode (FTO) involving a dye-sensitized solar part as photo-electrons generator, combined with the electroactive and high dielectric SrF$_2$-PVDFNC film as storage unit. Electroactive β phase crystallization and large interfacial polarization between the SrF$_2$ NPs and poly (vinylidene fluoride) matrix result in high dielectric value ~ 5141 of the nanocomposite film. The device is charged by up to 1.48 V under visible light illumination (~110 mW/cm$^2$) with a constant discharge current density ~1.64 mA/cm$^2$. The photogenerated charge and power density of the photo-power cell are found to be ~24.3 W/m$^2$ and 2400 C/m$^2$ respectively. Our photo-power cell shows maximum areal specific capacitance ~1600 F/m$^2$ with 92% energy storage efficiency. The achieved overall efficiency (%) is found ~ 2.57. The charging-discharging stability is also tested for a long time span of 12 weeks. The efficacy and durability of our fabricated device has also been demonstrated by glowing up 24 commercially available blue light emitting diodes for 10 days with same intensity by charging one time under light.

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1. Introduction

A major concern for sustained development of our modern society is renewable energy creation and reservation [1]. Replacement of conventional fuel sources like coal, petrol, gas etc. is very important for the overall improvement of global economy due to the increased ecological impairment and fast exhaustion of fuel resources [2–5]. Under this circumstances, the researchers worldwide are looking for a simple, cost-effective and scalable mechanism for clean and reliable energy extraction from natural energy resources like wind, sunlight, water, biomass energy etc. with simultaneous storage of that energy in the same unit [6–8].

Recently, fabrication of piezoelectric nanogenerators, solar cells, metal O$_2$ batteries have proved very promising for harnessing electrical energy from clean and renewable sources like living systems, mechanical, solar and chemical respectively [9,10]. More recently hybrid devices capable of both electrical energy generation and storage with superior power density and stability have been reported with success [11–13]. In such hybrid devices, an extra storage unit is not necessary, which makes them not only more cost-effective but also promising in the field of portable electronics.

Development of nanocomposite materials based photo-power units have been gaining interest among researchers for example, a poly (vinylidene fluoride) (PVDF)/ZnO nanocomposites (NCs) based photo-power unit was proposed by Zhang et al. (energy density $E = 1.4$ mW h kg$^{-1}$) [14]. Later, perovskite structure and dye-sensitized solar cell based few photovoltaic cells with storage ability was reported [15–17]. Wee et al. suggested an organic photovoltaic integrated with carbon nanotube based supercapacitor with low power loss [18]. Power density, storage stability and durability of an integrated photo-power unit are the necessary factors for practical utilization. Proper designing and integration of dye-sensitized solar energy conversion part and high dielectric polymeric film into a single unit has potential to surpass the critical deficiency associated with traditional power utilization [14]. Our
previous attempts of designing such type of two electrode photo-power cells with high dielectric PVDF composites demonstrated both energy conversion as well as storage in one unit [19,20].

Electroactive and high dielectric polymer NCs have been progressively appreciated by the scientists due to their versatile utilization in the domain of piezoelectric nanogenerators, capacitors, thin film transistors, grid levelling, rail runs, non-volatile memories, sensors, actuators and also in biomedical fields [21–24]. Though, extensive researches have been carried out to design PVDF based piezoelectric nanogenerators [25,26], but the utilization of high dielectric PVDF or PVDF based NCs as energy storage unit in hybrid photo-power banks are very limited to the best of our knowledge.

PVDF ([-CH2-CF2-]n) and its copolymers are very promising electroactive semicrystalline thermoplastic polymer with superior piezoelectric, pyroelectric, ferroelectric properties, thermal stability and chemical resistance. Five crystalline polymorphic phases α, β, γ, δ and ε have been recognized in PVDF amongst which β phase is the most electroactive [27–30]. Existence of matrix of the orthorhombic unit cells and all trans(TTTT) planar zigzag chain conformation in electroactive β polymorph leads to optimum dipolar moment per unit cell and consequent highest piezoelectric, ferroelectric, pyroelectric and dielectric properties in comparison to the other phases of PVDF [27,28].

Herein, we have designed a cost-effective, simple, highly efficient and durable two electrode light sensitive energy storage device (LSESD) by using in situ synthesized SrF2 nanoparticles (NPs) incorporated giant dielectric PVDF thin film (named as PSR) associated with an organic dye, phenosafranine-polyvinyl pyrrolidone (PSF-PVP) film and zinc oxide (ZnO) and titanium dioxide (TiO2) NPs as a solar part (Fig. 1). This solar component and the high dielectric PSR film forms the key part of solar energy conversion and photo-generated electrical energy storage respectively.

2. Experimental

2.1. Materials

The materials that are used in our present work are poly(vinylidene fluoride) (PVDF) pellets (Aldrich, Germany, $M_w$: 275,000 GPC, $M_n$: 71,000), Strontium nitrate (CDH, India), Ammonium fluoride (Merck, India), dimethyl sulfoxide, (DMSO) (Merck, India), polyvinyl pyrrolidone (PVP) (Loba Chemie), ZnO and TiO2 NPs, Ortho-phosphoric acid (H3PO4) (Merck, India), Phenosafranine (PSF) (Sigma Aldrich, Germany), FTO coated glass (Sigma Aldrich, Germany).

2.2. In situ synthesis of SrF2/PVDF nanocomposite films

Initially, a solution of PVDF (4%) in DMSO was prepared by...
continuous stirring at 60 °C. Then, Sr(NO₃)₂ and NH₄F were dissolved in PVDF-DMSO solution separately with different molar concentrations (0.01–0.2 M) and corresponding double molar concentrations respectively. Addition of NH₄F into Sr(NO₃)₂/PVDF solution resulted in gradual formation of SrF₂ NPs in the polymeric solution under magnetic stirring at 80 °C. The SrF₂/PVDF NC films were obtained by solvent casting method via evaporation of the solvent in an oven at 80 °C in a dust free atmosphere for 24 h. Simultaneously, pure PVDF film was also prepared under same conditions. For characterization, all the films were stored in a vacuum desiccator. The average thickness of the films was ~20 μm. The different concentrations of SrF₂ NPs in PVDF matrix (0.01 M – 0.2 M) have been tabulated in Table S1 (See supporting information).

2.3. Photo-power cell fabrication

Firstly, the dye solution (20 mg/ml PSF and 100 mg/ml ZnO and TiO₂ NPs in 10% mass PVP solution) were prepared in distilled water by continuous stirring at room temperature. Next the dye solution is drop casted on a FTO coated glass (well cleaned by acetone) and vacuum desiccated at ambient temperature (~25 °C) till the solu-

tion becomes viscous. Thereafter, the previously prepared PVDF/SrF₂ NC (0.2 M) is the melting enthalpy of 100% crystalline PVDF with a value 104.6 J/gm.

The optical absorption and emission properties of all samples were investigated by UV—visible spectrometer (Lambda 25, Perkin Elmer, USA and UV – 3101 PC, Shimadzu) and Cary Eclipse fluorescence spectrophotometer (Agilent Technologies) respectively.

The morphological characteristics of the thin films were studied by field emission electron microscope (FESEM) (INSPET F50, Netherland).

To study the dielectric properties of the composite samples, a digital LCR meter (Agilent, E4980A) was used. The variation of the capacitance (C) and tangent loss (tan δ) with frequency were recorded in the frequency range 20 Hz to 2 MHz at room temperature applying 1 V ac voltage across the two opposite surfaces of the samples. Dielectric constant (ε) and the ac conductivity (σac) of the samples are evaluated using,

\[
\varepsilon = \frac{Cd}{\varepsilon_0 A}
\]

where, d and A are the thickness and area of the samples respectively and f is the frequency in Hz applied across the samples and ε₀ is the permittivity of the free space with value 8.854 × 10⁻¹² Fm⁻¹ [27].

The photovoltaic studies of photo-power cell were carried out using a digital multi-meter (Agilent U1252A) and an Electrometer (Keysight- B2985A).

3. Results and discussions

The fabrication of LSESD and its digital photograph have been represented schematically in Fig. 1. We have used the in situ SrF₂/PVDF NC thin film as the active material for storing the photo-generated electrons or charge carriers by the adjacent solar part. Our study involves detailed characterization and optimization of the optical and dielectric properties of the in situ SrF₂/PVDF NC thin films and then fabrication of a self-charged prototype photo-power pack for ensuring best performances under practical condition.

3.1. Characterization and optimization of SrF₂/PVDF NC film for LSESD

3.1.1. X-ray diffraction (XRD) analysis

The crystalline behaviour of the samples and the formation of SrF₂ NPs are investigated by X-ray diffraction technique as depicted in Fig. 2a. The formation of the nanosized SrF₂ NPs within the polymer matrix is confirmed by the peaks around 2θ = 26.7, 30.8, 44.2 and 52.4 corresponding to (111), (200), (220), (311) (JCPDS No. 06-0262) crystal planes in the XRD patterns of the NCs. Diffraction peaks at 2θ ~ 17.4° (100), 18.2° (020), 19.6° (021) and 26.3° (201,310) are observed for pure PVDF which correspond to nonpolar α-crystallite.

The existence of prominent diffraction peak at 20.6° (020) in SrF₂/PVDF NC films confirms the nucleation of electroactive β-crystallites due to the catalytic effect of the SrF₂ NPs in PVDF matrix [27,28,31,32]. It is also noticeable that electroactive β polymorph nucleation in the SrF₂/PVDF NC films considerably increased with concentration of the SrF₂ NPs up to 16.4 vol %. The ratio of the intensity I₂θ.6 and I₁θ.2 (Fig. 2b) is calculated to measure the amount of α and β phase in the pure PVDF and SrF₂/PVDF NC films [33,34]. The maximum obtained value of this ratio is – 9.01 for 16.4 vol % (PSR 0.15).

3.1.2. Fourier transform infrared spectroscopy

Fourier transform infrared (FTIR) spectra of pure PVDF and NPs
doped films are shown in Fig. 2c. From the spectrum of pure PVDF, the absorbance bands are observed at 489 cm⁻¹ (CF₂ wagging), 533 cm⁻¹ (CF₂ bending), 615 and 764 cm⁻¹ (CF₂ bending and skeletal bending), 795 and 975 cm⁻¹ (CH₂ rocking) due to the α phase and a small absorbance band at 840 cm⁻¹ (CH₂ rocking, CF₂ stretching and skeletal C–C stretching) which is for the β phase [26–28]. However, in the composite film, the α phase gradually diminishes and the electroactive β phase related characteristic absorbance bands became prominent at 475 cm⁻¹ (CF₂ deformation), 510 cm⁻¹ (CF₂ stretching), 600 cm⁻¹ (CF₂ wag) and 840 cm⁻¹ (CH₂ rocking, CF₂ stretching and skeletal C–C stretching) [35]. The intensity of the characteristic bands of β phase shows maximum at 16.4 vol % doping of SrF₂ NPs and then decreases for higher concentrations [36].

By using eqn (1), we have calculated the relative fraction of electroactive β phase content (F(β)) in the NPs doped PVDF films and the variation of F(β) (%) with dopant concentration has been illustrated (volume %) in the Fig. 2d [37,38]. The maximum F(β) value achieved is 80% for the 16.4 (volume %) doping concentration.

3.1.4. Field emission scanning electron microscopy

Surface morphological and microstructural characterization of the unblended PVDF and in situ synthesized SrF₂ NPs incorporated PVDF films are studied in detail by the field emission scanning electron microscopy (FESEM) as shown in Fig. 3. The FESEM image of the pure PVDF thin film represents spherulites with diameter in the range ~50–70 μm confirming the presence of the α polymorph in major amount. But for the NPs modified PVDF thin films, the presence of the comparatively smaller size spherulites (average diameter ~10–15 μm) confirms the nucleation of the piezoelectric β polymorphs [42,43]. The formation of SrF₂ NPs (average diameter ~30–40 nm) in PVDF matrix are also confirmed by FESEM images. The NPs are grown in the PVDF matrix and uniformly distributed within the NC films upto 16.4 vol% (PSR0.15). Homogeneous growth and distribution of the SrF₂ NPs in the polymer matrix up to PSR0.15, leads to maximum electroactive β crystals due to optimum interfacial interaction between the NPs surfaces and the polymer chains. For higher loading concentration, agglomeration of NPs is observed within the polymer matrix thus reducing the interfacial interaction areas with the polymer chains.

3.1.5. Dielectric properties

The variation of the dielectric constant and tangent loss of samples as a function of SrF₂ NPs content at 20 Hz and 2 MHz are shown in Fig. 4a and b. The dielectric constant of the samples have been increased slowly upto PSR0.10 sample with increasing content of the NPs at 20 Hz and a sharp increment in dielectric value is observed at 16.4 vol% of SrF₂ NPs content indicating realization of percolation threshold value for the NC samples. For higher concentrations of the NPs the dielectric gradually decreases. However, the trend in dielectric pattern at 2 MHz is different (Fig. 4b) than at low frequency as dielectric properties at higher frequency is mainly dipolar. We have achieved a remarkable value of dielectric constant ~5.14 × 10³ for PSR0.15 at 20 Hz and 40 at 2 MHz for the same film which are 570 times and 10 times larger than that of the dielectric constant of the unblended PVDF film respectively. The tan δ values of the PSR thin films is increased almost linearly with the increasing SrF₂ content upto PSR0.15 and then decreases for both frequencies (Fig. 4a and b). This variation of the dielectric value may be explained by Maxwell–Wagner–Sillars (MWS) interface polarization [44–47]. Unification and localization of the charge carriers along with the mobilized ions increased at the interfaces of the NPs and PVDF matrix due to the inclusion of the conducting SrF₂NPs within the insulating plastic polymer which consequently leads to the giant enhancement of the dielectric value for the NC samples at a certain doping of the NPs i.e., percolation threshold value (16.4 vol %) [27,48]. The ac conductivity also increases with increasing SrF₂ NPs concentration in PVDF matrix up to 16.4 vol % and also with frequency (Fig. 4c).

Dependence of the dielectric constant, ac conductivity and tangent loss (tan δ) on the frequency of pure PVDF and NPs doped PVDF films are graphically illustrated in Fig. 4d–f. A decrease in the dielectric constant is observed for the SrF₂/PVDF NC samples with increasing frequency but the change in dielectric value with frequency for pure PVDF thin film is very low (Fig. 4d). This issue may be explained directly by the MWS principle [44,47]. The large dielectric constant value at lower frequency is mainly due to the existence of large interfacial polarization i.e., high surface accumulation of charges and the interaction between the short range dipoles at the conducting interfaces of the NCs under externally applied alternating electric field. When the frequency is increased, a restricted movement of the space charges and the low accretion of the charges with frequency within the samples result the failure of the MWS effect and reduction of dielectric value [44,48].

The dependence of the tangent loss (tan δ) with frequency is shown in Fig. 4e. A sharp increment in the tan δ value is seen up to a certain value and then the value sharply decreased with frequency i.e. the dielectric relaxation peaks are observed at different lower frequencies for the NPs modified PVDF thin films upto 16.4 vol % NPs concentration. This tan δ peaks or Debye-like relaxation process occurs in the samples due to large dielectric response at that frequency. The noticeable fact is that the tangent loss peaks are shifted towards higher frequency with increasing loading concentration of the NPs due to the strong dipolar relaxation associated with the SrF₂ NPs embedded in PVDF matrix [34,49]. The frequency dependency of ac conductivity of pure PVDF and SrF₂ NPs incorporated PVDF thin films are presented in Fig. 4f. The ac conductivity of the NPs loaded PVDF samples are almost invariant at low frequency region (20 Hz–1 KHz) and then increased linearly at high frequency zone. The MWS effect and dipolar relaxation mode are the reason for the increment of the ac conductivity whereas the presence of frequency independent plateau is responsible for the constancy of conductivity i.e. dc conductivity at low frequency region. But, for pure PVDF the ac conductivity is increased with frequency throughout the frequency range suggesting purely ac plateau nature of the conductivity [27,34].
Fig. 2. (a) XRD patterns of pure PVDF and SrF$_2$—PVDF NC thin films. (b) Ratio of I$_{20.5}$ and I$_{18.3}$ of the samples. (c) FTIR spectra of pure PVDF and PSR thin films. (d) $\beta$-phase content with increasing SrF$_2$ content from IR spectra. (e) DSC thermographs of pure PVDF and SrF$_2$ NPs incorporated PVDF thin films, the calculated (f) fusion enthalpy and (g) degree of crystallinity of the films from DSC thermographs and (h) TGA thermographs of pure PVDF and PSR samples for investigating the thermal stability of the samples.
3.2. Optical properties

Fig. 5a and b represent the UV–Vis and photoluminescence (PL) spectra of the samples respectively. A broad absorption peak centred at 215 nm (~5.76 eV) is observed for SrF2 NPs doped films with continuous increment of intensity with NP concentrations implying the uniform distribution and strong interaction between NPs and polymer matrix as shown in Fig. 5a. Whereas, no such characteristic peak was observed for the PVDF film in the UV–Vis spectrum. Fig. 5b, shows two strong peaks and a comparably slight small peak at 441 nm (blue), 536 nm (green) and 489 nm (blue) respectively at excitation wavelength $\lambda_{ex} = 202$ nm in the photoluminescence spectra of the NPs incorporated PVDF films. Due to the intimate and uniform conjugation of the SrF2 NP within the polymer matrix, the intensity of the peaks gradually increased with the loading concentration of the NP. But for the same excitation wavelength, pure PVDF has no such characteristic peak. This UV absorption phenomenon of the composite films and corresponding blue and green emission (visible region) make this material a potential candidate for light emitting diode (LEDs), UV protector and sensor applications [50].

3.3. Mechanism and photovoltaic performances of the LSES D

Detailed characterization of the samples shows the PSR0.15 film as an appropriate candidate for device fabrication (Fig. 1) due to its superior dielectric value. The LSES D is fabricated by assembling a photoelectrode (FTO) involving phenosafranine-polyvinyl pyrrolidone with TiO2 and ZnO NPs in the presence of ortho-phosphoric acid (H3PO4) as a photo-electrons generator, combined with the electroactive and high dielectric in situ synthesized SrF2–PVDF NC thin film (PSR0.15) for photo-electron storage. Similarly, another device was also fabricated by replacing the PSR0.15 with pure PVDF film having same dimension. A 40 W tungsten bulb covered with ultra-violet and infrared light eliminator filters is used to charge the device with a power of 110 mWcm$^{-2}$.

Considering the working principle of dye-sensitized solar cell, the proposed mechanism of our device may be divided into two processes, generation of the photoelectrons and storage of the photo-generated charge carriers. The proposed working mechanism of our LSES D is schematically shown in Fig. 6a. A composite mixture of PSF-PVP-ZnO-TiO2 dye solution in aqueous acidic medium (H3PO4) has been used as a solar part for photoelectron and hole generation by absorbing photons under light illumination. While the storage mechanism is mainly controlled by the electroactive and high dielectric PSR0.15 thin film of dimension 0.3 cm x 0.3 cm and thickness 15 μm. Here, FTO coated glass and aluminium acted as the working and counter electrode respectively. The HOMO/LUMO energy configuration of PSF, ZnO NPs, TiO2 NPs and UV–Vis absorption spectrum of the entire dye solution is shown in Fig. 6b and c respectively. Photoelectron generation activity by visible light (i.e. photons) determines the efficiency of the solar part i.e. the inorganic-organic dye composite film casted on the working electrode FTO. The combination of two NPs with PSF-PVP has been used as dye-sensitized solar part due to their excellent optoelectronic, photosensitivity and catalytic characteristics which are essential for improving the photons to electricity conversion efficiency of our prototype LSES D [51,52]. PVP and H3PO4 act as a solid polymer electrolyte and electron suppler within the dye film. [53].

When the device is introduced under visible light (~110 mW/cm$^2$), the PSF molecules are excited by absorbing photon ($h\nu$) and the electrons are jumped from HOMO state (~5.2 eV) to LUMO state of the PSF (~ ~ 3.07 eV). The excited photoelectrons are injected to the conduction band of TiO2 (~ ~ 4.21 eV) NPs arising the oxidation of the dye molecule (PSF$_{ox}$).

$$\text{PSF} + h\nu \rightarrow \text{PSF}^* \rightarrow \text{PSF}_{ox} + e^-$$

Then, the injected photoelectrons in the conduction band of
TiO$_2$ NPs defuse through the adjacent ZnO NPs (~ 4.3 eV) to FTO (~ 4.4 eV). The main role of TiO$_2$ and ZnO NPs in solar unit are to tunnel the electrons through their conduction band to the FTO (Fig. 6b). Later, the electrons travel through the external circuit towards the counter electrode (Al) and accumulate at the junction between the Al and high dielectric PSR0.15 thin film as well as at

Fig. 4. Variation of dielectric properties of pure PVDF and SrF$_2$ NPs incorporated thin films with SrF$_2$ NPs content (a, b) dielectric constant and tangent loss and (c) ac conductivity. The frequency dependency of (d) dielectric constant, (e) tangent loss and (f) ac conductivity of the samples.

Fig. 5. (a) UV-Visible absorption spectra and (b) Photo luminescence spectra (λex = 202 nm) of pure PVDF and SrF$_2$ NPs doped PVDF thin films (PSR0.01, PSR0.05, PSR0.10, PSR0.15 and PSR0.20).
the interfaces of the NPs and polymer matrix. Meanwhile, the photo-generated holes in PSF are replenished by the donating electrons from neighbouring solid polymer electrolyte film (PVP and H3PO4) which reduces the dye (PSF⁺) to its ground state (PSF). In this process PVP and H3PO4 are oxidized.

\[
\text{PSF}_\text{ox} + \text{PVP} \rightarrow \text{PSF}_\text{red} + \text{PVP}_\text{ox}; \\
\text{PSF}_\text{ox} + \text{H}_3\text{PO}_4 \rightarrow \text{PSF}_\text{red} + [\text{H}_3\text{PO}_4]_\text{ox}
\]

Continuous oxidation of PVP and H3PO4 drains the electrons resulting a net positive charge at the interfaces of the counter electrode and dielectric film (PSR0.15). And the counter electrode (Al) is negatively charged due to the large accumulation of the photo-electrons. Therefore, an effective potential difference is developed between the two electrodes which generates photo-voltage in photo-power cell [19,20,54–56]. When, the device is fully charged i.e. the open circuit voltage (Voc) becomes ~ 1.48 V, the oxidized dye PSF⁺ is ceased to be reduced back to its original state and the current flow in the device is dropped to zero. The current-voltage (J-V curves) and self-charging-discharging phenomena (V-t graph) of our LSESD are illustrated in Fig. 7a–d. A high short circuit current (Isc) (~3.98 mA/cm²) is measured during charging process for the LSESD containing PSR0.15 [57]. Whereas, Isc is ~0.26 mA/cm² for pure PVDF based LSESD (Fig. 7a and b). Under light illumination, the open circuit voltage (Voc) ~ 1.48 V is obtained within 1 min for the LSESD involving PSR0.15 as storage part which is 1.5 times larger than that of the pure PVDF based LSESD. Closer observation of the self-discharge curve (shown in Fig. 7c and d) signifies the superior storage stability of the PSR0.15 based photo-power cell than the pure PVDF based device as the V-t discharge characteristic is almost time independent for a long time. The discharge time i.e. internal electron recombination span is very low for the pure PVDF based device (Fig. 7d) due to the very small value of dielectric constant of the pure PVDF film. These lead to lower charge storage capacity and stability of that device for practical utilization. The internal discharging action of the photo-power cell is totally performed in a dark atmosphere.

In dark atmosphere, the photo-electrons generation process has been ceased as there is no photon to excite the PSF molecules. However, high dielectric PSR0.15 thin film behaves like an insulating medium and the lower conductivity nature of the PSR0.15 thin film allows very few electrons to pass through the insulating medium for recombination with holes. Thus a very small voltage drop and a small discharge current (~160 µA) is observed due to recombination of electrons and holes under this condition.

The performances of the two fabricated devices are investigated by discharging fully charged devices with a constant discharge current density ~1.64 mA/cm² for LSESD constructed using PSR0.15 and 1.07 mA/cm² formed using pure PVDF film under the dark ambient (Fig. 7e and f). The discharge time is found to be ~146 s and 12 s for the PSR0.15 and pure PVDF incorporated LSESD respectively. An estimated value of the stored charge density, specific areal capacitance, energy density and power density of our LSESD have been calculated from the discharge V-t graph (Fig. 7e and f) by using the following equations

\[
Q = \int I_{\text{dis}} \, dt \\
C = \frac{Q}{dV} \\
E_{\text{output}} = \frac{1}{2} CV^2 \\
P = \frac{V}{t_{\text{dis}}}
\]

where, \(Q\), \(C\), \(E_{\text{output}}\), \(P\), \(I_{\text{dis}}\) and \(dV\) are the charge density, specific areal capacitance, output energy density, power density, discharge current density, discharge time and voltage difference respectively.

A highest charge storage capability of ~2400 C/m² has been achieved for our LSESD with PSR0.15 with superior energy and power density ~500 mW h/m² and ~ 24.3 W/m² respectively (Fig. 8a). The storage ability i.e. specific areal capacitance(C) is found to be ~1600 F/m² (Fig. 8a). These values are significantly high compared to previously reported photo-power units or photo-supercapacitors (See comparison in Table S2 in supporting information). The output characteristics are also calculated for the device fabricated using pure PVDF film and shown in Fig. 8b.
To find out the energy conversion efficiency ($\eta_{\text{conversion}}\%$), we have used the following equation,

$$\eta_{\text{conversion}}\% = \frac{P_{\text{out}}}{P_{\text{in}}} \times 100 = \frac{V_{\text{OC}} \times I_{\text{SC}} \times FF}{P_{\text{in}}} \times 100$$  \hfill (9)

$V_{\text{SC}}$ = open circuit voltage, $I_{\text{SC}}$ = short circuit current, $FF$ = fill factor, $P_{\text{in}}$ = the incident light power (110 mW/cm$^2$) [51].

Where, $FF = \frac{V_{pp} \times I_{pp}}{V_{OC} \times I_{SC}}$  \hfill (10)

$V_{pp}$ = voltage power point, $I_{pp}$ = current power point, where $I_{SC}$ and $I_{pp}$ are in mA/cm$^2$, $V_{OC}$ and $V_{pp}$ are in V, $P_{in}$ and $P_{out}$ are in mW.

By using equations (7), (9), (11) and (12) the energy storage efficiency is evaluated as

$$\eta_{\text{storage}} = \frac{\eta_{\text{overall}}\%}{\eta_{\text{conversion}}\%}$$  \hfill (11)

Overall efficiency. $\eta_{\text{overall}}\% = \frac{E_{\text{output}}}{E_{\text{input}}} \times 100$  \hfill (12)

where the input energy density $E_{\text{input}} = P_{\text{in}} \times t_f$, $t_f$ is the charging time [58].

The values of energy conversion, storage and overall efficiencies and the output characteristics (energy density, power density and storage ability) of the devices constructed with both PSR0.15 and pure PVDF thin films are graphically represented in Fig. 8c and d respectively.

We have achieved 92% energy storage efficiency associated with 2.78% energy conversion and 2.57% overall efficiency.
which are very promising and much better than the previously reported studies (Table S2). The efficiency values of PSR0.15 based LSESD are also many times larger than that of the efficiencies of the pure PVDF based LSESD (Fig. 8d). Our photo-power cell developed with PSR0.15 as storage part exhibits superior recyclability and storage stability for long times. We have investigated its performances for 12 weeks without any noticeable degradation in the output characteristics, storage ability and efficiencies. Finally to analyse the self-charging ability (V-t curve), storage stability and practical application as photo-power bank, we have connected three LSESD made of PSR0.15 serially. This serially connected combo pack have been charged upto ~4.1 V under light illumination within a short time span of 1 min (Fig. 8e). And the combo power bank is able to light up 24 serially connected blue LEDs continuously up to 10 days with same intensity demonstrated schematically in Fig. 8f and Video S1 (Supplied as supporting data). Here, we have used a fundamental and low cost technique to design a very simple and single hybrid self-charging energy storage device with very efficient energy creation and storage capability. This power pack has significant potential due to its superior performance and charge storing stability for acceptance in clean renewable energy developing areas for overcoming the daily energy demands.

Supplementary video related to this article can be found at https://doi.org/10.1016/j.electacta.2018.06.054.

4. Conclusions

In summary, we have grown SrF$_2$ NPs via in situ process within the PVDF matrix to develop a simple and cost-effective self-charged photo-power system efficient in energy harvesting and storage. In situ growth of SrF$_2$ NPs in PVDF matrix improves the electroactive β-phase formation (~80%) and the dielectric value of the PVDF from 9 to 5141. The maximum open circuit voltage ~1.48 V and short circuited current density ($I_{sc}$) ~ 3.9 mA/cm$^2$ are obtained from PSR0.15 based LSESD under the photo illumination with superior power density (~24.3 W/m$^2$) and storage capability (~1600 F/m$^2$). The maximum energy conversion, storage and overall efficiency of the self-charged LSESD are calculated to be ~2.78, 92 and 2.57% respectively. The storage potentiality of our LSESD is much higher than that of the earlier reported photo-induced power cells. We have also demonstrated the practical utilization of our photo-power unit lightening commercially available 24 blue LEDs with same intensity for 10 days by charging one time. Such high storage efficiency combined with stable reusable capacity of our self-charging photo-power cell has potential application possibilities in the field of portable and large scale energy demanding areas of our modern society.

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