Conducting Poly(aniline)-Electrical Charge Transportation

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ABSTRACT

Conductive polyanilines are synthesized by doping with inorganic and organic acids, namely Hydrochloric acid (HCl) and ±10-camphor sulfonic acid (CSA). The direct current (DC) conductivities ($\sigma_{DC}$) are found to be about $9.5 \times 10^{-8}$, 1.8, and 95.8 S/cm for PANI base, PANI (HCl) and PANI (CSA), respectively. $\sigma_{DC}$ is measured down to a temperature of $\sim 100$ K and the apparent change in the activation energies are found to be 98.16, 74.40, and 57.24 meV for PANI base, HCl, and CSA dopings respectively. $\sigma_{DC}$ is less temperature dependent near room temperature, further decrease in temperature the $\sigma_{DC}$ is strongly dependent. Upon the inspection of AC conductivities ($\sigma_{AC}$) versus frequency curves, it can be inferred that the conduction process is noticeably influenced upon doping and within the dopants. $\sigma_{AC}$ has shown classical plateau (DC-AC crossover) region, nonetheless shifted crossover frequency (critical frequency) upon doping is rather interesting. Critical frequencies ($\omega_c$) are obtained from universal power-law for all samples. The variation in the dielectric properties can be attributed to the dopant incorporation. In material characterization, successful doping is corroborated by FTIR, UV-vis spectroscopy and slight influence upon doping can also be seen in thermal properties. Intense photoluminescence (PL) peaks at 322.5, 581.4 and 644.2 nm are observed. PANI (CSA) exhibited highest peak intensity followed by PANI (HCl) and PANI base.

Keywords: Protonation; Composites; Electrical Transportation; AC Conductivity

1. Introduction

In general, conducting polymers are a special class of organic polymers that have alternate single-double bond conjugation and can conduct electricity. Among all the conducting polymers polyaniline (PANI) demonstrates outstanding properties due to its environmental stability, redox reversibility, high electrical conductivity and ease of synthesis which drives it towards potential electrical device applications [1-5]. It is agreed that in most of its applications the behaviour of the conductivity is a long standing problem. The improvement on electrical properties of doped PANI reflecting the conditions of preparation/dopant incorporation is of fundamental importance. In comparison with metal oxides, acid doped conducting polymers influence more on electrical properties. Nevertheless the conductivity needs to be improved to reach lab scale device fabrication. Interest has focused upon the discovery of doping by acids with certain functionalized protonic acids may be used to render the emeraldine salt (ES) form of the polymer processable from solution [6,7]. Heeger and co-workers have have reported the use of functionalized protonic acids to dope emeraldine base (EB) and, simultaneously, to make soluble the resulting PANI complex in common organic solvents [8]. This procedure increases the PANI compatibility with bulk polymers with similar molecular structures. In addition it can strongly influences the electrical properties of a device. Consequently a wide range of conductivities are obtained by doping with various protonic acids [9,10]. A common and well-known inorganic acid namely hydrochloric acid (HCl) doped PANI is soluble in pyrrolidine and concentrated acids with improved electrical conductivity [11]. On the other hand, the organic acid that confers certain solubility and high conductivity on PANI is ±10-camphor sulphonic acid (CSA) [12] where doped in polyaniline presents a high conductivity ($>1 \times 10^7$ S·cm$^{-1}$) in the presence of m-cresol solvent. It provides high conductivity and crystallinity values to the PANI-CSA complex system, conferring on the polymer a metallic like-behavior [13]. In the recent years, HCl and CSA doped PANi have been used in Poly(methylmethacrylate) PMMA composites to produce highly conductive electrospun nanofibers by our group previously [14-16], in which the investigation devoted to microwave hall mobility and electrical properties. These results may open novel fundamental and applied frontiers [17] in

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diverse fields. Considering the potential applicability as conducting nanowires and rather deep understanding of blends as a whole can be realized when the detailed characteristics of conducting component (doped PANI) is known. At this stage, authors are motivated to excavate the electrical properties in doped and undoped PANI in “bulk” rather than nanowires. Doped PANI as a fundamental conducting species, its various properties are rather interesting and believed to be essential to understand.

Present article provides simultaneous comparison of two different acids doped PANI and their optical, photoluminescence, electrical (AC, DC) and dielectric properties.

2. Experimental

2.1. Materials and Methods

Polyaniline base (Mw = 65,000), Hydrochloric acid (HCl), ±10-camphor sulfonic acid (CSA), m-cresol, and N-Methyl-2-Pyrrolidone (NMP) were obtained from Sigma Aldrich. All the chemicals were used as received.

The complete doping procedure has been described in our previous publications [14,15]. As briefly, 1 g of PANI base added in 40 ml of HCl stirred at 0 to −5˚C for 4 - 5 h. The fine dispersed particles were separated by centrifugation at 10,000 rpm for 10 min and vacuum dried for 48 h, the obtained powder was ready for further use. Similarly, 1 g of PANI base, 1.32 g CSA and 30 ml m-cresol grounded together in homogenizer and stirred with initial heat 50˚C for about 45 min and dried for 1 day to get fine powder, was used further characterization.

Scanning electron microscope (SEM) images of powder samples was obtained from JEOL JSM-6701F SEM. UV-visible absorption spectrum was recorded from dispersion in NMP using Shimadzu UV-3600, UV-Vis-NIR spectrophotometer. Photoluminescence was carried out using (Fluorolog, Horiba Jobin Yvon) axial S channel and 300 nm of wavelength was used for excitation. FTIR spectrum was recorded using Nexus 670 spectrometer. The thermal properties were obtained from DTA-TGA, SDT-2960, TA Instruments, while samples were heated in a ceramic crucible at a rate of 10˚C/min. The electrical leads were attained from the pellets with Ag paint. DC electrical conductivity (σDC) of PANI base is measured under two probe method, whereas PANI (HCl) and PANI (CSA) were measured under four probe configuration. In the former case, voltage is applied (lab-built voltage source) across the pellet and the current is measured (Keithley, 6514 system electrometer). Then σDC (=d/RA) was calculated from the measured values of resistance (R), cross sectional area (A) and thickness (d) of the pellet. Whereas in the latter case, the current (Keithley 6221) was applied at two probes and voltage (Agilent 34401A) was measured across the remaining two probes, and hence the conductivity (σ = ln2/πdR) is estimated, where symbols carry the previous meanings. By using a low temperature cryostat, temperature dependent σDC is recorded. The AC response of the pellets was obtained from Schlumberger Solartron SI 1260 electrochemical impedance spectroscopy in 1 Hz to 12 MHz range which is pre-calibrated with standard resistor and capacitor.

2.2. Characterization

Typical SEM images are shown in Figure 1 and the average particle sizes (using UTHSCSA Image tool 3.00 program) were determined to be about 100, 170, and 250 nm for PANI base, PANI (HCl), and PANI (CSA) respectively.

It is rather hard to distinguish the individual particles or grains from the images. However, Danming et al. [18], have noticed no significant effect of HCl doping on the particle size. The particle size is left undetermined in the present context as it is appropriate not to comment on the average size of the particles, nonetheless the morphology is more or less similar to those previously reported [19].

UV-visible absorption spectra are shown in Figure 2. Two distinct peaks are observed at two different wavelengths, centered at ~3.75 and ~2.0 eV which are characteristics of PANI. The first peak (~3.75 eV) indicates the π-π* transition in the benzenoid rings, which is almost unaffected upon doping. This band also relates to the extent of conjugation between adjacent phenyl rings in the polymer chain. The second band (~2.0 eV) is due to the exciton absorption of the quinoid rings and which are attributed to the inter chain or intra chain charge transport [20]. This second absorption band for pristine PANI is at ~1.81 eV which is blue-shifted upon CSA doping (to ~2.09 eV), in contrast, for HCl doping, though it is not very explicit band, it can be inferred that it is certainly

![Figure 1. SEM images of (a) PANI base (b) PANI (HCl) and (c) PANI (CSA).](image-url)
red-shifted (to ~1.57 eV). The blue shift upon doping essentially indicates the increased oxidation state and hence better conductivity. Thus it is expected that CSA doping should possess higher conductivity than HCl doping. Also sulphonic acid units may interact with amine/imine hydrogens, which enhances the electrical properties of polymers [19,21]. However, red-shift indicates the improved emeraldine structure, consequently further improvement in the degree of doping. Similar set of implications can be seen in the literature [22] while PANI is doped with MnO₂.

PL spectra were measured for all the three samples in the range of 300 - 800 nm and shown in Figure 3. The wavelength of excitation chosen for all the samples is 300 nm. This is because of π-π* transition of the benzenoid unit is responsible for PL in PANI [23]. Three excitation peaks were observed for all the samples at 322.5, 581.4 and 644.2 nm. PANI (CSA) shows the highest PL intensity followed by PANI (HCl) and PANI base, respectively. PANI (CSA) shows better PL intensity, the possible reason was benzenoid and quinoid units are more orderly arranged in it which favors the formation of excitons and increase in delocalization length of a singlet exciton [23,24]. Hence, one should expect higher photoluminescence emission from PANI (CSA) which is having higher extent of π conjugation. PANI (HCl) shows two extra emission peaks at 425.9 and 451.0 nm compared to PANI (CSA) and PANI base, the possible reason may be the presence of lighter dopant ions (Cl⁻) in PANI (HCl) compared to with higher charge mobility compared to the heavier dopant ions (SO₃⁻) in PANI (CSA).

FTIR studies are known to be useful to identify chemical bonds, functional groups and the components of unknown sample mixtures. In the spectra every wavelength of light absorbed is characteristic of a specific chemical bond. The spectra are shown in Figure 4, in all the samples, the peak present at 1559 and 1484 cm⁻¹ is due to the C-C stretching of quinoid and benzenoid rings respectively. This unaffected band upon doping indicates that the aromatic ring is retained in the polymer, which is corroborated by UV-vis in which the exciton band is almost unvaried with regards to the position of the peak. The bands at 1300 and 1240 cm⁻¹ are due to the C-N stretching of the benzenoid ring. The peak observed at 1139 cm⁻¹ is similar to those reported earlier and is associated with vibration mode of N=Q=N which is an electronic band (Q refers to the quinonic type rings). The presence of a band at 805 cm⁻¹ is attributed to the out of plane deformation of C-H in benzene ring. A small peak observed in case of PANI (CSA) at 805 cm⁻¹ corresponds to sulphonic acid group and the observed peak positions are consistent with the literature [19].

Thermograms from the samples are depicted in Figure 5. The derivative of the weight loss versus temperature (not shown here) is employed to identify the distinct peaks during heating, which are assignable to the decomposition of particular constituents in the composite. In Figure 5, a small fraction of weight loss at less than 100°C is observed in all samples, which is due to the evaporation or expulsion of the absorbed water. As such it is hard to remove the moisture from the samples completely during the synthesis as interpreted by Feng and Xue [25] in the case of PANI in water-oil emulsion.
The second weight loss observed nearly at 255°C for PANI (HCl) and is attributed to the evaporation of the dopant. This intermediate rather small decomposition step is observed in the case of PANI (CSA), which starts at 309°C and stretches till 335°C, further it decomposes rapidly. Although an intermediate degradation step can be seen more explicitly for HCl doping around 497°C, it merges with that of pure PANI as that of CSA doping at ~515°C, which is the final decomposition associated with polymer backbone, PANI. Thus the doped samples depicted slightly lower decomposition temperatures corresponding to a lower thermal stability unlike the case of plasticized PANI (HCl), where higher thermal stability can be expected [26].

3. Results and Discussion

3.1. DC Electrical Transportation

$I^V$ curves are shown in Figure 6 and corresponding conductivities are tabulated in Table 1. From Figure 6, linear nature of $I^V$ curve close to zero bias indicates that the charge transfer is a straightforward consequence of ohmic behavior (without any breakdown) associated with hopping mechanism [27], indicating that the measurement was performed in a field independent conductivity region. Hence the electrical conduction is through electrons, and trapped ions, for example, H$^+$ and Cl$^-$ ions in the case of HCl doped PANI. The conductivities of the doped samples have depicted an improvement of ~10 orders in magnitude when compared with that of pristine PANI [8, 28], also consistent with those earlier reported [19, 29]. This improvement in the conductivity is originated from the additionally generated extended states and charged defects with electronic structures, which are evident in various bands from UV-vis, and PL spectra. Additionally, a significant increase in conductivity with acid dopants is also observed in other composite systems such as PANI (HCl)/Polyvinyl alcohol and PANI (CSA)/PMMA [30-32]. CSA doped PANI has shown highest conductivity among the other samples of ~95.8 S/cm, may be due to its additional feature of CSA where an interaction between the sulphonic acid groups with amine/imine hydrogens can be expected [33].

Since $I^V$ characteristics are not stand-alone to explain the conduction process, or the entire electric nature of sample, we have carried out temperature dependent conductivity and plotted the variations in Figure 7 (ln $\sigma_{DC}$ vs $1/T^{1/4}$). In the intrinsic region close to the room temperature (RT), the conductivity curves are not simply straight lines. Two regions of different slopes for PANI (HCl) are observed in contrast to the other two samples. This transition of such gradient is rather slow till ~200 K, further $\sigma_{DC}$ decreases with the temperature more rapidly. Below 200 K the behavior of the different samples is not very distinct, except CSA doped sample tails off slightly at ~130 K. This implies that at ~200 K, we can see a transition between the conduction processes explicitly for HCl doped PANI (Figure 7(b)). Above 200 K, the conduction is associated with polaron transport in the extended state.
At lower temperatures, the probability of thermal release becomes rapidly smaller, so that at a temperature below 200 K, it is more likely that polaron hops to a neighbouring localized state. Therefore at RT, these materials show semiconducting nature. In such case, if the conduction mechanism is due to the carriers excited beyond the mobility edge into non-localized or extended states, the DC conductivity can be expressed \[34\] as Equation (1):

\[
\sigma_{\text{DC}}(T) = \sigma_0 \exp\left(-\frac{E_a}{k_B T}\right)^{1/n}
\]

where \(n\) represents the dimensionality (\(n = 2\) for 1D, \(n = 3\) for 2D, and \(n = 4\) for 3D) of the variable range hopping conduction. Further, the pre-exponential factor \(\sigma_0\) is the conductivity at RT, \(E_a\) is the activation energy, which is temperature independent, \(k_B\) is Boltzmann constant, and \(T\) is absolute temperature.

The temperature dependence of DC conductivity indicates that the observed semiconducting nature is due to only a small fraction of delocalized carriers. Using Equation (1), activation energies are determined for three samples (see Table 1). Surprisingly, the lowest is yielded for PANI (CSA) and the highest for PANI base. These activation energy values suggested that the hopping conduction is the prominent conduction mechanism in these types of systems. Hopping mechanism requires low activation energy with higher dielectric permittivity. Due to this \(E_a\), electrical conductivity is expected to be weakly temperature dependent around 300 K \[35\]. In this context, it is important to mention that the linearity in the graphs (Figure 7) may indicate 3D hopping mechanism \[34\] nevertheless it is also explained based on percolation theory \[36\].

### 3.2. AC Electrical Studies

Impedance spectroscopy results usually involve capacitance and rarely involve inductance and it has become customary to plot the real and imaginary parts of the impedances as shown in Figure 8. Figure 8(a) depicts a typical depressed semicircle for PANI base and fitting of RC circuit is shown in Figure 8(b). The fittings for PANI (HCl) and PANI (CSA) are not shown here. Nyquist plots for PANI (HCl) and PANI (CSA) are shown in Figures 8(c) and (d), respectively. The high intercepts \[37\] of Nyquist plots are equivalent to DC resistance (at low frequencies), using these values the estimated conductivity \(\sigma_0\) values are tabulated in Table 1.

#### Table 1. Calculated DC, AC conductivities (S/cm) and activation energies \(E_a\) (meV), and \(\omega_c\) in rad/sec.

<table>
<thead>
<tr>
<th>Sample</th>
<th>(\sigma_{\text{DC}})</th>
<th>(E_a)</th>
<th>(\sigma_{\text{AC}})</th>
<th>(\sigma_0)</th>
<th>(s_1)</th>
<th>(s_2)</th>
<th>(\omega_c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PANI base</td>
<td>(9.5 \times 10^{-8})</td>
<td>98.16</td>
<td>(-13.8 \times 10^{-8})</td>
<td>(2.73 \times 10^{-7})</td>
<td>0.92 ± 0.02</td>
<td>-</td>
<td>(-250)</td>
</tr>
<tr>
<td>PANI (HCl)</td>
<td>(-1.8)</td>
<td>74.40</td>
<td>(-2.9)</td>
<td>0.268</td>
<td>0.44 ± 0.02</td>
<td>-</td>
<td>(-2 \times 10^6)</td>
</tr>
<tr>
<td>PANI (CSA)</td>
<td>(-95.8)</td>
<td>57.24</td>
<td>(-240)</td>
<td>(2.94 \times 10^3)</td>
<td>0.12 ± 0.002</td>
<td>0.04 ± 0.002</td>
<td>(-25 \times 10^3)</td>
</tr>
</tbody>
</table>

Figure 8. Nyquist plot of (a) PANI base (b) Z-view fitting for PANI base (c) PANI (HCl) and (d) PANI (CSA).
Figure 9, shows log-log plot of AC conductivity ($\sigma_{AC}$) against frequency ($\omega$) for the three samples, which have shown two regions of diverse nature. Above a characteristic frequency, $\omega_c (=2\pi f_c)$, (DC-AC crossover) the conductivity increases with frequency. The frequency independent part $\sigma_0$ increases strongly with different dopings, which is offsetting the critical frequency $\omega_c$ to higher values [38]; where, $\omega_c$ is determined on the basis of a description provided by Kilbride et al. [39]. The highly disordered structure of PANI is well known that it is bulk conductivity, when protonated, has electronic character. Further, we can see from the UV-vis spectra, the various allowed transitions of electrons among different states. It can also be noted that the doped samples have shown significantly different absorption curves from the undoped sample, i.e., additional electronic states have been formed due to doping. Due to this, the inter-chain distances may be reduced. A shorter inter-chain distance is beneficial for the inter-chain hopping of charge carriers, bringing out a much higher electrical conductivity which is the case with HCl & CSA doped PANI.

The observed AC conductivity follows universal power law [40] of the following form

$$\sigma_{AC}(\omega) = \sigma_0 + a\omega^\beta$$

Above $\omega_c$, PANI base and HCl doped PANI are governed by one critical exponent, whereas CSA doped PANI has shown two exponent (see red and blue lines in Figure 9, Table 1 for critical exponents). The influence of doping can be seen explicitly on the transition frequency ($\omega_c$). For instance, CSA doping, it has occurred around 4 kHz, whereas for HCl doping it shifted towards much higher frequencies.

HCl doping has shown rather interesting feature which is in clear contrast to that of base and CSA dopings, such as the $\sigma_0$, which is constant for a wide range of frequencies (1 Hz to ~16 kHz) and rises sharply producing highest critical exponent ($0.44 \pm 0.02$) among other samples. While, $\sigma_{AC}$ of other two samples (PANI base and CSA doped) also increase with frequency after reaching certain value of frequency resulting in a symmetric hopping model [40,41]. This is because of localized charge carrier motion makes it possible to take maximum advantage of the well conducting regions, while at lower frequencies charge transport must extend over longer distances and is limited by poorly conducting regions. Since CSA is heavier molecule than HCl, the later can align much efficiently and easily to the electric field in high frequencies [42]. This can be noticed in Figure 9 in which the capacitive contribution to $\sigma_{AC}$ from CSA doping appeared in much lower frequency than that of HCl doping. Upon further increase in frequency, CSA doped sample has shown a trend which may be a saturation which is described by another exponent ($S_2$).

3.3. Dielectric Behaviour

The electric permittivity parameters ($\varepsilon'$ and $\varepsilon''$) obtained from the real and imaginary parts of impedances are shown in Figure 10 as a function of frequency. The values are seen to decrease with increasing frequency, associated with the polarizability mechanism. It is well known that the doped PANI system contains two types of charged species, one polaron/bipolaron system which is mobile and free to move along the chain, the others are bound charges (dipoles) which have only restricted mobility and account for strong polarization in the system. Above a certain frequency, the real permittivity ($\varepsilon'$) exhibits plateau region, this is may be due to space charge polarization and it reaches saturation due to further increase in the frequency [19,43]. This is due to the fact that the conductivity of the sample increased upon doping and acting as better dielectric systems in lower frequencies, which is indeed significant. Nevertheless upon crossing a critical frequency, $\varepsilon'$ is approaching zero.

This onset frequency is similar for PANI base and PANI (HCl) in contrast to that of CSA doped PANI which is slightly higher. Similar feature is noticed in the

![Figure 9. Frequency dependent conductivity for (a) PANI base, (b) HCl and (c) CSA doped PANI. $\omega_c$ indicates the DC-AC transition, where as different critical exponents are shown as $S_1$ and $S_2$ with fittings in red and blue color respectively.](image-url)
case of $\varepsilon''$ for all the samples. The decrease in dielectric loss is not symmetric for all the three samples on the logarithmic scale. The frequency of maximum loss shifts towards higher frequency with acid dopants. [44] As we can see from Figure 10(a), the dielectric constant of $(\varepsilon')$ of PANI base has a maximum of $2.5 \times 10^2$ F/cm, whereas for PANI (HCl) and PANI (CSA) (shown in Figures 10(b) and (c)) has increased ~5 and ~8 orders respectively. Moreover, the electric permittivity for the PANI (CSA) shows higher values compared to PANI (HCl) and PANI base, which is due to the local orientation of the individual dipole moment associated with the relatively large and rigid CSA dopant [44]. To analyze the electric conductivity relaxation process completely, the electric permittivity is converted into electric modulus [43], because the electric permittivity is not alone sufficient to explain in depth electrical properties of conjugated polymers.

The calculated real and imaginary parts of electric modulus are plotted in Figure 11. Real and imaginary modulus reaches zero at lower frequencies, indicating that the electrode polarization imposes negligible contribution to $M'$ which can be ignored. $M'$ is steady till 10 Hz, 20 kHz, and 400 kHz for PANI base, PANI (HCl), and PANI (CSA) respectively, beyond which it increases rapidly. From Figure 11(a), one can infer that there are unexpected peaks arise due to the cumulative dipole moments of the main chain [44]. The frequency dependence of the imaginary part of the modulus is also shown on the respective plots. The peaks observed at 0.5 MHz, 0.7 MHz and 4 MHz for PANI base, PANI (HCl), and PANI (CSA) respectively. It suggests that the increase in the dopant molecule size lead to appearance of $M''$ is shifting towards higher-frequency region, i.e., the DC conductivity has enhanced. $M'$ and $M''$ of PANI base and doped PANI are increased abruptly with increasing frequency. In the Figure 11(a), the $M'$ and $M''$ of PANI base
shows non monomodal distribution of the electric modulus. This may also be due to the complicated conduction process or various phases. But in PANI (HCl) and PANI (CSA), this multiple conduction process (Figures 11(b) and (c)) is not present in contrast to PANI (CSA) in which we have used m-cresol in the synthesis[43,44] the $M''$ increased more slowly. $M''$ variation with the frequency gives information about different conduction conditions such as hopping and dimensionality of the conduction process, which are depend on the conjugated state, chain structure, and doping level from different dopants. Therefore, a proper selection of dopant would supply suitable application of conductive polyaniline.

4. Conclusion

The presence of the dopants and un-effected aromatic ring of polyaniline were confirmed by UV-vis, FTIR respectively. Although there is no improvement in the thermal stability, the dopent disassociation from the matrix can be seen from the thermal analysis. The conductivity is enhanced upon doping due to generation of extended states in doped molecules or by changed defects with electronic structure. The structural and charge transfer processes are explained with FTIR and UV-vis spectroscopy. Intense PL is observed in CSA doped PANI followed by HCl doped PANI and PANI base. Henceforth these materials should also expect to show good electroluminescence (EL) as well. Thus CSA doped PANI is a promising material with respects of applications as emissive layers in organic or polymer light emitting diodes. Activation energy being an empirical parameter is employed to characterize the exponential temperature dependence of the change in the conductivity. The decrease in activation energy is noticed with acid dopants indicates that it would be a very good conducting material. Dielectric responses of polyaniline salt films, made from two different methods, were compared to investigate the effect of a dopant on electrical properties. Henceforth, we anticipate that these HCl and CSA doped PANI would be good candidate for electrical device fabrication.

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REFERENCES


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