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Carbonization of Polymers of Intrinsic Microporosity to Microporous Heterocarbon: Capacitive pH Measurements

Naiara Hernandez 1, Jesus Iniesta 1, Vicente Montiel Leguey 1, Robert Armstrong 2, Stuart H. Taylor 2, Elena Madrid 3, Yuanyang Rong 3, Rémi Castaing 3, Richard Malpass-Evans 4, Mariolino Carta 4, Neil B. McKeown 4, and Frank Marken*3

1 Departamento de Química física e Instituto Universitario de Electroquímica, Universidad de Alicante, Apartado 99, 03080 Alicante, Spain
2 School of Chemistry, Cardiff University, Main Building, Park Place, Cardiff CF10 3AT, UK
3 Department of Chemistry, University of Bath, Claverton Down, Bath BA2 7AY, UK
4 School of Chemistry, Joseph Black Building, David Brewster Road, University of Edinburgh, Edinburgh, Scotland EH9 3JF, UK

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Proofs to F. Marken (f.marken@bath.ac.uk)
Abstract

A nitrogen-containing polymer of intrinsic microporosity (PIM-EA-TB-H2; nitrogen adsorption surface area 846 m² g⁻¹) is vacuum carbonized at 700 °C and thereby directly converted into a microporous heterocarbon (cPIM; N₂ adsorption surface area 425 m² g⁻¹). Nitrogen functionalities in the polymer backbone are retained in the heterocarbon and appear responsible for unusual time-, electrolyte-, and pH-dependent properties. Electrochemical characterization suggests a high specific capacitance (typically 50 F g⁻¹) but only after prolonged immersion in aqueous HClO₄. The time-dependent increase in capacitance during immersion is assigned to slow hydration and ingress of HClO₄ into hydrophobic micropores (H₂SO₄ or H₃PO₄ are more hydrophilic and much less effective). Once hydrated, the microporous heterocarbon exhibits pH-dependent capacitance “switching” over a wide pH range and analytical applications as “capacitive” pH sensor are proposed.

Graphical Abstract:

Keywords: conformal carbonization; pH sensing; micropore hydration; voltammetry; heterocarbon
Introduction

Novel carbon and nano-carbon materials provide an important driving force for the development of new (electro-)analytical methodology and sensor devices [1]. Particularly exciting are the recent development of “nano-dot” carbons [2], functionalized nanoparticles [3], the use of grafting to provide surface modified carbon sensors [4], hyper-crosslinked porous carbons [5], and the pyrolytic formation of microporous carbon substrates for electroanalysis [6].

Mesoporosity is considered to be crucial in sensor design [7] but less accessible microporosity in the < 1nm range may also provide selectivity in particular for smaller analytes, e.g. for protons (vide infra). Pyrolytically produced carbons [8], tailored microporous carbons [9], and metal-organic framework-based microporous heterocarbons [10] are just some examples of a rapidly extending range of microporous heterocarbon materials now available to the analytical researcher. “Nitrogen-doped” heterocarbons and nanocarbons are of interest, for example, due to their catalytic properties [11] or in sensing applications [12]. In this report we explore the properties of a microporous nitrogen-containing heterocarbon obtained via vacuum carbonization of a novel polymer of intrinsic microporosity, PIM-EA-TB-H2 (see Figure 1).

The measurement of pH (or proton activity) is routinely based on potentiometric (e.g. glass membrane with a proton adsorption-dependent equilibrium potential) electrodes [13].
Although ubiquitous and fully established, these types of potentiometric pH measurements can be problematic when other adsorbing/surface active species are simultaneously present in the sample. This is particularly true in real biological samples, colloids, surfactant containing solutions etc. It has therefore been suggested that amperometric sensors (e.g. based on the limiting current response of a microelectrode [14] or on dual-plate microtrench electrodes [15]) offer advantages for proton concentration determination at least over limited proton concentration ranges. A new class of “reference-less” pH sensors based on the voltammetric response of pH-dependent surface redox systems has been developed by Compton and coworkers based for example on quinone functionalities grafted onto glassy carbon or graphite surfaces [16,17,18,19,20]. Other types of pH sensor concepts have been proposed, for example a pH-dependent impedance response was reported for aligned single walled carbon nanotubes [21]. Carbon composite materials are often employed in sensor development [22] and a review has appeared summarizing pH sensing carbon materials and composites [23]. For most voltammetric pH sensor systems the measurement of a voltammetric current response is linked to a potential read out (relative to an internal standard) to give the pH reading. The use of a purely capacitive current signal (reported here) based on the charging/discharging currents (although briefly mentioned previously for core-shell carbon nanoparticle systems [24]) appears to be new and potentially very robust (there is no need for potential control) and there are opportunities for simple nano-sensor device concepts.

Pyrolysis of polymer precursors provides versatile access to both non-porous and porous electrode materials [25]. High quality sensor films can be produced for example from thin
films of photo-resist polymer coatings [26,27]. Only recently has the carbonization of a polymer of intrinsic microporosity (PIM) been reported [28] to give interesting heterocarbon materials with both morphology and microporosity being retained during the carbonization process. This is in marked contrast to many other carbonization processes where post-treatments are applied to create/modify the desired microporosity. PIM materials [29] have emerged as a molecularly highly rigid polymer systems with the ability to absorb and separate gases [30], to sense gases [31], and they are currently under intense investigation for applications in electrochemistry [32].

The vacuum carbonization of PIM-EA-TB (a polymer of intrinsic microporosity –PIM- with ethano-anthracene –EA- structural components and Tröger base –TB- functionality [27]) at 500 °C was shown to yield electrically conducting carbon materials without change in shape, morphology, or in cumulative pore volume. It was recognized that the rigid molecular structure allows cross-linking with loss of some small molecular components, but without significant changes in polymer backbone and in morphology. In a follow-up study PtCl$_6^{2-}$ was adsorbed into the PIM-EA-TB precursor and in a “one-step” vacuum carbonization a heterocarbon with embedded platinum nanoparticles was obtained [33]. Due to the rigid molecular structure, the size range of the resulting platinum nanoparticles was uniform and the surface of platinum was “free” of capping agents, and although encapsulated, immediately highly reactive in electrocatalysis without further pre-treatments. In another recent study on PIM carbonization, Kim et al. [34] have shown that also the prototypical polymer of intrinsic microporosity PIM-1 [35] can be carbonized (at 1100-1300 °C) to give thin microporous carbon membranes with improved ion rejection
and flux properties for water purification under reverse osmosis conditions. Indeed, water flux and good salt rejection were reported to be better than those for current state-of-the-art water purification membranes.

In this study we explore the carbonization of PIM-EA-TB-H$_2$ (a polymer of intrinsic microporosity–PIM– with ethanoanthracene–EA– molecular motif linked via Tröger base–TB–functionality: see structure in Figure 1). This polymer is closely related to the previously reported polymer PIM-EA-TB [36], differing only in the lack of methyl groups at the bridgehead (9,10) position of the EA unit. PIM-EA-TB-H$_2$ is readily prepared in three simple steps from the cheaply prepared Diels-Alder adduct of ethylene and anthracene. A heterocarbon powder sample is obtained at 700 °C vacuum carbonization conditions and employed as ink applied to glassy carbon in electrochemical studies. Unique properties such as slow electrolyte-dependent hydration and pH-dependent capacitance responses are assigned to the presence of micropores and N-hetero atoms throughout the microporous carbon. Applications in “capacitive” pH sensing are suggested.

**Experimental**

**Chemical Reagents**

All solutions were prepared with deionized water of resistivity not less than 18.2 MΩ cm$^{-1}$ at 25 °C from a Thermo Scientific water purification system. Isopropanol, perchloric acid (70%), sulfuric acid (97%), phosphoric acid (85%), Nafton-117 solution (5% wt), sodium dihydrogenphosphate (99%) and sodium phosphate dibasic heptahydrate were purchased
from Sigma-Aldrich and used without further purification. PIM-EA-TB-H$_2$ was obtained following a literature recipe [36].

Briefly, the synthesis of PIM-EA-TB-H$_2$ was performed by nitration of the ethanoanthacene hydrocarbon, followed by reduction to give the required 2,6(7)-diaminoethanoanthracene monomer, which was readily polymerised by the action of dimethoxymethane in trifluoroacetic acid as previously reported for PIM-EA-TB [37].

**Instrumentation**

Electrochemical measurements were performed with a potentiostat system (IVIUM Compactstat) or a µAutolab type III potentiostat/galvanostat controlled by Autolab GPES software version 4.9 for windows XP in a conventional three electrode cell, with a Pt wire as a counter electrode, a KCl-saturated calomel reference (SCE, Radiometer, Copenhagen) as reference electrode, and a glassy carbon disk electrode (GC, BASi) with 3 mm diameter as working electrode. The heterocarbon powder was applied to the electrode as an ink (*vide infra*).

Morphologies of the PIM-EA-TB-H$_2$ before and after the carbonization were analyzed with a JEOL SEM6480LV scanning electron microscopy (SEM). Raman spectroscopy was carried out on a Renishaw inVia system with $\lambda = 785$ nm excitation. Thermogravimetric analysis (TGA) data were collected on a Setaram Setsys Evolution TGA instrument under Ar from 20 °C to 700°C. For nitrogen adsorption analysis, samples (ca. 0.025 g) were degassed (120 °C, 6 h) prior to analysis. Analyses were carried out (for three samples: (A)
PIM-EA-TB-H$_2$ treated with aqueous 0.1 M HCl, rinsed and dried; (B) PIM-EA-TB-H$_2$ treated with aqueous 0.1 M NaOH, rinsed and dried; (C) carbonized PIM-EA-TB-H$_2$) at 77 K on a Micromeritics 3Flex with P$_0$ measured continuously. Free space was measured post-analysis with He. Using nitrogen as the adsorbate gas at 77 K, an 80 point physisorption analysis was undertaken and the data analyzed by the DFT method (N$_2$ at 77 K, cylindrical pore, NLDFT equilibrium model).

Elemental analysis (obtained by Butterworth Laboratories Ltd., London in triplicate) suggest for sample (A) PIM-EA-TB-H$_2$ treated with aqueous 0.1 M HCl, rinsed and dried C 75.50 % H 6.15 %, N 9.47 %; sample (B) PIM-EA-TB-H$_2$ treated with aqueous 0.1 M NaOH, rinsed and dried C 77.61 % H 6.01 % N 9.78 %; and sample (C) carbonized PIM-EA-TB-H$_2$ C 83.19 % H 2.46 % N 5.67 %.

**Procedure for Electrode Preparation**

Carbonization of the PIM-EA-TB-H$_2$ powder was performed at 700 °C for 3 h under vacuum (Edwards oil pump at ca. 4 mbar) in a custom-made quartz tube employing an Elite Thermal Systems Ltd. oven. An amount of 2 mg of the black powder (cPIM) obtained from the PIM-EA-TB-H$_2$ carbonization and 100 µL of 0.25 wt% Nafion solution were ground together (mortar and pestle) and dispersed in 1 mL of isopropanol. The resulting ink was dispersed by sonication for 15 minutes to form a homogeneous ink. Different amounts of the ink (typically 3 µL = 5.5 µg cPIM carbon) were placed onto the 3 mm diameter glassy carbon disk electrode (GC, BASi) and dried under ambient conditions.
Results and Discussion

Characterization of Carbonized Polymer of Intrinsic Microporosity (cPIM)

The polymer PIM-EA-TB-H$_2$ is obtained as a pale-yellow powder and when vacuum carbonized at 700 °C a black hetero-carbon is obtained (see Figure 1A,B). Based on the elemental CHN analysis (see experimental) the highly porous polymer before carbonization is likely to contain captured guest molecules, which lead to some deviation from the ideal calculated elemental mass C 83.8 % H 5.9 % N 10.3 %. In particular, uptake of acids such as HCl lead to a decrease in carbon (C 75.50 % H 6.15 % N 9.47 %), but there is clear evidence for the presence of nitrogen. A treatment with aqueous 0.1 M NaOH followed by rinsing and drying changes the composition analysis (C 77.61 % H 6.01 % N 9.78 %) indicative of removal of at least some of the guest species. The vacuum carbonization to give cPIM increases the carbon content (C 83.89 % H 2.46 % N 5.67 %) and decreases the hydrogen content. The nitrogen content is lowered, but mainly retained, indicative of a polymer back bone staying intact. Data from thermo-gravimetric analysis (Figure 1D) shows that a minor release of water occurs around 100 °C (i) and the release of small structural components (e.g. ethylene from a retro-Diels-Alder reaction) occurs in the range 250-350 °C (ii) with onset of carbonization occurring from 380 °C (iii) to 460 °C (iv) with a final weight loss of approximately 25% at 700 °C. A hypothetical reaction scheme [28] is shown in Figure 1C suggesting some cross-linking as the carbonization progresses. Details of the carbonization chemistry are complex and temperature-/time-dependent and therefore the reaction scheme in Figure 1C is speculative. However, most
of the polymer backbone material is suggested to remain structurally intact to give a porous nitrogen-containing hetero carbon without chemical post-treatment with interesting implications for application of the heterocarbon.

**Figure 1.** (A,B) Photographic images for PIM-EA-TB-H\(_2\) before and after vacuum carbonization. (C) Hypothetical reaction scheme to rationalize weight changes during carbonization. (D) Thermogravimetric analysis (TGA) data for PIM-EA-TB-H\(_2\) under argon (see text).
Analysis of Raman data (Figure 2A,B) suggests that at an excitation wave length of 785 nm some fluorescence occurs in the background for the PIM-EA-TB-H₂ spectra, which are otherwise dominated by spectral lines at 800 cm⁻¹ and 1350 cm⁻¹ similar to those seen for PIM-EA-TB [28]. After 700 °C vacuum carbonization only the prominent Raman peaks for graphitic materials (no fluorescence) are observed at 1329 cm⁻¹ (D-peak, disordered amorphous carbon) and at 1594 cm⁻¹ (G-peak, graphitic carbon) [38]. The D-peak seems to be composed of overlapping peaks. Further deconvolution of the Raman spectra for cPIM shows that four Gaussian-shaped bands are present: G and D bands at 1329 cm⁻¹ and 1594 cm⁻¹, a shoulder present at the left of the D band at 1178 cm⁻¹ due to the TPA band (related to trans-polyacetylene-like structures, [39,40]) and the A band at 1512 cm⁻¹, which contributes to the intensity between G and D band and which is observed in other types of disordered carbons [33]. However, the D’ band, which is related to defect-induced breaking of translational symmetry, is not present in the spectra. There is also very little evidence for the 2D peak usually seen at 2770 cm⁻¹ for graphitic or carbon nanotube materials. The intensity ratio \( I_D/I_G = 1.17 \) is consistent with a high level of defects and some similarity of the spectrum exists in particular with glassy carbon materials [41].

Scanning electron microscopy images of the morphology of the polymer starting material (PIM-EA-TB-H₂) and of the product after 700 °C vacuum carbonization (cPIM) are shown in Figure 2C-2F. Figures 2C and 2D are obtained at lower magnification and Figures 2E and 2F show the morphology for the polymer precursor and for the cPIM product at higher magnification. As reported recently for a related PIM material [28], due to retention of the structural backbone there is very little change in morphology upon vacuum carbonization.
Figure 2. (A,B) Raman spectra obtained with $\lambda = 785$ nm for PIM and cPIM materials. (C,E) Scanning electron microscopy (SEM) images for PIM at low and at high magnification. (D,F) SEM images for cPIM at low and at high magnification. (G) SEM image for the cPIM ink ($3 \mu L = 5.5 \mu g$ carbon) applied to the 3 mm diameter glassy carbon electrode.
In order to obtain further insight into the micropore structure powder samples were investigated with nitrogen adsorption methods (BET) before and after carbonization. Figure 3A shows a comparison of isotherms comparing PIM-EA-TB-H₂ (i) treated with aqueous 0.1 M HCl (causing some swelling and protonation [42]), (ii) treated with aqueous 0.1 M NaOH (to remove unwanted protons and impurities; BET data for “as synthesized” PIM-EA-TB-H₂ are not shown but were similar to those for NaOH treated material), and (iii) a sample after vacuum carbonization. After carbonization nitrogen gas adsorption is lowered. The BET surface area (see Figure 3B) is reduced from 846 m²g⁻¹ for pure PIM-EA-TB-H₂ (somewhat lower when compared to PIM-EA-TB [28,41]) to 425 m²g⁻¹ for cPIM. DFT analysis of the pore size distribution (Figure 3C,D) suggests that changes in cumulative pore volume due to vacuum carbonization are relatively small (when compared to changes induced by protonation), which is in agreement with previous observations with a related polymer, suggesting only minor changes in morphology. A significant loss of very small micropores appears to occur upon carbonization, which could be due to cross-linking and loss of flexibility in the polymer chains.
Electrochemical Properties of Carbonized Polymer of Intrinsic Microporosity I:

Time Effect and Capacitance

In order to investigate the electrochemical properties of the carbonized material, cPIM, an ink based on isopropanol, 0.023 wt% Nafion (or 0.23 μg/μL), and cPIM (1.8 μg/μL) was prepared and applied to 3 mm diameter glassy carbon substrate electrodes. Typically a 3 μL volume of ink was deposited to give approximately 5.5 μg cPIM deposit (see SEM image in Figure 2G). When immersed into aqueous 0.1 M HClO₄ and when employing
cyclic voltammetry (see Figure 4A) an increase in capacitance is immediately observed upon immersion. However, gradually with time a much stronger increase in capacitance was noted. The value for the capacitance is estimated here from the capacitive background current at 0.25 V vs. SCE in the cyclic voltammogram based on specific capacitance = \( \frac{I_{\text{cap}}}{(\text{scan rate} \times \text{weight})} \). Figure 4C shows a plot of specific capacitance increasing over a period of 24 hours. This gradual increase is assigned here to a gradual penetration of protons and hydration into the microporous heterocarbon interior. Carbon particles are not uniformly sized (see Figure 2G) and in particular bigger particles may contribute to this effect more significantly.

The effect of scan rate on the voltammetric response (Figure 4B) is consistent with that for a capacitive current signal with approximately linear increase of current with scan rate. In order to investigate the effects of the amount of ink deposit, experiments were repeated with a scan rate of 5 mV s\(^{-1}\) for (i) 2.8, (ii) 5.5, (iii) 9.1, (iv) 12.7, (v) 16.4 μg cPIM ink on a 3 mm diameter glassy carbon electrode (see Figure 4D,E). The specific capacitance after 24 h submersion appeared reproducible up to approximately 10 μg cPIM ink deposit, but beyond this thickness a decrease is observed possibly linked to insufficient inter-particle conductivity in thicker film deposits. Therefore, experiments were performed with less than 10 μg cPIM loading.
Figure 4. (A) Cyclic voltammograms (10 mV s\(^{-1}\), third cycle) for cPIM (5.5 µg deposited onto a 3 mm diameter glassy carbon electrode) immersed in aqueous 0.1 M HClO\(_4\). The influence of the time submerged in solution on the voltammetric response is shown for (i) bare glassy carbon background, (ii) 5 min, (iii) 1 h, (iv) 12 h. (B) Cyclic voltammograms (scan rate (i) 1, (ii) 5, (iii) 10, (iv) 50 mV s\(^{-1}\)) for a 24 h submerged 5.5 µg cPIM coated electrode. (C) Plot of the specific capacitance (estimated from cyclic voltammograms) versus submersion time. (D) Cyclic voltammograms (scan rate 5 mV s\(^{-1}\)) for (i) 2.8, (ii) 5.5, (iii) 9.1, (iv) 12.7, (v) 16.4 µg cPIM deposited onto a 3 mm diameter glassy carbon electrode and immersed into 0.1 M HClO\(_4\). (E) Plot of apparent specific capacitance versus amount of cPIM deposit.
The slow increase in specific capacitance appeared to be linked to gradual hydration/protonation of the cPIM heterocarbon particles and additional experiments were performed exploring the effect of the electrolyte on this wetting process. Initial experiments in Figure 4 were performed in 0.1 M HClO$_4$ and in Figure 5 shows data for (A) 1 M HClO$_4$, (B) 1 M H$_2$SO$_4$, and (C) 1 M H$_3$PO$_4$ electrolyte. The effect of the HClO$_4$ concentration on the hydration process appears insignificant, but the effect of the electrolyte anion is pronounced.

When comparing data for HClO$_4$ and H$_2$SO$_4$, the perchloric acid solution appears to hydrate the cPIM much more rapidly. In contrast H$_3$PO$_4$ shows very little increase, even after long immersion periods. Additional experiments in non-acidic aqueous media suggest that acidic conditions are required for the specific capacitance to increase. Mechanistically, it can be suggested that protonation of nitrogen sites in the cPIM heterocarbon is involved in the internal wetting process. This is coupled to uptake of anions into small micropores and cavities. Strongly hydrated anions such as SO$_4^{2-}$ or PO$_4^{3-}$ are detrimental in this case, but the weakly hydrated perchlorate anion may be much more suited to bind into hydrophobic micropores and drive the internal uptake of electrolyte and water into cPIM micropores. **Porosity effects in the cPIM material electrochemistry appear to be dominated by very small pores and by ingress of protons into deeper regions of the heterocarbon material. This effect was not observed for cPIM obtained from PIM-EA-TB [28] and is likely to be linked to the higher carbonization temperature, here 700 °C compared to previously 500 °C [28].**
Figure 5. Cyclic voltammograms (scan rate 50 mV s$^{-1}$) for 5.5 µg cPIM on glassy carbon immersed for (i) bare glassy carbon background, (ii) 2 minutes, (iii) 10 minutes, (iv) 1 h, (v) 12 h in (A) 1 M HClO$_4$, (B) 1 M H$_2$SO$_4$, (C) 1 M H$_3$PO$_4$. (D) Comparison of cyclic voltammograms (scan rate 50 mV s$^{-1}$, solid bare glassy carbon, dotted 5.5 µg cPIM) immersed 12 h in (i) HClO$_4$, (ii) H$_2$SO$_4$, (iii) H$_3$PO$_4$. (E) Cyclic voltammograms (scan rate 10 mV s$^{-1}$) for 5.5 µg cPIM immersed 12 h in 0.1 M HClO$_4$ and then immersed (for 5 minutes each) sequentially in (i) 0.1 M HClO$_4$, (ii) 0.1 M phosphate buffer pH 7, (iii) 0.1 M HClO$_4$, (iv) 0.1 M phosphate buffer pH 7, (v) 0.1 M HClO$_4$. (F) Schematic drawing to illustrate internal protonation and wetting of cPIM micropores.
It is interesting to ask whether the hydration is permanent and whether a change in pH can reversibly or irreversibly change the specific capacitance of the cPIM material. In Figure 5E data for an experiment based on switching the pH is shown. A glassy carbon electrode was coated with 5.5 μg cPIM and then initially immersed into aqueous 0.1 M HClO₄ for 12 h. The specific capacitance reaches 55 F g⁻¹. When immersed into aqueous 0.1 M phosphate buffer pH 7 the capacitance immediately drops by an order of magnitude. Perhaps surprisingly, the capacitance is immediately switched back “on” by re-immersion into 0.1 M HClO₄ (Figure 5E). Therefore the internal hydration appears to be retained even if H⁺ and ClO₄⁻ ions are removed (Figure 5F). This switching process is reversible and an equilibration time of approximately 5 minutes appeared sufficient for a stable capacitance response reading to be obtained. This suggests that the state of protonation of the microporous cPIM material quickly responds to external pH conditions and with the change in internal protonation the specific capacitance dramatically changes even when the state of hydration is not changing. The internal protonation appears to be linked to an apparent increase in the capacitive surface area as indicated schematically in Figure 5F. A potential application of this phenomenon is envisaged in “non-potentiometric” pH sensing.

**Electrochemical Properties of Carbonized Polymer of Intrinsic Microporosity II.: pH Effects**

In order to further explore the effects of pH, electrochemical impedance experiments were performed. A glassy carbon electrode loaded with 5.5 μg cPIM and pre-immersed into
aqueous 0.1 M HClO₄ solution is employed. The electrode is kept in 0.1 HClO₄ and only for measurements transferred into aqueous 0.1 M phosphate buffer media. Figure 6A shows impedance data obtained at 0.25 V vs. SCE for a frequency range from 10 kHz to 0.1 Hz represented as Nyquist plots. An approximate equivalent circuit model to fit the data was based on an external resistance $R$, in series with a Warburg diffusion element to account qualitatively for internal ion and charge diffusion, and a series capacitance $C$ to account for the charging of the cPIM heterocarbon material. Data extracted for $R$ and for $C$ are shown in Figure 6B and 6C, respectively.

![Figure 6](image)

**Figure 6.** (A) Nyquist plots for impedance measurements (10 kHz to 0.1 Hz, 50 mV amplitude) for 5.5 μg cPIM on glassy carbon immersed for 2 minutes in 0.1 M phosphate buffer at various pH values. Before the measurement (12 h) and in between measurements the electrode was immersed in 0.1 M HClO₄. Lines indicates fits for a simple RWC model. (B) Plot of the resistance $R$ versus pH. (C) Plot of the capacitance $C$ versus pH. (D) Plot of the inverse time constant $(RC)^{-1}$ versus pH.
The series resistance shows an almost linear correlation to pH over a pH 4 to pH 12 range. The capacitance C appears independent of pH in the alkaline range, but then rapidly increases in the acidic pH range. At pH 2 the specific capacitance is estimated 70 F g\(^{-1}\) with further increase in the acidic range probably leading to super capacitor performance. It is interesting to combine R and C into the characteristic frequency (RC\(^{-1}\)) as shown plotted versus pH in Figure 6D. Again, a close to linear correlation with pH over a broad range of pH values is observed.

This “capacitive” pH reading is likely to be insensitive to colloid and surfactant adsorption and other interference which affect potentiometry. The application of a capacitive current response (capacitive pH sensor) based on the internal micropore protonation of cPIM particles could be robust and versatile and further study of this material is in progress. The properties of the cPIM material as well as the sensor architecture need to be further developed and optimized to be capable to work for example in complex media such as wound dressings.

**Conclusion**

It has been shown that the vacuum carbonization of the polymer of intrinsic microporosity PIM-EA-TB-H\(_2\) yields a novel microporous heterocarbon with properties very closely related to those of the starting polymer. The polymer backbone and porosity are maintained. Nitrogen in the molecular structure is retained (in part) during the 700 \(^{°}\)C carbonization process and protonation of nitrogen in micropores has been suggested to be
responsible for unusual phenomena such as a slow electrolyte dependent hydration process and a “capacitive” pH response. The immersion time and pH-dependent capacitance has been explained with ingress of protons associated with electrolyte anions into micropores with the associated increase in the apparent capacitive surface area. The protonation/deprotonation processes appear to be fast when compared to the wetting/dewetting processes of micropores. In future this will allow a simple impedance-based pH probe to be constructed without the need for a reference electrode and with performance characteristics that are likely to be complementary (and applicable in complex media) to those of existing pH sensing equipment.

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