Asymmetric Supercapacitors: Optical and Thermal Effects When Active Carbon Electrodes Are Embedded with Nano-Scale Semiconductor Dots

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Abstract: Optical and thermal effects in asymmetric supercapacitors, whose active-carbon (AC) electrodes were embedded with nano-Si (n-Si) quantum dots (QD), are reported. We describe two structures: (1) p-n-like, obtained by using a polyethylimine (PEI) binder for the “n” electrode and a polyvinylpyrrolidone (PVP) binder for the “p” electrode; (2) a single component binder—poly(methyl methacrylate) (PMMA). In general, AC appears black to the naked eye and one may assume that it indiscriminately absorbs all light spectra. However, on top of a flat lossy spectrum, AC (from two manufacturers) exhibited two distinct absorption bands: one in the blue (~400 nm) and the other one in the near IR (~840 nm). The n-Si material accentuated the absorption in the blue and bleached the IR absorption. Both bands contributed to capacitance increase: (a) when using aqueous solution and a PMMA binder, the optical-related increased capacitance was 20% for low n-Si concentration and more than 100% for a high-concentration dose; (b) when using ion liquid (IL) electrolyte, the large, thermal capacitance increase (of ca. 40%) was comparable to the optical effect (of ca. 42%) and hence was assigned as an optically induced thermal effect. The experimental data point to an optically induced capacitance increase even in the absence of the n-Si dots. Overall, the experimental data suggest intriguing possibilities for optically controlled supercapacitors.

Keywords: asymmetric supercapacitors; polymeric-doped supercapacitor electrodes; nano-semiconductor dots embedded in active carbon electrodes; supercapacitors: optical effects; supercapacitors: thermal effects

1. Introduction

Supercapacitors, S-C [1–3]—capacitors that take advantage of the capacitance at the interface between an electrode and an electrolyte—have been found in a large range of energy applications—least of all, optical modulators [4], and as buffering elements to subdue demand fluctuations in digital power networks [5,6]. A subclass of these is asymmetric S-C—capacitors made with two types of electrodes [7–9]. Asymmetric S-C are mostly fabricated as pseudo-supercapacitors—S-C whose capacitance is associated with a chemical reaction at the electrolyte/electrode interface. Pseudo-supercapacitors are aimed at increasing the operation potential range [10–14] via electrochemical means. To these, one ought to add dye-sensitized solar cells [15,16], a special class of optically powered electrochemical energy sources with two distinct electrodes and an electrolyte that mediates the reacting ion species. Here, we describe carbon-based, optically controlled asymmetric S-C that do not exhibit a chemical reaction at their electrolyte/electrode interfaces; namely, we focus on basic, electrochemical double-layer (EDL) supercapacitors. In the past, nano particles were incorporated within active-carbon electrodes, yet meant for only electronic purposes [17–19]. Our intent is to gain a basic understanding of the optical and related thermal effects when incorporating n-Si dots in active carbon (AC) based electrodes. We describe two S-C types, which are both embedded with n-Si dots: polymeric-doped AC electrodes...
via polyvinylpyrrolidone (PVP) and polyethylenimine (PEI) binders, and non-doped AC electrodes with a poly(methyl methacrylate) (PMMA) binder.

2. Materials and Methods

The basic optoelectronic cell is composed of two transparent electrodes, either indium tin oxide (ITO, sheet resistance Rsqr = 20 Ohms) or fluorinated doped tin oxide (FTO, sheet resistance Rsqr = 30 Ohm) films on glass substrates. The electrodes were facing each other to form a parallel plate capacitor. The electrodes were coated with active-carbon (AC) film (either produced by American Hardwood, AH, or by General Carbon Company, GCC). In the case of p-n cells, the AC was incorporated in binders similarly to functionalize carbon nanotubes [20–25]. While the AC is not a semiconductor material, nevertheless, it was hypothesized that the small AC domains would make it susceptible to polymeric doping.

**PVP and PEI Binders, p-n-like Cells:** In the case of a p-n-like cell, the AC (various concentrations in the range 100–200 mg/mL) in methanol was dispersed using a sonicator with a horn antenna. The n-Si dots, of size less than 100 nanometers and at various concentrations in the range of 1–10 mg/mL, were mixed in mostly the PVP “p-type” material. The concentration range of the PVP was 20–40 mg/mL but larger than 10% by weight. Two molecular weights have been considered for the polymers: low m.w. of 25 kD and high m.w of 630 kD. Two molecular weights were also considered for PEI: low m.w. of 25 kD and 50% water diluted 600 kD. High m.w polymers were used previously [24] with CNT and they were thought to have better AC “doping” than the low m.w. polymers. However, preliminary results with high m.w polymers exhibited large equivalent series resistance (ESR) and relatively small capacitance values. Therefore, we focus here on low m.w. polymer binders. The irradiation on n-Si embedded PEI resulted in pseudo-capacitor behavior (Figure 1d) and, hence, n-Si were incorporated into either the PVP or the PMMA. The slurries were drop-casted on the transparent electrodes and were dried out at 90 °C for 30 min. The entire structure was held by a strong clip, leaving an exposed surface for light illumination. The illuminated (exposed) area was smaller than the entire area of the S-C. Typically, the area for p-n samples was 1 × 1 cm² whereas the AC/PMMA samples were larger, ca. 2 × 2.5 cm². The capacitance change of a given sample was assessed as a relative value: under light irradiation and without it. Sample temperature increased as a result of the optical absorption. A baseline for capacitance change due to pure thermal effect was established.

![Figure 1. Cont.](image-url)
Figure 1. (a,b) Cyclic voltammetry (CV) curves at scan rate of 0.1 V/s for AC/PVP-AC/PEI low m.w. cell with ion liquid (IL) electrolyte and Al current collectors. (a) When the positive lead of the potentiostat was connected to the AC/PVP side and the negative lead was connected to the AC/PEI side. (b) When the leads have been switched: the positive lead of the potentiostat was connected to the AC/PEI side and the negative lead was connected to the AC/PVP side. (c) Charge-Discharge (CD) curves: red curve is when the positive lead is connected to AC/PVP; blue curve is when the positive lead is connected to AC/PEI. (d) Incorporating n-Si in both electrode material and, more specifically, in AC/PEI resulted in a pseudo-capacitor behavior when using an aqueous electrolyte. (e,f) 30 × 30 micron$^2$ AFM scanned images of AC/PVP/n-Si and AC/PEI electrodes, respectively, on FTO. The color height bar extends from −6 (dark) to 5 (bright white) microns and their respective linear scans. The respective linear scan along 50 microns is shown below each image. The y-scale (height) is from −4 to +4 microns.

Film surface was analyzed by Atomic Force Microscope (AFM) as shown in Figure 1e,f. Film thickness was assessed by measuring the substrate weight difference upon film deposition, ΔW, and taking into account the fraction weight, f, of each film component: d$_{\text{eff}}$, the effective film density in g/cc, is defined as d$_{\text{eff}}$ = f$_1$d$_{\text{AC}}$ + f$_2$d$_{\text{polymer}}$. The film thickness was derived from, t = ΔW/(A·d$_{\text{eff}}$), with A being the film area. For example, for a p-n-like film and neglecting the small weight of n-Si, f$_2$/f$_1$ = 0.15. The density of AC was measured as 0.35 g/cc and the density of PVP is 1.2 g/cc. Therefore, d$_{\text{eff}}$ = 0.85 × 0.35 + 0.15 × 1.2 = 0.48 g/cc. The weight difference was 3 mg. Thus, t = 3/(480 × 1$^2$)~63 microns.
**PMMA Binder:** The adhesion of PEI and PVP to the conductive glass with IL was good; the films sometimes peeled off when aqueous cells were taken apart. PMMA binder worked better under these circumstances with both IL and Na2SO4 electrolytes. The n-Si were incorporated within the electrode facing the light source. Typically, a 200 mg/mL AC produced by GCC with a 20 mg/mL PMMA binder on FTO or ITO was used. The 2 mg/mL n-Si was added to the slurry with the other components in Anisole, sonicated with a horn antenna and drop-casted on the conductive glass substrate. The film was dried out in an oven at 90 °C for 30 min. With a PMMA binder, \( \frac{f_2}{f_1} = 0.1, A \approx 5 \text{ cm}^2, \Delta W = 34 \text{ mg}; t \approx 156 \text{ microns}. \)

**Electrolytes:** 98% ionic liquid (IL, Alfa-Aesar) [26–28] and 1 M Na2SO4 were used as electrolytes. For IL, we used 1-n-Butyl-3-methyl-imidazolium hexafluoro-phosphate. The ionic liquid was soaking 0.1-mm-thick lens tissues (Bausch & Lomb) used as separators. Using a hydrophilic nano-filtration polyamide filter (TS80, Sterlitech) proved to be too resistive for the as-is IL; both the tissue and the membrane worked well with the 1 M of Na2SO4 electrolyte.

**Electrochemical Measurements:** Measurements were carried out with a Potentiostat/Galvanostat (Metrohm). The samples were irradiated with a 75-W incandescent light bulb situated 30 cm above the samples. The light intensity of the entire radiation spectra (from the visible to the IR) was measured with a bolometer and was assessed as 30 mW/cm². A calibrated homemade hot plate, which was interfaced with a thermocouple, was used for the thermal experiments. A second thermocouple assessed the temperature precisely at the sample surface.

**Electrical Measurements:** Current-voltage plots (I-V curves) on dry films were obtained with a sensitive, 100 fA, computer-controlled dedicated system (Keithley).

**Optical Transmission Measurements:** A computer-controlled monochromator (SPEX), which was interfaced with a white light source, a chopper and a Si detector, was used to assess the optical transmission of each film on a glass substrate. The transmission value was assessed as the signal obtained with the film on the glass slide divided by the signal obtained with only the glass slide.

3. Results and Discussion

3.1. Film Characterizations

**Asymmetric Cells:** The asymmetry of the capacitive element is demonstrated in Figure 1a,b. Shown are cyclic voltammetry (CV) curves at a scan rate of 0.5 V/s for a cell made with concentrations of AC, PVP, PEI, as follows: 100 mg/mL, 20 mg/mL and 20 mg/mL with the lower m.w. polymers. The “p”-like electrode was made of AC/PVP on Al and the “n”-like electrode was made of AC/PEI also on Al. The AC was made by AH and the electrolyte was IL soaking a tissue separator. The curves were mirror images of each other when the positive and negative leads were switched; namely, the effect may not be attributed to Schottky barrier at the contacts. Figure 1a was obtained when the positive lead of the potentiostat was connect to the AC/PVP electrode (denoted as \( p^+ \)) and the negative lead of the potentiostat was connected to the AC/PEI side (denoted as \( n^- \)). Figure 1b was obtained when lead connections were switched—namely, \( n^+ - p^- \).

Charge-discharge (CD) curves (Figure 1c) conveyed the same message: the rise time is shorter and the discharge time is longer for \( p^+ - n^- \), whereas the reverse is true when the lead connections are switched. Granted that the S-C is less than perfect; it was of the order of tens of micro-F for this 0.5 cm² capacitor. Figure 1d exhibits a pseudo-capacitive nature when the n-Si is incorporated in the AC/PEI electrode.

The film surface is shown in Figure 1e,f. The AFM scan in a cell of 30 × 30 microns² exhibited a grain width of ca. 15 microns and height in the range of ±4 microns (limited by the machine specifications). Overall, it seems that the grain size here is of the order of 15 microns.

**Optical Transmission and I-V Curves:** The optical transmission through the various material components is shown in Figure 2a. Since the films’ thickness varied and we are only interested in the spectral shape of the curves, each transmission curve was normal-
ized to its peak transmission. The glass slide signal was referenced to the transmission through air; all other curves were referenced to the signal of their substrate—a glass slide. The transmission of a glass slide is fairly constant throughout the spectral range between 400 and 900 nm. The transmission of the ITO and FTO on glass is flat throughout the visible with an overall transmission coefficient of 0.8; PMMA has also a flat transmission in the visible with a transmission coefficient of 0.9 (not shown).

![Graph](image)

**Figure 2.** (a) Normalized optical transmission through the various film components used in the experiments. The peak transmission was set to 1 for each curve in order to accentuate the spectral response. The transmission of the glass slide was referenced to air. All other transmission values were referenced to the transmitted signal through the glass slide. (b) I-V curves for AC/PVP and AC/PVP/n-Si under illumination of 30 mW/cm² light bulb and in the dark (room lighting). The inset shows the measurement arrangement. The effect is small but nonetheless measurable. (c) The setup for measuring conductivity with and without illumination (the light irradiates the film from above). (d) A picture of AC/PVP film bridging a scratched ITO glass substrate.

The yellowish n-Si powder absorbs heavily in the blue green spectral range, as is typical for these nano-scale dots. The indirect bandgap of Si at 1.1 microns turns into a direct bandgap and its absorption is blue-shifted when the dot size becomes smaller. The AC film exhibited two distinct absorption bands regardless of maker (the one depicted in the picture has been produced by American Hardwood (AH)). One absorption band is in the blue, centered at 460 nm, and the other is in the near IR, centered at 840 nm. PVP absorbs in the deep blue and portrays flat absorption for wavelengths between 500 and 900 nm. The absorption of AC/PVP followed that of only AC. Most notable is the transmission of PVP/n-Si (not shown) and AC/PVP/n-Si. The absorption peak at 840 nm disappeared, leaving only an absorption peak in the blue. Additionally noted is the blue shift for the 460 nm absorption line to below 400 nm with a combined effect that is larger than for either components. The behavior at 840 nm may be explained if the absorption line is attributed to impurity doping or surface states. Electrons were transferred from the AC (donors) to the n-Si (acceptors) and the transition was bleached. The behavior near 400 nm is more complex and could involve dipole coupling between the AC and n-Si species.

In Figure 2b, we show the I-V curve of illuminated and non-illuminated dry films made of AC/PVP and AC/PVP/n-Si on ITO. The setup for the conductivity measurements is shown in Figure 2c and a picture of a scratched sample is shown in Figure 2d. As shown,
a 1-mm scratch was made in the ITO layer, preventing electrical conduction across it. The deposited film was bridging the gap and enabled conduction (Figure 2c,d). There are two takeaways from the experiments: (1) the ITO/AC films exhibited Ohmic contacts; and (2) the film conductivity has increased under white light illumination. The films’ thicknesses and widths were not the same, which explains the difference in the curves’ slopes. Nevertheless, we can assess the relative conductivity \( (\sigma_{\text{illuminated}} - \sigma_{\text{dark}})/\sigma_{\text{dark}} = 5.5 \times 10^{-2} = 5.5\% \) for the AC/PVP/n-Si and \( 3 \times 10^{-2} = 3\% \) for the AC/PVP film. As expected, the n-Si has improved the film conductivity under light even though its concentration was of the order of a few percent compared with a typical 20% conductive additives in commercial S-C material [28].

3.2. Supercapacitors under Light ON and Light OFF Conditions

In the remainder of this paper, we will examine the optical and related thermal effects in these films. In Figure 3, we show CV and CD curves for a p-n cell—p(AC/PVP/n-Si)-n(AC/PEI)—under light ON and light OFF. The CV curves are asymmetric when light is OFF (room light) and ON. The sample was deposited on ITO with a lens tissue separator soaked with IL. Capacitance increase can be observed in Figure 3a. The relative capacitance increase should be normalized by the respective differences between the illuminated and non-illuminated areas: \([(C/A)_{\text{illum}} - (C/A)_{\text{dark}}]/(C/A)_{\text{dark}} = 0.42 \) or a 42% increase under illumination. Upon illumination, one may observe a tilt of the plot axis towards larger current values. Such tilt may be attributed to a larger film conductivity and a lower ESR. The latter is corroborated by CD data (Figure 3b). Capacitance here is calculated for the discharged branch as \( C_{\text{eff}} = (I_0/V_0/\tau) \), which replaces the usual linear expression, \( C = I_0/(dV/dt) \). Here, \( I_0 \) is the (constant) discharge current, \( 1/\tau \) is a single decay rate that approximates the discharged branch, \( V_0 \) is the voltage difference (\( V_0 = 1 \) V in our case) and \( C_{\text{eff}}, \) the effective capacitance across the various film regions as it is gradually discharged. The relative capacitance change under illumination is ~37%. Overall, the ESR is quite large and may be attributed to the large impedance of the current collector (ITO; \( R_{\text{sqr}} = 20 \) Ohms compared to 2.7 mOhms of Al), use of IL and non-optimized binder to AC ratio.

![Figure 3. (a) A picture of a p(AC/PVP/n-Si)-n(AC/PEI) cell with tissue-soaked IL electrolyte on FTO glass. The cross-section of the p- and the n-electrodes is 1 cm². (b) CV at a scan rate of 0.5 V/s under illuminated and non-illuminated conditions. (c) Corresponding CD curves. A small ESR decrease is noted under illumination. Note the rotated curve in (a) and the increase in both the charge and discharge time constants in (b) under the illumination with white light.](image)
3.3. Thermal Considerations during Light ON and Light OFF Conditions

3.3.1. PMMA Binder with Aqueous Solution

Let us start with the simpler system using a single electrode binder (PMMA), where the electrode facing the light source is embedded with n-Si dots. In Figure 4a,b, we present two CV plots: Figure 4a has been obtained when heating the sample from 23 to 35 °C, at the rate of 0.1 °C/s, while collecting the CV data continuously. The light was OFF and the curve served as a reference. The AC manufacture was GCC and the electrolyte was 1 M Na₂SO₄. The loading of the n-Si was 2 mg/mL. Note the rotation of the curve as the sample heats up; its waist at zero potential has changed only slightly. The relative change from the first scan to the last was <2%. Figure 4b exhibits two CV curves for light OFF and light ON. Upon illumination, the temperature at the sample surface has elevated to 27 °C. The relative capacitance change for the ON/OFF case was 20% (including the fact that the exposed illuminated area is smaller than the area of the entire S-C), clearly larger than the thermal reference. While the curve has rotated, its waist has increased too. Finally, CV curves were obtained with AC electrodes and only PMMA as a binder (namely, without n-Si). As seen from Figure 4c, there is a small optical effect when considering the smaller illuminated area of the S-C, while the temperature at the sample surface has reached 30 °C under optical illumination. Heating of the AC electrode even without n-Si is attributed to an overall AC scattering/absorption. Thus, while the optical-related heating is substantial, the increase in local polarization is small.

![Figure 4](image-url)

**Figure 4.** CV curves for AC in PMMA binder with 1 M Na₂SO₄ electrolyte soaking a TS80 membrane and n-Si loading of 2 mg/mL. Curves were obtained at a scan of 0.1 V/s. (a) Calibration curve when the sample was heated when light was OFF from 23 to 35 °C. Note the rotation of the curve as the sample heats up; its waist, however, at zero potential barely changed. (b) CV curves for light OFF (room lighting) and light ON. While the curve has rotated, its waist has increased too. (c) AC with PMMA binder yet without n-Si has exhibited a small optical effect if we take into account the smaller area exposed to the light.

An easy calibration method may be made with the one S-C cell. In that case, one compares the effect of illumination for two cases: the case when the electrode containing n-Si
dots is facing the light source and the case when the other electrode, the one without
the dots, is facing the light source. This experiment is presented in Figure 5. Both electrodes
were deposited on an ITO with Rsqr = 5 Ohms (Huanyu). The electrolyte was Na₂SO₄ and
the cell has reached 26 °C in both cases during illumination. The concentration of the n-Si
dots was 10 mg/mL, substantially larger than all other cases described before. Specifically,
when the electrode without n-Si was facing the light source, the relative OFF/ON change
was ca. 11% (22% when considering the smaller light exposed area); it was 56% (112% when
considering the smaller light exposed area) for the electrode containing the n-Si dot and
facing the light source. Based on Figures 4c and 5a, one may postulate that there could be
an optical effect even in the absence of n-Si dots due to absorption of AC in the blue.

![Figure 5](image)

**Figure 5.** Measuring the cell capacitance when the electrode without the n-Si dot is facing the
light source, (a), and when the electrode containing the dots is facing the light source. In both cases,
the surface of the sample reached 26 °C. The n-Si loading was 10 mg/mL, (b).

### 3.3.2. PMMA Binder with Ion Liquid Electrolyte

In order to assess whether the capacitance increase is due to the resulting heating
of the electrolyte through light absorption, we repeated the previous measurements and
replaced the aqueous solution with an IL electrolyte. The AC electrode here was embedded
with a higher n-Si concentration—10 mg/mL. Figure 6a presents the capacitance increase
upon heating from 23 to 45 °C at a rate of 0.1 °C/s. Here, too, the light was OFF and the
heated sample served as reference. The CV data were continuously collected throughout
the heating process. Figure 6b shows the related CV curves upon light ON and light OFF.
A large capacitance increase is noted: the relative capacitance increased by 50% when the
sample was heating up from 23 °C to 37 °C (Figure 6a). Similarly, a ca. 50% capacitance
increase and related temperature increase is noted based on Figure 6b. Thus, and unlike
Figure 4, we cannot separate here the optical effect from the electrolyte-related, thermal
effect. Similar conclusions could be drawn when a lower concentration of n-Si was used.

### 3.3.3. p-n Binders with Ion Liquid Electrolyte

CV were obtained for p-n-like cell (n-Si in AC/PVP and AC/PEI, respectively). The electrolyte was IL. In Figure 7a, we present a CV curve when heating the sample
from 23 °C to 35 °C while collecting the CV data continuously, all while the light source
was OFF. The heating rate was 0.1 °C/sec. The relative change from the first scan to the last
was ca. 40%. Figure 7b exhibits two CV curves for light OFF and light ON. Upon illumination,
the temperature at the sample surface has elevated to 30 °C. The relative capacitance change
for the ON/OFF case was 35%. Using the calibration curve of Figure 7a, the thermal
effect between 23 and 30 °C may account for only 23% of the capacitance increase while the
remaining 13% could be attributed to the optical effect. This last value is similar to the value
obtained for the aqueous system. Nonetheless, since the p-n is a more complex system
than the one involving PMMA, and recognizing that the conductivity of PVP changes upon
light illumination (Figure 2b), more data are needed to affirmatively determine the portion
of the optical effect with IL electrolytes. We point to recent studies that showed that the
The thermal sensitivity of IL could be substantially decreased by using a proper mixture [27] and could be used here to accentuate the optical effect; such a study is beyond the scope of the current manuscript.

![Figure 6](image_url) CV curves for AC embedded with high concentration of n-Si (10 mg/mL) with a PMMA binder. The IL electrolyte was soaking a lens tissue. Curves were obtained at a scan of 0.1 V/s. (a) Calibration curve when the sample was heated when light was OFF from 23 to 45 °C. Note the rotation of the curve as the sample heats up. (b) CV curves with light OFF (room lighting) and light ON. The sample heated up from 23 °C to 37 °C upon illumination.

![Figure 7](image_url) CV curves for p-n-like cell with IL electrolyte. Curves were obtained at a scan of 0.1 V/s. (a) Calibration curve when the sample was heated when light was OFF from 23 to 35 °C. (b) CV curves when the white light illumination was OFF and ON. Overall, the thermal effect from the IL dominated the capacitance increase.

### 3.3.4. p-n Binders with Aqueous Solution

The thermal response of p-n-like cells may be corroborated by using Al electrodes, 1 M \( \text{Na}_2\text{SO}_4 \) electrolyte and eliminating the n-Si QDs from the electrode composition. Figure 8 shows CD curves for the p(AC/PVP/n-Si)-n(AC/PEI) sample. The sample was heated by a 75-W lamp (although the Al current collector prevented any light penetration) and the temperature was recorded at the sample surface by using a thermocouple point probe. The AC was manufactured by GCC. The <2% difference in the discharge time may be more to do with the instability of the binder in the aqueous electrolyte than with the heat. Therefore, we reiterate that aqueous solutions at the small temperature range studied have little thermal effect on the cell capacitance.
Figure 8. CD curves at two temperatures for S-C, p(AC/PVP/n-Si)-n(AC/PEI) interfaced with Al electrodes and Na₂SO₄. The decay times for the discharge are shown in the figure. The slight change (<2%) is attributed to imperfect contacts. No major capacitance change is noted when using the aqueous solution.

3.4. The Effect of IR Absorption Band at 840 nm

As discussed earlier, the incorporation of n-Si dots bleached the absorption band at 840 nm. Further corroboration may be obtained when measuring the capacitance while using a yellow optical filter. The filter transmits wavelengths larger than the cut-off wavelength of 550 nm, thus eliminating the blue band from the white-light spectra. One has to factor in the reduced overall light intensity when the light source is interfaced with such a filter. The intensity of the light source interfaced with the filter was 70% of the total white light output (both visible and IR) as measured by use of a bolometer. Thus, if the intensity-related effects are linear, the capacitance increase due to the optical effect and/or thermal effect would be 70% of the effect without the filter.

Figure 9a shows that a p(AC/PVP/n-Si)-n(AC/PEI) on FTO with IL electrolyte, illuminated with a filter-interfaced white-light source, exhibited the same characteristics as non-illuminated S-C. On the other hand, a p(AC/PVP)-n(AC/PEI) on FTO (without n-Si) in Figure 9b showed that the characteristics of light ON with and without the filter are the same; both were different from the light OFF situation. When using the filter, we eliminated the heating through blue light absorption but absorption was still present through the IR band.

All of the above may be summed up as follows: the IL was heated up through only the blue band absorption when the electrode contained n-Si; the n-Si eliminated the IR absorption band and the filter eliminated the blue absorption band. In the absence of n-Si, the sample was heated through both the blue and the IR bands; while the filter eliminated...
the absorption in the blue, the electrolyte was still heated through the absorption in the IR. In both cases, there was an increase in the relative cell capacitance.

4. Conclusions

Asymmetric S-C, embedded with n-Si QD, have shown a large capacitance increase. For aqueous cells, this was due to local polarizations even in the absence of n-Si dots. Cells interfaced with IL electrolyte exhibited a capacitance increase that was attributed to optically induced thermal effects in the electrolyte itself. Overall, such optoelectronic elements exhibit promise for novel, optically controlled supercapacitors.

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