ORIGINAL PAPER



A comparative study of the effect of organic dopant ions on the electrochemical and chemical synthesis of the conducting polymers polyaniline, poly(o-toluidine) and poly(o-methoxyaniline)

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Abstract

Electroactive polymers such as polyaniline and its derivatives have many potential applications, but the nature of the dopants can have an adverse effect on their properties. Polyaniline (PANI), poly(o-methoxyaniline) (POMA) and poly(o-toluidine) (POT) have been synthesised using appropriate organic acid dopants (2-naphthalenesulfonic acid and 1,5-naphthalenedisulfonic acid) via electrochemical and chemical polymerisation methods, where the later was achieved at pH (1-2) in the presence of ammonium persulfate as an oxidant. The effects of the monomers and organic dopants on the physiochemical properties and morphological surface of the polymers were studied by means of Fourier transform infrared spectroscopy, electrical conductivity measurements, thermogravimetric analysis and scanning electron microscopy (SEM). The above films were also synthesised electrochemically whilst varying conditions such as scan rate, scan number and in aqueous solution of organic acids using cyclic voltammetry techniques. Cyclic voltammetry experiments have been employed to analyse the growth behaviour and electrochemical stability of films in the background electrolytes (monomer-free). It was found that PANI salts displayed higher doping levels and conductivity while POMA and POT salts were found to show lower conductivity and doping levels, respectively. Furthermore the electrochemical response of films was different in the stability and this may be because of the methoxy (-OCH₂) and methyl (-CH₂) groups located at the ortho position of the aromatic ring structure of the monomers (o-methoxyaniline and o-toluidine). The -OCH₃ and-CH₃ groups have significant steric effects and can reduce to form long chains of polymers and also decrease conductivity. The SEM showed that the morphologies of the prepared conducting polymers were different, which was associated with the nature and size of the dopant and monomer present in the bath solution.

 $\textbf{Keywords} \ \ Conducting \ polymers \cdot Poly(o\text{-toluidine}) \cdot Poly(o\text{-methoxyaniline}) \cdot Dopant \ ions \cdot Chemical \ and \ electrochemical \ polymerisation$

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Introduction

Conducting polymers such as polyaniline and its derivatives (Qu et al. 2020), polythiophene (Bazzaoui et al. 1994), polypyrrole (Omastová and Mičušík 2012) and poly(3,4ethylenedioxythiophene) (Gruia et al. 2020) are used in a number of fields in industry such as corrosion protection of metals (Sazou and Deshpande 2017), conducting textiles (Tseghai et al. 2020), biosensors and gas sensors (Yin et al. 2006), electrocatalysis (Gustafson et al. 2014), electrochromic display devices (Tian et al. 2017), supercapacitors (Deshpande et al. 2013) and in rechargeable batteries (Hong et al. 2020). This is due to their excellent features such as low cost, environmental stability, light weight and good electrical properties. In addition, they can be prepared easily from their monomers by electrochemical or chemical polymerisation (Hillman et al. 2017; Ismail et al. 2019b). Although there are many applications for conducting polymers, especially in electromechanical and electronics applications, there are certain limitations to their use such as poor electrochemical stability, low conductivity, insolubility in common organic solvents and their infusibility. Thus, some electroactive polymer structures are difficult to fabricate via electrochemical or chemical methods (Hong et al. 2020). Polyaniline (PANI), and its substituted derivatives such as poly(o-methoxyaniline) (POMA), also known as poly(o-anisidine), and poly(o-toluidine) (POT) (the ring substituted with alkyl and alkoxy) are considered a significant family of conjugative polymers and they can be prepared by either chemical or electrochemical polymerisation methods (Mattoso and Bulhoes 1992; Wei et al. 1989). Generally, the chemical polymerisations of aniline, o-methoxyaniline and o-toluidine (Structures of monomers are shown in Fig. 1) are carried out in aqueous/organic or ionic liquid solutions using an oxidant and strong inorganic or organic doping acids (Huang and Wan 1999). By contrast, in the electrochemical method an electrochemical cell is needed, where a thin coating of polymer can be deposited on the surface of a substrate (Alabdullah et al. 2020; Ismail et al. 2019a). In both systems, the polymerisation of the

relevant conjugated monomer can be achieved, and charge carriers are produced utilising particular dopants. The conductivity of the resultant polymer occurs as a result of electron transfer through the conjugated structure. However, the choice of appropriate dopant and monomer are important factors in the production of a polymer with good electrical conductivity and processability.

In this regard, conducting polymers have been investigated by many researchers in order to optimise properties such as electrical conductivity and electrochemical response without adding fillers (Delvaux et al. 2000). The addition of certain substances directly to the polymer matrix or during the polymerisation process can cause changes to the electrical properties of the polymer. The influence of inorganic and organic acids as dopant ions on the electro/chemical growth, morphology, molecular structure and the optical and conductivity properties of polyaniline, poly(o-toluidine), poly(o-methoxyaniline) films has been reported (Borole et al. 2004). The type of dopant and thus technique used to prepare the polymer can also have a significant effect on the polymer's properties (Alesary et al. 2018). Therefore, the degree of polymerisation and type of dopant are very important factors as they can affect the charge carriers, morphologies and conductivity of the polymer, but this relies on the concentration of monomer, polymerisation technique and kind of dopant species used (Zhang et al. 2005). Thus, the stability of redox reactions is considered to be of particular significance to the polymerisation process (Chujo 2011).

Many researchers have attempted to enhance the quality of conductive polymers and to improve their electrochemical stabilities, but major challenges still remain in this field. Recent studies focus on the use of inorganic acids (hydrochloric acid (HCl), nitric acid (HNO $_3$), chloric acid, HClO $_3$ and sulphuric acid (H $_2$ SO $_4$)) as dopant materials in the preparation of conducting polymers (Borole et al. 2002). However, most polyaniline salts doped with small size dopants (i.e., inorganic acids) have low thermal stability, low solubility in conventional organic solvents and are hazardous materials. In order to overcome these drawbacks, other organic acid dopants can be used instead of HCI and $\rm H_2SO_4$ and this issue has received

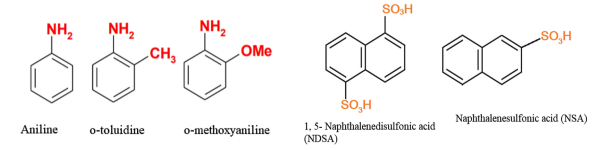


Fig. 1 Structures of monomers and organic dopant ions used in this work



less attention in contrast to the inorganic acids. Previous studies reported that the introduction of -SO₃H in the aniline ring can increase the conductivity and optimise the growth rate for polyaniline compared to other acids, where the reaction rate followed the order: $H_2SO_4 > H_3PO_4 \approx HF > HCI > HNO_3 > H$ $C1O_4 > HBF_4 > CF_3CO_2H$ (Borole et al. 2002). Therefore, in this study, 2-naphthalenesulfonic acid (NSA) and 1,5-naphthalenedisulfonic acid (NDSA) (the structures of the NDSA and NSA dopant ions are shown in Fig. 1) were selected as large organic sulphonic acid dopants having -SO₃H groups as the best anion dopant effecting the growth rate of polyaniline. In addition, the presence of the two sulfonic acid groups in NDSA helps achieve the doping for different polymer chains of the conducting polymer and creates a 3D network structure. In this way, the structure becomes more crystalline, which increases its conductivity, as reported in this paper. For the 2-naphthalenesulfonic acid, the dopant has one proton to dope the polyaniline single chain at one time.

The main purpose of this paper was to study the influence of certain organic sulphonic dopants on the conductivity, thermal stability, morphological characteristics and electrochemical behaviour of the PANI, POT and POMA polymers. Thus, the effect of the substituted methyl (–CH₃) and methoxy (–OCH₃) groups present at the ortho position in PANI on the physicochemical properties of polymer (mentioned above) can further be determined. To the best of our knowledge, this work has not been attempted in previous studies. Two different types of dopants have been used in the preparation of six conducting polymer compounds chemically. These compounds are PANI-NDSA, POT-NDSA, POMA-NDSA, PANI–NSA, POT-NSA and POMA-NSA. Fourier transform infrared (FTIR), thermogravimetric analysis (TGA) and scanning electron microscopy (SEM) were used to characterise the various species. The second part involved the preparation of PANI, POT and POMA doped with NDSA and NSA individually via electrochemical methods (cyclic voltammetry). The electrochemical stability of the polymers so prepared was characterised in their monomerfree organic dopant acids. The conductivities of the polymers have been measured for both methods (chemical and electrochemical). The polymer was deposited on the special Pt working electrode (diameter 1 cm); for more details, see the conductivity measurements in the results and discussion, and this is considered to be a key point in this study.

Experimental

Materials

Aniline, o-toluidine and o-methoxyaniline monomers were obtained from Sigma Aldrich (≥99%), Ammonium peroxydisulfate (APS) of analytical reagent grade was obtained from Fisher Chemicals (98%) and used as an oxidant for

chemical polymerisation. 1,5-naphthalenedisulfonic acid (NDSA) and 2-naphthalenesulfonic acid (NSA), used as dopants, were obtained from Sigma Aldrich at purities of 97% and 70%, respectively. All chemical materials and solvents were used as received without further purification.

Chemical polymerisation

Six samples of polymers were synthesised by separately dissolving 5 ml of each monomer (aniline, o-methoxyaniline and o-toluidine) in 82 ml of 1 M of each organic acid (NDSA and NSA) and stirring for 30 min in an ice bath. The ratio of 1 M organic acids to aniline was 0.75. The oxidant material was ammonium persulfate (APS), where 9.33 g of this oxidant was placed in the container and dissolved in 100 ml of distilled water. The molar ratio between the oxidant and each monomer was 4:3 (Alesary et al. 2018; Ismail et al. 2019b). The oxidant liquid (APS) was dropped into each of the monomer solutions made previously. The polymerisation process was performed in an ice bath at a temperature of 1–2 °C with stirring, and the pH of the bath solution was 1–2. The colour of the solution started to change to blue 15-20 min after adding the oxidant and then progressively became dark green. In all the experiments, the polymerisation process was performed under the same conditions of use and the reaction time was 4 h. The final compounds were filtered using a Buchner funnel, followed by washing with approximately 2 L of distilled water followed by acetone, and then dried in a vacuum oven at 50 °C.

Electrochemical polymerisation

The electrochemical polymerisation was performed by cyclic voltammetry (CV) in a cell with three electrodes using an Autolab PGSTAT20 potentiostat. A platinum (Pt) electrode (1 mm dia.) and Pt flag (surface area 2 cm²) were used as the working and counter electrodes, respectively, with an Ag/AgCl reference electrode made from an Ag wire placed in saturated KCl solution. The Pt working electrode was polished using 0.3-µm alumina paste prior to each experiment. The electropolymerisation was performed using a 0.2-M solution of each monomer (aniline, o-toluidine and o-methoxyaniline) separately in aqueous solution containing various organic acids (NDSA and NSA were used separately; see the legends of figures for each experiment) at 1 M concentrations. The electrode was scanned over a potential range from -0.2 to 1.1/1.2 V at a scan rate of 20 mV s⁻¹ for 10 cycles, with all experiments carried out at 25 ± 2 °C.

After the completion of electrochemical deposition in each experiment, the electrode was disconnected and removed from the monomer electrolyte. Then, each polymer



film was rinsed and cycled (10 cycles; ν 20 mV s⁻¹) in the same background electrolyte (monomer-free) at room temperature (25 ± 2 °C) for characterisation.

Instrumentation

Fourier Transform Infrared (FTIR) spectroscopy was performed using a Perkin Elmer Spectrum One Fourier transform infrared spectrophotometer over a frequency range of 500–4000 cm⁻¹ with the polymer dispersed in a KBr disk. The morphologies of the conducting polymers were characterised by scanning electron microscopy (SEM) using a FEI SIRION SEM instrument. SEM analyses were achieved under vacuum (10⁻⁵ Pa) with an accelerator voltage in the range of 5 to 10 keV due to the polymers being relatively soft materials. Thermogravimetric analysis (TGA) was used to measure the thermal stabilities of the conductive polymers powders in platinum crucibles via a Mettler Toledo 44 TGA/DSC 1 STARe system.

The conductivity of all polymers synthesised by chemical or electrochemical polymerisation was measured by a four-probe method using an APTFE cell. The dry powder polymer was pressed into discs (13 mm diameter) using a compressor (10 tonnes), and the polymer thicknesses measured using a digital micrometer. The conductivity was recorded using a Keithley195 current source and a Keithley197 electrometer controlled via a computer. Finally, the van der Pauw (Banaszczyk et al. 2010) Eq. (1) was used to calculate the conductivities of the polymers, as follows:

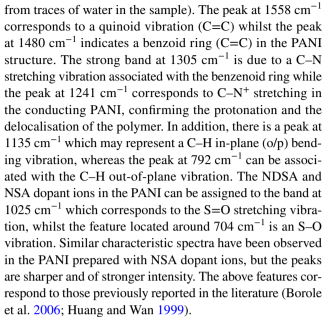
$$\sigma = \frac{\ln 2}{\pi R d} \tag{1}$$

where σ represents the conductivity of the polymer (in S/cm), R is the gradient of the current versus voltage graph and d is disc thickness of the polymer.

Results and discussion

FTIR spectroscopy of polymers doped with NDSA and NSA

FTIR spectroscopy was used to analyse polymeric samples that were synthesised from two types of organic dopant ions using 1,5-naphthalenedisulfonic acid (NDSA) and 2-naphthalenesulfonic acid (NSA). Polyaniline, poly(o-methoxyaniline) and poly(o-toluidine) were synthesised using chemical polymerisation in this study. Figure 2a shows the infrared spectra of the main vibrational features associated with PANI-NDSA and PANI-NSA. In the case of PANI-NDSA, there is a small peak at 3210 cm⁻¹ which can be attributed to an N-H stretching vibration while the band located at 3435 can be attributed to -OH groups (which could possibly be



It is worth noting that the FTIR spectra of the poly(otoluidine)-NDSA and poly(o-toluidine)-NSA showed more peaks and with higher intensities compared with the PANI conducting polymers. The same characteristic spectra were observed in both polymers (i.e., with NDSA or NSA), but there are small shifts in the locations of various peaks, as highlighted in detail in Fig. 2b. Figure 2b shows strong peaks at 3449 cm⁻¹ and 3045-3150 cm⁻¹, which can be attributed to -OH and N-H stretching vibrations, respectively. The bands at 2893 cm⁻¹ in POT-NDSA and at 2912 cm⁻¹ in POT-NSA were due to C-H stretching, the bands at 1549 cm⁻¹ were assigned to vibrations associated with quinoid rings (C=C), while the peak at 1498 cm⁻¹ can be attributed to benzoid ring-related vibrations. It should be noted that the peak at 1383 cm⁻¹ can be assigned to the methyl group adjacent to the benzene ring of o-toluidine (Yalcınkaya et al. 2010). The strong band located at 1095 cm⁻¹ corresponds to the biopolaronic structure (C-H out of plane o/p bending) in the conjugated POT-NSA, but this band is extremely weak in POT-NDSA and hence is not observed. The S=O stretching and S-O vibrations for NDSA and NSA dopant ions in the POT can be assigned to the bands observed at $1039-1029 \text{ cm}^{-1}$ and $759-667 \text{ cm}^{-1}$, respectively. These bands are in agreement with those of chemically synthesised poly(o-toluidine) (Elmansouri et al. 2007; Yalçınkaya et al. 2010).

The FT-IR spectra of poly(o-methoxyaniline) samples doped with NDSA and NSA in Fig. 2c showed similar peaks for both compounds with associated small shifts in their locations. The intensities of the bands in these samples are lower as compared to those found in the poly(o-toluidine) samples. The N-H stretching bond appeared at 3210–3212 cm⁻¹ as a small band. It is interesting to note from Fig. 2c that the intensity of the N-H peak is smaller



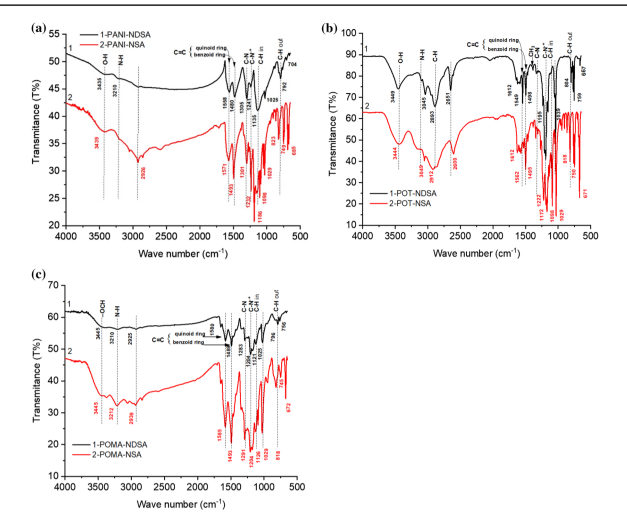


Fig. 2 FTIR spectra of polymers doped with NDSA and NSA chemically synthesised with $(NH_4)_2S_2O_8$ as polymerisation initiator: a PANI-NDSA (black line) and PANI-NSA (red line); b POT-NDSA

(black line) and POT-NSA (red line); ${\bf c}$ POMA-NDSA (black line) and POMA-NSA (red line)

than that observed in poly(o-toluidine) doped with NDSA/ NSA. This is because of the type of polymer used, which has an -OCH₃ group that can impede the growth reaction, and which seems to be associated with the N-H stretching vibration. A small band at 2925–2939 cm⁻¹ can be assigned to a C-H stretching bond associated with the NDSA and NSA dopants in POMA-NDSA/NSA. The peaks at 1588 and 1489 cm⁻¹ correspond to quinoid (C=C) and benzoid vibrations, respectively. The bands at 1283 cm⁻¹ and 1204 cm⁻¹ are due to C-N and C-N⁻⁺ stretching, respectively. The C-H out-of-plane (o/p bending) vibration appeared at 1174 cm⁻¹. There are two bands at 1121 cm⁻¹ and 1025 cm⁻¹ that could be attributed to the alkyl-O group in POMA. The mediumintensity band at 1121 cm⁻¹ suggests polaron bonding, whereas the small band at 791 cm⁻¹ is related the 1, 2 disubstituted terminal rings' o/p bending vibration. A weak band at 1655 cm⁻¹ was noted in POMA-NDSA and POMA-NSA that could be due to OCH₃, due to its partial oxidation in the electrolyte during preparation. The peaks mentioned above are in agreement with those previously reported in the literature (Christinelli et al. 2017; Mettai et al. 2018; Zhang et al. 2005).

Conductivity measurements

The measurement of the conductivities of polymers doped with NDSA and NSA dopant ions was one of the main purposes of this study, which depends on the type of polymer and dopant used. Table 1 shows the conductivities of polymers doped with NDSA and NSA by chemical and electrochemical methods. In the instance of electrochemically prepared polymer films, the polymer was deposited on the special Pt working electrode (diameter 1 cm). The counter electrode was a Ti mesh (30 mm×40 mm) coated with iridium oxide. The polymer powder was collected from the



surface of the electrode which, after drying, was used for numerous experiments and pressed into IR discs.

It is clear from Table 1 that the conductivities of polymers prepared by electrochemical polymerisation are greater than for their analogues prepared by the chemical method due to small fibrous chains in the uniform arrangement of the monomer structures which have an effect on the number of charge carriers and thus affect the conductivity (Delvaux et al. 2000; Li et al. 2011). The highest conductivity (0.65/0.98 S cm⁻¹) was obtained for PANI doped with NDSA, while the conductivities of POT-NDSA and POMA-NDSA were found to be 8.87×10^{-6} S cm⁻¹ and 5.21×10^{-4} S cm⁻¹, respectively. Generally, the conductivities of the PANI films in this work were less than PANI products that had been doped with inorganic acid (HCl, H₂SO₄) (Cao et al. 1989; Catedral et al. 2004). This is related to of the kind of dopant ions used in the polymerisation process, which have large molecular sizes compared to HCl or H₂SO₄ used in the previous work. The proton released from the dopant acid is considered to be of particular importance due to the fact that this proton has a significant influence on the level of protonation. NDSA and NSA dopant ions need a longer time to release protons compared to HCl or H₂SO₄ dopants, which is one of the reasons for the low conductivities of these polymers.

In this work, the conductivities of both poly (o-toluidine)-NDSA and poly(o-methoxyaniline)-NDSA were less than the conductivity of polyaniline-NDSA; this may be because of the methoxy (-OCH₃) and -CH₃ groups located at the ortho site of the aromatic ring structure of the monomers. These groups cause significant steric effects which can impede the polymerisation. This can lead to a decrease in the conductivity of such products compared to PANI (which does not have an alkyl-substituted group). Another possible reason for their low conductivity is the structure of the monomer used in the synthesis of the polymer (Cao et al. 1989). The structures of the monomers and dopant ions used are illustrated in Fig. 1. It is clear from Table 1 that the conductivities of POMA doped with NDSA or NSA are greater than that of POT doped with the same ions. This can be attributed to electronic effects (inductive effect and conjugative effect) and steric effects due to–OCH₃ and–CH₃. It is well known that the electronic effect of the methyl group is weak; however, its steric effect is stronger than that of the methoxy group due to the ease of rotation of the O–C bond in methoxy which reduces the steric effect. Therefore, the strong steric hindrance of CH₃ in POT results in more product crosslinking, shorter chains and lower doping levels compared to POMA. Previous studies have confirmed that the conductivity of the polymer is decreased with decreasing length of chain due to the reduction in electron movement along the chain (Jamal et al. 2008). The effects described could be the reasons why POT shows a lower conductivity than POMA when they are doped with identical dopant ions.

In the case of the conductivities of polymers doped with NSA, the conductivity of PANI-NSA was 0.033 S.cm⁻¹ while the conductivities of POT-NSA and POMA-NSA were 2.78×10⁻⁶ S cm⁻¹ and 1.3×10⁻⁵ S cm⁻¹, respectively. Accordingly, we can see that conductivities of the products of PANI, POT and POMA doped with NSA are less than those of the same products when doped with NDSA, despite the size of the NSA dopant ion being smaller than that of NDSA. This is because the NSA dopant reacts weakly with the monomers during polymerisation and needs a longer period to release protons. Therefore, this could be a reason for the decreased effectiveness of the electron transfer process in the POT film.

Thermogravimetric analysis

Thermogravimetric analysis (TGA) monitors the change in weight of a polymer as function of temperature change. This is included in the investigation of the polymer decomposition and to determine the amount of water loss as well as other volatile substances (Aphesteguy and Jacobo 2007). In this work, the TGA technique was used to identify the thermal stabilities of the polymer samples synthesised from different monomers and organic dopant ions. Polyaniline, poly(o-methoxyaniline) and poly(o-toluidine) were synthesised using chemical polymerisation in this study. Figure 3 shows TGA thermograms of polyaniline, poly(o-toluidine) and poly(o-methoxyaniline) doped individually with NDSA

Table 1 Conductivities of polymers doped with NDSA and NSA at room temperature (25±2°C)

Polymers	Dopant ion	$\sigma_{\rm RT}$ (S cm ⁻¹) measurement chemical method	$\sigma_{\rm RT}~({\rm S~cm^{-1}})$ measurement electrochemical method
Polyaniline	NDSA	6.50×10^{-1}	9.82×10^{-1}
Polytoluidine	NDSA	8.87×10^{-6}	1.21×10^{-4}
Polyanisidine	NDSA	5.21×10^{-4}	5.86×10^{-3}
Polyaniline	NAS	3.30×10^{-2}	1.2×10^{-1}
Polytoluidine	NAS	2.78×10^{-6}	6.83×10^{-5}
Polyanisidine	NAS	1.30×10^{-5}	5.13×10^{-4}



and NSA. Three weight loss steps were observed in the PANI-NDSA thermogram. The first step was due to organic solvent loss, which started at 52 °C; an organic solvent, acetone), was used to rinse the polymers, whereas the second step is attributed to dopant ion evaporation at 188 °C. The last step, which occurred at 366 °C, is associated with the degradation of the polymer itself. The weight loss steps for PANI-NSA were found to occur at 90 °C, 243 °C and 452 °C due to the removal of moisture, dopant anion evaporation and degradation of PANI-NSA, respectively. POT products also showed three weight loss stages as a function of temperature, as described in Table 2.

However, the TGA thermograms of poly(o-methoxyaniline) in Fig. 2 showed just two stages of weight loss. The first stage of weight loss appeared at 122 °C, which is caused by the removal of the dopant (NDSA/NSA). The second stage is related to the polymer degradation, which occurs at 323.97 °C. It is noted that the absence of water in these polymers could be because the polymer was dried at 50 °C in the oven; further details about the degree of weight loss for polymers are shown in Table 2.

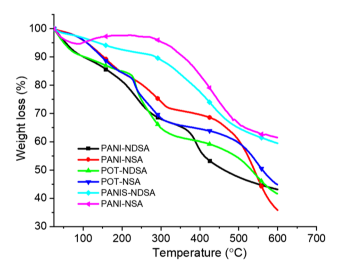


Fig. 3 TGA thermogram of polymers doped with NDSA and NSA chemically synthesised with $(NH_4)_2S_2O_8$ as the polymerisation initiator

chemically synthesised with $(NH_4)_2S_2O_8$ as the polymerisation initiator

Morphological characterisation

Figure 4 shows the morphologies of polyaniline, poly(otoluidine) and poly(o-methoxyaniline) doped with NDSA/ NSA as produced via chemical synthesis (use of (NH₄)₂S₂O₈ oxidising agent). From Fig. 4a, it is clear that the morphology of PANI-NDSA has two irregular-shaped regions consisting of fibrous particles and small agglomerated particles, while the morphology of POMA-NDSA (Fig. 4c) shows a larger aggregation of particles than that obtained in PANI films, producing a large lumpy morphology. The morphology of POT-NDSA and POT-NSA, as illustrated in Fig. 4b and e, respectively, resemble large plates but are uniform structures with no noticeable changes of shape observed between them. However, the surface morphologies of PANI and POMA are completely changed when doped with NSA, as shown in Fig. 4d, and f, respectively. This confirmed that the doping agents are of great significance with regard to modification of the morphology in terms of the resultant surface and size of composites (Alesary et al. 2018). We can see that the morphology of PANI-NSA, unlike that seen for PANI-NDSA, consists of one region of agglomerated particles, while there are two regions in the morphology of POMA doped with NSA, the first region being agglomerated with very small amorphous particles whilst the second does not have a clear surface and is coarse. It can be concluded that changing the monomer and dopant ion has an important influence on the morphology of the resultant polymer salts (Alesary et al. 2018; Yalçınkaya et al. 2010).

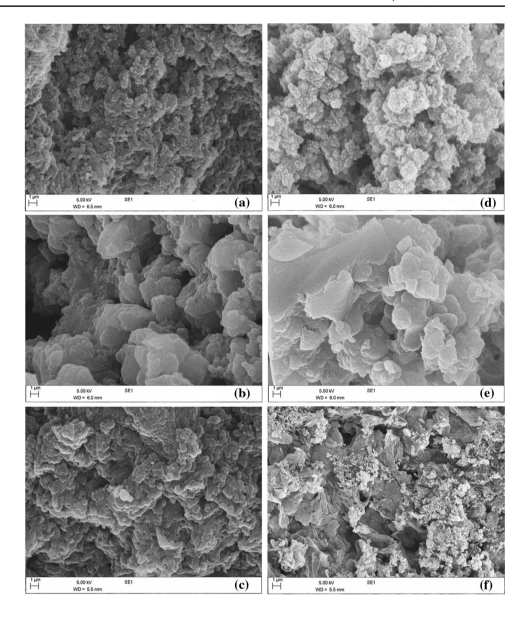
The morphologies of all polymer salts show obvious changes due to the use of electrochemical polymerisation, as shown in Fig. 5. In this method, the shape of PANI-NDSA resembles lines or homogenous chains and it seems that PANI-NDSA has a large porous surface compared to that seen for PANI-NSA. This could explain why the conductivity of PANI with NDSA/NSA dopant ions as synthesised by electrochemical polymerisation is greater than that of polyaniline produced by chemical polymerisation with

Table 2 Stages of weight loss for the polymers

Conducting polymers	Temperature loss of water/solvent (°C)	Temperature loss of dopant anions (°C)	Temperature backbone degradation (°C)
PANI-NDSA	52	188	368
PANI-NSA	90	243	452
POT-NDSA	40–70	225	426
POT-NSA	75	220	459
PANIS-NDSA	122	_	327
PANIS-NSA	90	_	355



Fig. 4 SEM photographs for polymers produced by chemical polymerisation under the conditions described in the experimental section; a PANI-NDSA, b POT-NDSA, c POMA-NDSA, d PANI-NSA, e POT-NSA and f POMA-NSA



dopant organic ions (NDSA and NSA), as shown in Table 1. The morphology of POT-NDSA (Fig. 5b) shows a uniform morphology having smaller, more globular plates than POT-NSA (Fig. 5e). The size of the particles in both is much smaller than was observed for POT-NDSA/NSA polymerised by the chemical method. The surface morphology of the POMA electrochemically produced with an NDSA dopant contained two regions which resemble chains agglomerated together and a lumpy structure, while POMA-NSA (Fig. 5f) shows a coarser surface than in POMA-NDSA and a different shape to the analogous polymer in Fig. 4f. The results of this study (Figs. 4, 5) confirm the presence of a significant difference in morphology of these polymers depending on the monomer, dopant ions and method of polymerisation used.

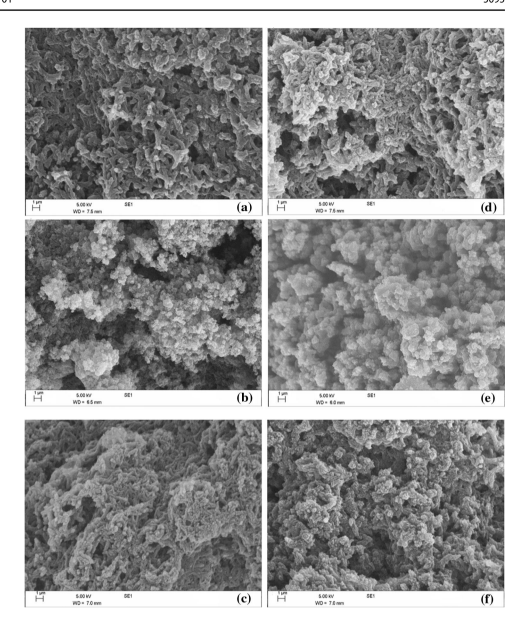
Cyclic voltammetry studies

Electropolymerisation of monomers

The electropolymerisation of different conjugate monomers were carried out on the Pt working electrode (1 mm dia.) in 1 M organic acid dopant (NDSA and NSA separately) solutions containing 0.2 M aniline, o-toluidine and o-methoxyaniline separately at a scan rate 20 mV s⁻¹ for 10 cycles using cyclic voltammetry (CV); the potential range used for PANI and POT films doped with NDSA/NSA was between - 0.2 and 1.1 V, while that for POMA-NDSA/NSA films was between - 0.2 and 1.2 V. All experiments were performed at room temperature (25 \pm 2 °C. The potential range used for PANI and POT films was - 0.2 to 1.1 V, while for



Fig. 5 SEM photographs for polymers produced by electrochemical polymerisation under the conditions described in Fig. 6; a PANI-NDSA, b POT-NDSA, c POMA-NDSA, d PANI-NSA, e POT-NSA and f POMA-NSA



the POMA film was -0.2 to 1.2 V. The reason for this is that the potential window for any conducting polymer varies depending on the structure of the deposited polymer. Further, it is related to the potentials at which nucleation and the oxidation of the monomer occur, the bounds of which define what is referred to as a nucleation loop. For PANI and POT films, the nucleation loop in the first cycle was at 1.0 V while for POMA was at 1.09 V, as shown in Fig. 6. POMA does not grow when the positive potential is less than 1.1 V under such conditions, and for this reason the potential range used for POMA was -0.2 to 1.2 V, in contrast to the PANI and POT films (-0.2 to 1.1 V). The nucleation loop vanished in the following cycles because of the persistent growth of the film preventing further nucleation in subsequent scans.

In all experiments, the growth of polymer films starts to develop significantly, resulting in an increase in the current peaks with an increasing number of cycles, as shown in Fig. 7. This implies that the charge and amount of polymer on the working electrode are increased with increasing cycle number (Nie et al. 2008). The results show that the electroactive behaviour of the PANI films doped with NDSA/NSA is greater than that for POT and POMA salts using the same dopants, while lower current peaks were observed for POT films compared with other polymers. This may be due to the strong steric effect and reduced electronic transfer that occur because of the alkyl group on the ortho site of the benzene ring of the monomer (toluidine), reducing the strength of the bonds between chains of poly(o-toluidine) compared to the polyaniline (Leclerc et al. 1993; Mello et al. 1995). It may be noted from the results that the rate of growth for NDSA dopant ions is greater than that for NSA under the same conditions (i.e., in the presence of the same monomer). This



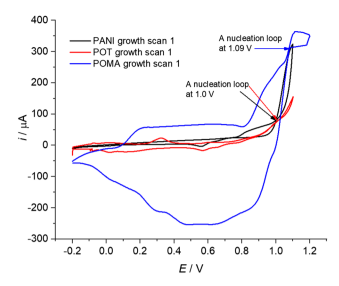


Fig. 6 The nucleation loop for the PANI, POT and POMA films for scan 1 of each of the polymers deposited in Fig. 7

suggests that the NDSA dopant ion tends to interact more strongly with the ion pairs of the anilinium radical cation (polaron) on the film chains compared to NSA.

Two main redox couples are observed in the CVs of PANI-NDSA/NSA, while more than four redox couples can be observed in the POT-NDSA/NSA and POMA-NDSA/NSA. In the PANI films, the first oxidation process (at ca. 0.33 V) corresponds to the leucoemeraldine base (fully reduced form) that converts to the protonated emeraldine (half oxidised form), while the second redox peaks (at ca. 0.94 V) are related to the protonated emeraldine converting to the fully oxidized form (pernigraniline structure), and the opposite for the reduction processes. The redox peaks in the intermediate region between 0.30 and 0.40 V may be associated with the crosslinked PANI salts, These results are analogous to those reported in the literature (Borole et al. 2004). In the cases of POMA and POT, there are three intermediate redox peaks, possibly indicating that the potential of redox peaks of the monomers is related to the type and size of the anion present and the increase in the number of redox peaks is strongly influenced by the size and nature of the species present in the electrolytic bath. It was observed that both the anodic and cathodic current peaks associated with the polymerization of anisidine and o-toluidine films are greatly influenced by the presence of organic sulphonic acids in the electrolytic bath.

The potential locations of the redox couples for POT-NDSA/NSA and POMA-NDSA/NSA polymers were significantly different to those of PANI-NDSA/NSA, as shown in Fig. 7a and d. The increased number of

potential peaks in the POT and POMA salts compared with PANI suggested that the electronic effect of the methyl substituent in POT and the methoxy substituent in POMA increased the extent of the crosslinking present. This could lead to a decrease in the doping levels and increase in the ratio of quinoid to benzenoid ring modes of POTs and POMA salts, producing a polymer with a low electrical conductivity, as reported in Table 1 where the quinoid ring in the PANI structure is more electronwithdrawing than the benzene ring, resulting in a reduction in the associated electrical conductivity (Abdiryim et al. 2005; Jamal et al. 2008). It is worth noting that the stability and reversibility of the PANI films and other films decreased in the electrolytes (monomer-free) with increasing thickness, i.e., with increasing scan number. Therefore, we deposited a thin layer of films in this study because the increased thickness of the deposited polymer is not electrochemically stable due to the reduced porosity of the polymer, which led to hindrance of the movement of cations and/or anions through the polymer network as described in our previous work (Ismail et al. 2019a).

The thicknesses (d) of all deposited polymers are reported in Table 3 and were estimated according to Eq. (2), whereas the charge (Q) in the coulomb (C) under the more cathodic peaks can be calculated from the integral of the cathodic current with time for the last scan of the deposited film. The thickness of the polymer can be measured based on the charge passed at the previous scan number.

$$d = \frac{QM}{2F\rho},\tag{2}$$

where Q is the value of overall charge for the electrodeposition polymer (see Table 3), ρ is the polymer density, M is the molar mass of the relevant monomer (in g/mol) and F is the Faraday constant (96,485 C/mol) (Ren et al. 2011; Shinde and Patil 2013). It is interesting to observe that the thickness of the polymers increases with the increase in the deposited charge, as shown in the Table 3.

From the results above, we can conclude that there are two important factors that can affect polymerisation rate, thickness and other properties of the polymer. First, the greatest polymerisation rate was found in NDSA compared to NSA under the appropriate conditions for the monomer used, strongly indicating it is affected by the nature of the dopant anion present in the electrolyte. Second, polymerisation rate can also be attributed to the type of monomer, where polymers formed from aniline in the presence of different dopant ions (NDSA or NSA) can have faster growth rates compared to the other polymers used in this study due to the absence of any steric effect from the methyl and methoxy groups.



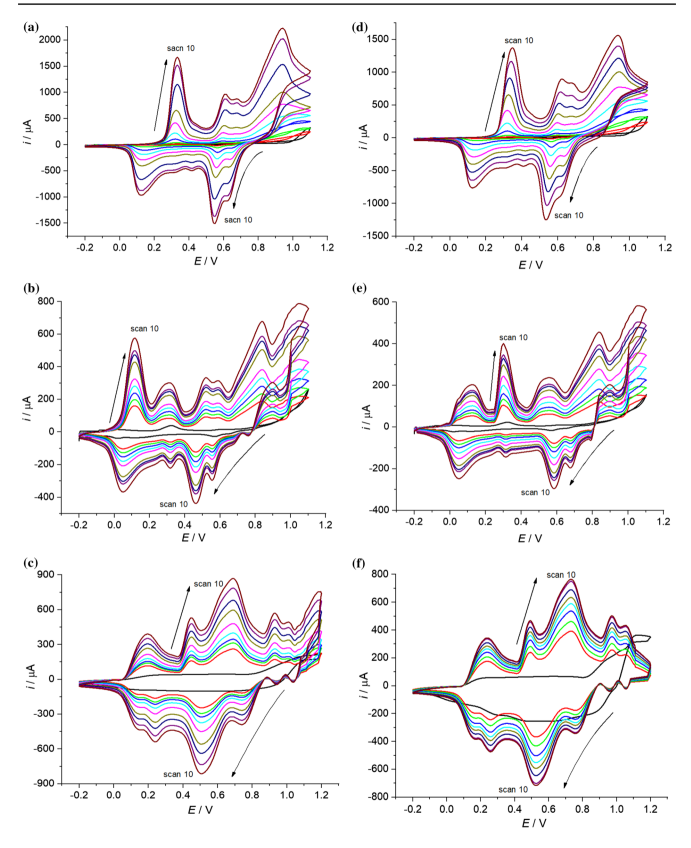


Fig. 7 The CV curves recorded during the electropolymerisation of the polymer films in 0.2 M monomer and 1.0 M organic acid (NDSA and NSA) for a potential window between -0.2 and 1.1/1.2 V at scan

rate of 20 mVs⁻¹ for 10 cycles; **a** PANI-NDSA, **b** POT-NDSA, **c** POMA-NDSA, **d** PANI-NSA, **e** POT-NSA and **f** POMA-NSA



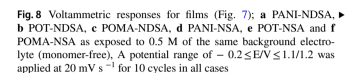
Table 3 Charge values and the thickness of polymer films deposited from different organic acidic electrolytes under otherwise identical conditions, as per Fig. 7

Samples	Q/C measured from cathodic current 10th cycle	Density of polymer (g cm ⁻¹)	Thickness of polymer (nm)
PANI-NDSA	0.02439	1.33	88.5
PANI-NSA	0.0208		75.5
POT-NDSA	0.00995	1.34 (Shinde and Patil 2013)	41.2
POT-NSA	0.00739		30.6
POMA-NDSA	0.02102	1.50 (Ren et al. 2011)	89.4
POMA-NSA	0.01853		78.8

Electrochemical response of polymers

After electrodeposition of the polymer film via cyclic voltammetry (Fig. 7), the redox behaviour and stability of the PANI, POT and POMA salts were investigated in similar background solutions (monomer-free) using cyclic voltammetry, as shown in Fig. 8. The concentration of the background solution was 0.5 M NDSA/NSA and the scan rate was 20 mV s⁻¹ in all experiments. The potential window used for PANI and POT films was between - 2.0 and 1.1 V while for POMA films was between -0.2 and 1.2 V. depending on the potential window of the deposited polymer. The current peaks of the films immersed in 0.5 M original organic acid electrolytes were close to the current peaks of the original films that were grown. As shown in Fig. 8a and d, two separate redox peaks can be observed in the electrochemical responses of the PANI films, and these are identical to the CV peak shape of PANI. This implies that electrochemical activity of PANI film is analogous to that of its parent PANI in both acid solutions (NDSA and NSA). The same, with regard to redox activity, was observed for the other polymers. In addition, the stabilities of all the polymers doped with the NDSA electrolyte was greater than those of the polymers doped with the NSA electrolyte. This is possibly due to polymer growth from NDSA being more conductive and the morphologies more porous than those observed with NSA.

It was found that the highest electrochemical stability corresponding to the redox peaks was observed for the POMANDSA film (60.30% charge redox retention) as exposed to 0.5 M NDSA, compared to the other polymers. On the other hand, the worst electrochemical stability was observed for the POT-NSA (30.41% charge redox retention) followed by POT-NDSA (43.31% charge redox retention) cycled in 0.5 M NSA and NDSA solutions, respectively. Herein, we observed a decrease in the peak currents as a function of increasing scan number; thus, there is a shift in the peak potentials and associated CV shapes were changed. This could be attributed to degradation of the film when switching to the overoxidation potential, which led to poor stability during redox cycling, as reported in previous studies (Bauermann and Bartlett 2005; Mažeikien and Malinauskas 2002).



These results show that the electrochemical stability of the polymer films in the electrolyte depends strongly on the type of polymer and also the nature of the anion present in solution. According to our experimental results, as reported in Table 4, the electrochemical stability of polymers prepared from different monomers and organic acidic dopants take the following order: POMA-NDSA > PANI-NDSA > POT-NDSA > POT-NDSA > POT-NSA.

Conclusion

In this paper, different conducting polymers have been successfully prepared from raw monomers (aniline, o-toluidine and o-methoxyaniline) by chemical and electrochemical polymerisation in the presence of 1,5-naphthalenedisulfonic acid (NDSA) and 2-naphthalenesulfoinc acid (NSA) dopant ions. The conductivities of POMA and POT doped with NDSA and NSA were found to be lower than the conductivities of PANI prepared using the same dopant ions. This is because of the type and level of protonation that were used in each case. The molecular sizes of o-toluidine (having a methyl group) and o-methoxyaniline (having a methoxy group) are nominally larger than that of aniline, resulting in a reduction in the growth of long chains of polymer and thus decreasing conductivity. The -OCH₃ and-CH₃ groups have significant steric effects and can reduce to form long chains of polymers and result in a decreased conductivity. It was found that the conductivity of polymers produced by electrochemical methods is greater than the analogous polymers synthesised via chemical polymerisation. The thermal stabilities of POT salts were higher than the PANI and POMA salts, where the degradation of POT salts was found to lie between 425 and 460 °C, while the degradation of PANI-NDSA and POMA salts was between 325 and 370 °C. SEM



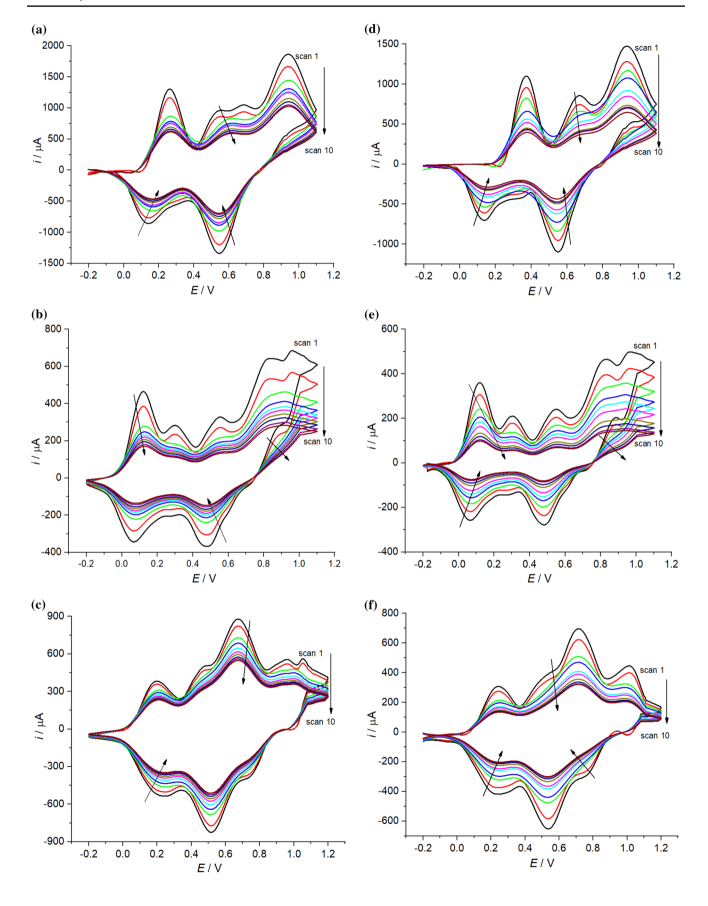




Table 4 Charge values for the electrochemical response for polymer growth in Fig. 7, as cycled in their background electrolytes (monomer-free) at a scan rate of $20~\text{mV}~\text{s}^{-1}$, as presented in Fig. 8. The

charge was calculated from the CV curves given in Fig. 8 by integration the current of the redox peaks with respect to time

Polymer films	Polymer cycled in electrolytes	Q/C redox 1st cycle	Q/C redox 10th cycle	% Q redox retention
PANI-NDSA	0.5 M NDSA	0.04342	0.02557	58.89
PANI-NSA	0.5 M NSA	0.03155	0.01521	48.21
POT-NDSA	0.5 M NDSA	0.01741	0.00754	43.31
POT-NSA	0.5 M NSA	0.01233	0.00375	30.41
POMA-NDSA	0.5 M NDSA	0.02552	0.01692	66.30
POMA-NDSA	0.5 M NSA	0.01912	0.00961	50.26

The charges found for cycles 1 and 10 are shown

shows a significant difference in the morphology of polymers depending on the monomer, dopant ions and method of polymerisation used.

The results show a rapid growth rate was observed in PANI films in the electrolytes containing NDSA and NSA under cyclic voltammetric conditions compared to POMA and POT, while it was found that the POMA-NDSA films had the highest electrochemical stability, recording ~ 66% retention of redox charge from the initial value. In contrast, POT salts showed lower electrochemical stabilities, recoding~30% of retention of redox charge from the initial value. We concluded that the methoxy and methyl groups in PANI derivatives (present at the ortho position in the phenylene ring) play a significant role in determining the physicochemical properties of the synthesised PANI derivatives. As a consequence, the methyl group was found to have a strong steric effect which led to POT salts becoming more oxidised than POMA salts. Thus, the nature of the anion present in the electrolyte and the type of method used strongly affect the subsequent polymerisation of the monomer.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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