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and complementary spectroscopic techniques (EDAX, Raman).



Controlled preparation of carbon nanotube-conducting polymer composites at the polarisable organic/water interface

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A R T I C L E I N F O

ABSTRACT

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1. Introduction

Intrinsically conducting polymers (CPs) have been the focus of great interest in many areas of science and technology during the last decades due to the attractive properties of these materials, i.e. the redox transformation can be used to induce conductance, spectral, mass, and volume changes [1,2]. These externally switchable changes are the basis of important applications of CPs such as electronic devices, sensors, smart windows, artificial muscles, and as membranes or modified electrodes [3-7]. The mechanical, thermal and electronic properties of CPs can be greatly enhanced by the formation of CP-carbon nanotube (CNT) composite materials: key factors are the method of preparation of the composite material, which in many cases can impart an useful anisotropic component to the physical properties of the composite [8,9]. One method of achieving such anisotropy is to grow the polymer materials around the aligned CNT "forest", where the latter is prepared via chemical vapour deposition (CVD) [10]. It has been shown that polymerisation can be advantageously achieved through electrochemical methods, where the CNT is used as the working electrode to produce an intimate contact between the carbon nanostructure and the polymeric deposit [4,6]. Single wall carbon nanotubes (SWCNTs) or their composites with CPs and metal nanoparticles can be characterised through Raman spectroscopy [11–14]. The most significant Raman bands of SWCNTs are the radial breathing mode (*RBM*), the tangential displacement mode (TG), which is also known as the G band, the

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disorder induced mode, which shows the density of the defects (D), and the high-frequency two-phonon mode (2D or G') [15].

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The electro-polymerisation of polypyrrole (PPy) at the interface between two immiscible electrolyte solutions

(ITIES) is reported. The approach is used to demonstrate the formation of a carbon nanotube (SWCNT)-

conducting polymer composite, by performing polymerisation in the presence of an assembly of SWCNT films.

The morphology of the SWCNT/PPy nanocomposites was determined using probe and electron microscopy

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There has been a recent upsurge in reports on the *in situ* assembly/ deposition of nanostructured materials (e.g. photoactive oxides, metallic nanostructures, and low-dimensional carbon species, such as SWCNTs and graphene (GR)) at the interface between two immiscible electrolyte solutions (ITIES) [16-20]. The ITIES is a special case of the generic liquid/liquid interface, where the presence of electrolyte in both phases provides external control over charge transfer, specifically ion, electron, and proton-coupled electron transfers [21-23]. Electropolymerisation at the ITIES has been reported: work on 2,2':5',2"terthiophene [24], was followed by studies of the influence of different polymerisation conditions and characterisation of this low oxidation potential thiophene derivative [25,26], and pyrrole derivatives (1methylpyrrole and 1-phenylpyrrole) [27]. The interfacial polymerisation of polypyrrole (PPy) and polyaniline (PANI) at the non-polarised, chloroform/water interface using ferric chloride (FeCl₃) as oxidant was reported in 2008 [28], and thereafter several papers described preparation of CPs at the liquid/liquid interface [29-33]. Numerous studies have also reported the formation of adsorbed carbon nanomaterial films via self-assembly, such as CNTs, and adsorbed films of few-layer GR or reduced graphene-oxide (rGO) at non-polarised, liquid/liquid interfaces, often with the specific aim of preparing transparent or modified carbon-coated electrode materials [34–37]. Recently we have extended this approach to the localisation and moreover the functionalisation of CVD GR at the ITIES [38,39]. The Zarbin group have demonstrated interfacial polymerisation using CNT or GR assembled at the non-polarised water/toluene interface to prepare carbon nanomaterial-PANI composites [40-42], mainly using a chemical oxidising agent, i.e. ammonium persulfate. The main advantages of the electro-polymerisation procedures, compared to the chemical route,

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consist of the possibility to control: the nucleation rate and growth by selecting the electro-polymerisation parameters; the thickness of the polymer films by the amount of charge passed during the deposition; and the CPs' morphology using suitable selection of an appropriate solvent and supporting electrolyte [7].

In this communication we report the interfacial electropolymerisation process for the controlled, potentiodynamic preparation of PPy films either at the bare ITIES or on the SWCNTs assembled interface. These free-standing PPy and SWCNT/PPy composite layers are transferable to solid substrates: these functional coatings are characterised using various microscopic and spectroscopic techniques.

2. Material and methods

Lithium chloride (LiCl, 99%); ammonium hexachloroiridate (IV) ($(NH_4)_2IrCl_6$, 99.99%); ammonium hexachloroiridate (III) monohydrate ($(NH_4)_3IrCl_6$, 99.99%); and the organic solvent 1,2-dichlorobenzene (DCB, 99%) were used as received, the pyrrole monomer (98%) was freshly distilled. The organic phase electrolyte bis(triphenylphosphoranylidene) tetrakis(pentafluorophenyl) borate (BTPPATPFB) was prepared as described elsewhere [43]. The potassium salt, KTPFB was obtained from Alfa Aesar, and the other chemicals were purchased from Sigma-Aldrich. Deionised water (18.2 M Ω cm resistivity), purified by a "PURELAB" Ultrafiltration unit (Elga Process Water), and was used for aqueous solution preparation.

Electrochemical experiments were performed using a four-electrode configuration [44] with an Autolab PGSTAT100 potentiostat (Metrohm-Autolab). The Galvani potential scale $(\Delta_o^w \Phi)$ was normalised in every case using the formal transfer potential of tetramethylammonium cation, which corresponds to $\Delta_o^w \Phi^\circ = 0.277 \text{ V} [45]$. The assembly of SWCNTs (synthesised by an arc discharge method, Sigma-Aldrich) at the organic/water interface is described elsewhere [44]. Briefly, sonication for 12 h using an ultrasonic bath (Elmasonic P60H) operating at 37 kHz, at 30% of its full power was used to prepare the SWCNT dispersion in DCB, the temperature was kept below 30 °C with a recirculating cooler system (Julabo F250) attached to the ultrasonic bath. Then, an aliquot of the dispersion with the organic electrolyte solution in DCB and an equal volume of aqueous phase were placed in contact and self-assembly of the SWCNT was induced using sonication (15 min). UV-Vis spectroscopy (DH-2000-BAL, Ocean Optics and USB2000 interface, Micropack GmbH) was used to measure the concentration of the SWCNT dispersions. Raman analysis (Renishaw RM 264 N94 (532 nm laser) spectrometer operating at power ≤ 0.3 mW), the SEM (Philips XL30 ESEM-FEG, operated at 10 kV) and the AFM (Bruker MultiMode 8, operated in "Peak Force" tapping mode with a silicon tip on a silicon nitride lever) characterisations were performed on the samples once transferred to a Si/SiO₂ wafer, following interfacial polymerisation, either for the PPy films or the SWCNT/PPy composite layers. The displayed errors are based on standard deviations from arithmetic averages of multiple measured values.

3. Results and discussion

3.1. Dynamic electro-polymerisation of PPy at ITIES

The successful interfacial electro-polymerisation of PPy film was achieved by a simple redox reaction between pyrrole (Py) oxidation to PPy (on the organic side of interface) and aqueous phase hexachloroiridate (IV) ($IrCl_6^2$), which is reduced to hexachloroiridate (III) ($IrCl_6^2$), Reaction 1).

$$IrCl_{6}^{2-}(aq) + Py(o) \rightarrow IrCl_{6}^{3-}(aq) + PPy(s) + H^{+}(aq)$$
(1)

The PPy films were prepared using dynamic electro-polymerisation, either *via* cyclic voltammetry at a 100 mV s⁻¹ scan rate or by potential step, from 0.1 M pyrrole (organic (o), DCB phase) and aqueous (aq)

phase containing 1 mM $IrCl_6^{2-}$ and 0.1 mM $IrCl_6^{3-}$ (composition of the cells given in Fig. 1). BTPPATPFB (10 mM) and LiCl (0.1 M) were used as (o) and (aq) phase supporting electrolytes, respectively. Four different dynamic interfacial polymerisation protocols were used: cyclic volt-ammograms (CVs) with 25 (Fig. 1A), 50 (Fig. 2B), 75 (Fig. 1B) cycles and the potential step method. The difference in the PPy films' and the blank CV's (blank = no pyrrole, Fig. 1 cell 1) electro-activity can be observed by comparing the last CVs of each polymerisation, *i.e.* the 25th (Fig. 1A-b) and 75th (Fig. 1B-b) cycles (Fig. 1 cell 2).

The main features of the PPy Raman spectra were assigned according to the literature [46] as follows (Fig. 1C, marked by asterisks): the ring deformation (933 cm^{-1}), the C—H in plane bending (984 cm^{-1}), the C—C backbone stretching (1318 cm^{-1}), the C=C backbone stretching (1562 cm^{-1}) vibration modes for the neutral form; and the C—H inplane bending (1048 cm⁻¹), N—H in-plane bending (1254 cm⁻¹), C—C backbone stretching (1333–1415 cm⁻¹), C=C backbone stretching $(1587-1587 \text{ cm}^{-1})$ for the cationic form, respectively, as the PPy films were partially doped both forms are present. In every case the transferred PPy films were shown to exhibit continuous coverage on the Si/SiO₂ substrate with the naked eye. The series of the SEM images of the transferred PPy films on Si/SiO₂ (Fig. 1D-F) show a trend in the extent of the coverage as a function of the number of the CV cycles, which were used for the electro-polymerisation of the PPy. Comparing the polymer films deposited using 25 (Fig. 1D), 50 (Fig. 1E), and 75 (Fig. 1F) cycles, the increase in coverage can be observed and moreover, a difference also can be seen in the morphology of the different PPy films. The initial deposits are globular, and consist of more discrete particles, whereas with further cycling the structure becomes more coherent, i.e. the PPy film in Fig. 1F is built up from flakes/layers, while applying the potential step method, more globular structure achieved.

3.2. Preparation of SWCNT/PPy composite at DCB/water interface

A schematic of the redox process and CVs of the electropolymerisation of PPy on the interfacial SWCNT film are shown in Fig. 2. During the reduction of the $IrCl_6^{2-}$ to $IrCl_6^{3-}$ on the aqueous side of the interface, the oxidation of the pyrrole occurred through the assembled SWCNT film at the interface, by analogy with similar redox reactions at the SWCNT assembled ITIES, e.g. Pd electrodeposition [44]. The optimised potentiodynamic polymerisation was used for the preparation of the single wall carbon nanotubes-based polypyrrole composite (hereafter, SWCNT/PPy), employing 50 CV cycles at a scan rate of 100 mV s⁻¹. The CV responses with and without the Py monomer are shown in Fig. 2B. The resultant SEM, AFM micrographs and Raman spectra of SWCNT/PPy and the comparison of the pristine (C) and PPy functionalised SWCNTs (F) are given in Fig. 2. The pristine SWCNTs formed bundles either on transfer to solid substrate or due to the presence of BTPPATPFB electrolyte (Fig. 2C), while in the case of the SWCNT/PPy, thinner or more individualised nanotubes are embedded in the PPy matrix (Fig. 2F), which can form a proper coating on the Si/SiO₂. This suggests that the drying process, rather than the removal of the electrolyte, is the cause of the bundling evident in Fig. 2C.

The AFM image of the composite (Fig. 2E) was used to estimate the thickness of the transferred SWCNT/PPy layer, found to be 31.8 \pm 0.3 nm. The diameter of the nanotubes in the case of the pristine SWCNT was 6.2 \pm 3.4 nm and the SWCNT/PPy was estimated to be in the range between 9.6 \pm 0.5 nm to 14.9 \pm 1.6 nm. The difference is attributed to the PPy "coating" deposited around the nanotubes between the pristine SWCNT and the composite ones. Fig. 2D-a and Fig. 2D-b, respectively, show the *RBM*, the two main components of the *G* band (*G*⁻ and *G*⁺), and the 2D band in Raman spectra of pristine SWCNT and SWCNT/PPy. The left inset in Fig. 2D presents the pristine (black) and composite (red) SWCNTs' spectra, due to the overlap of the *D* and *G* bands of the SWCNT and the significant peaks of the PPy, it is hard to detect all of the polymer peaks. Nevertheless, the ring deformation

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