Nitrogen-doped coatings on carbon nanotubes and their stabilizing effect on Pt nanoparticles†

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A homogeneous coating of nitrogen-doped carbon on carbon nanotubes is performed using ionic liquids. The N-doped material is employed as a support for nanoparticles. Electrochemical degradation behavior is monitored in situ and compared to an unmodified material. The strongly enhanced stability is explained on the basis of a Pt–nitrogen interaction.

Here, we present a simple, but very effective strategy to mitigate Pt NP coarsening in supported high surface area fuel cell electrodes. Our study is based on the hypothesis that surface heteroatoms, such as nitrogen, mediate the enhancement of platinum adsorption on the surface of carbon materials and slow down or even suppress its growth and surface area degradation. Nitrogen doping of carbon electrodes can be achieved via a variety of synthesis methods including pyrolysis and CVD of nitrogen rich precursors and post-synthetic methods like ion bombardment or thermal treatment in the presence of reactive nitrogen compounds such as ammonia. The method we have used throughout this study does—in contrast—not change the backbone of the material itself, but applies a thin coating of N-doped carbon on the surface of the electrocatalyst support via a post-synthetic method. On the one hand, this allows for the use of cheap and widely available bulk carbons as host material and on the other hand offers the possibility of a more direct comparison of doped and non-doped catalyst supports. Herein we show that by our method, growth and coarsening of Pt nanoparticles during electrochemical potential cycling are largely suppressed. We also show that the presence of N-C coatings does not compromise the electrocatalytic activity of Pt nanoparticles. The N-C coatings were formed from ion liquid precursors, which can wet the surface of the pristine carbon support and subsequently be pyrolyzed into nitrogen-rich carbon coatings. Particle growth on the pristine and N-C coated carbons was monitored using time-resolved small angle X-ray scattering (in situ SAXS), a non-invasive powerful, yet rarely used method to monitor growth trajectories of electrocatalytic particles in real time.

Ionic liquids are established precursor compounds in the field of materials chemistry, but just recently they have been applied as precursors for nitrogen-doped carbon. As liquids with negligible vapour pressure, such precursors can be easily processed and transformed into N-doped carbon materials by simple annealing under inert gas atmosphere. N-doped carbons derived from 3-methyl-N-butyl-pyridinium-dicyanamide (3MBP-dca) exhibit interesting properties: even at carbonisation temperatures of 1000 °C a high nitrogen content of approx. 10 wt% is observed in the resulting carbon material. The nitrogen atoms are incorporated into the carbon backbone in pyridinic or quaternary graphitic bonding environments within a structure that is dominated by graphitic microdomains. Furthermore, remarkable electric conductivities were reported, reaching the range of highly ordered graphite. Thus IL-derived...
N-doped carbons have been envisaged as interesting materials for electrochemical applications.

Unfortunately the usage of ILs as carbon precursors is accompanied with relatively high costs especially in comparison to the low price of most other carbon precursors. Thus to exploit the favourable properties of IL derived carbons in a most economical way, it was envisaged to prepare just thin coatings of the IL derived carbons on a convenient bulk carbon material. Such a coating also offers the opportunity to combine the properties of the IL derived carbon with the ones of the host. One interesting example has recently been presented using IL-derived N-doped carbon - Li2Ti3O12 composites as cathode material in lithium ion batteries. 30

In this study the IL 3MBP-dca was used to prepare a N-doped carbon (N-C) coating on the walls of carbon nanotubes (CNTs). 26,27 CNTs have been selected as a host, as their defined morphology allows for a good control of the homogeneity of the coating. Furthermore a preferable interaction of CNTs with ILs is known from the recent literature. 31 It should be however noted that in principle all other forms of carbons, like activated charcoal, should be as well feasible as host materials. Prior to the application of the N-doped carbon coatings, the CNT supports were surface oxidised using sulfuric acid in order to increase the number of polar end groups. This served to even further enhance the interaction between the polar ILs with the CNT surface. IL precursor concentrations were carefully adjusted to enable a homogeneous coating and to minimize any intertubular residues (see ESI† for detailed experimental procedures). An optimized 30 mol% of 3MBP-dca in water was applied in the wetting of the CNTs and subsequent formation of an N-doped carbon layer by thermal treatment under inert gas at 1000 °C. Higher concentrations of ILs or even pure ILs led to strong intertubular bulk carbon contributions, while lower concentrations did not yield sufficiently and homogeneously coated materials. The resulting composite material (henceforth referred to as “N-CNT”) was analysed by means of transmission electron microscopy (TEM), nitrogen sorption and X-ray photoelectron spectroscopy (XPS). Nitrogen sorption isotherms (see Fig. S1, ESI†) reveal a decrease of the surface area upon hydroxylation of the CNTs, probably due to increased intertubular interactions. In N-CNTs this effect is no longer observed due to the N/C-modified CNT surfaces. The micrograph in Fig. 1a further evidences no significant amounts of excess intertubular bulk residue. A similarly heat treated non-modified CNTs material, henceforth referred to as “CNTs”, were used as a benchmark support in this study.

The XPS survey scan in Fig. 1 shows peaks for C, O and N with concentrations of 97, 2 and 1 at.%, respectively. The nitrogen content may seem small at first glance. Nevertheless it has to be considered that a coating of carbon doped with ~10 wt% N has been applied on a carbon surface; thus higher N-contents can hardly be expected. High resolution analysis of the N1s region shows 3 components with binding energies of 286.8 eV, 287.9 eV and 288 eV. A specific assignment to functional groups, however, was not possible, because oxygen- and nitrogen-containing groups lead to very similar chemical shifts in that region.

To study the effect of N-doped interlayers on the electrochemically induced growth and surface area degradation of catalytic NPs, Pt nanoparticles were deposited 32,33 on the benchmark CNTs as well as on the N-CNTs support materials. Particle growth trajectories were measured in situ using SAXS. SAXS offers a time resolved analysis of the particle size distribution, characterized by the mean diameter and dispersion, the number density and the volume fraction of scattering centers. 34 Fig. 2a shows the high resolution TEM images of the as-prepared Pt-NP supported on the uncoated CNTs. The particles are homogeneously distributed on the support; a mean particle size of 1.7 nm was obtained from the log-normal distribution of the particle size resulting from counting 200 particles on TEM images (Fig. 2b).

![Image](image_url)

**Fig. 1** Transmission electron micrographs of N-CNTs with a chemical formula of the IL-coating precursor. XP survey spectrum of coated nanotubes with a highly resolved N1s spectrum.

**Fig. 2** High resolution TEM images of (a) 20 wt% Pt/(unmodified) CNTs and (b) a 17 wt% Pt/N-CNTs catalyst. (b) and (c) show the TEM based size histograms together with a log-normal fit. Histograms were based on 200 particles. (c) and (d) show experimental SAXS profiles with a log-normal fit for the 20 wt% Pt/CNTs and the 17 wt% Pt/N-CNTs, respectively. Insets in (c) and (f) show the extracted bimodal particle size distributions plotted on two separate y scales.
Probing a much larger particle population than TEM, SAXS provides statistically more reliable size information. Fig. 2c shows the background-corrected scattering curve of the pristine powder and its fit using a spherical particle form factor. The inset presents the corresponding bimodal particle size distribution (psd). The psd is characterized by a narrow distribution of smaller (~2.0 nm) and larger particles (~6.7 nm).

A similar analysis for the as-prepared Pt-NP supported on the N-CNTs is shown in Fig. 2d, e and f. The TEM derived mean particle size of the NPs was 1.9 nm, while SAXS data again suggested a bimodal size distribution clustering around 2.6 nm and 5.8 nm.

The crystallinity of the supported particles was probed by X-ray diffraction (Fig. S2, ESI†). Crystallite mean size analyses (2.1 nm for Pt-NPs on CNTs and 2.8 nm for Pt-NPs on N-CNTs) are reasonably consistent with TEM and SAXS-derived values.

Finally, the as-prepared CNT and N-CNT supported Pt-NPs powder catalysts were used to prepare a liquid catalyst ink suspension, cast into a thin catalyst electrode film and subsequently subjected to an electrochemical voltage cycling protocol to evaluate differences in the stability of the two nanoparticle catalyst-support couples. Emphasis was placed on the evolution of the mean particle size, the ECSA and the Pt-mass based catalytic activity for the oxygen reduction reaction (ORR) to water. This reaction occurs at fuel cell cathodes, the electrode potential of which can vary between 0.5 V and 1.5 V versus the hydrogen anode depending on the fuel cell operating conditions.4 Linear sweep voltammetry (Fig. S3a and S3d, ESI†) revealed Pt-mass based activities of 0.087 A mg⁻¹ for Pt-NPs supported on CNTs and 0.082 A mg⁻¹ for Pt-NPs on N-CNTs, which are in agreement with those of commercial carbon-supported Pt electrocatalysts.6 This shows that the N-C coatings had no significantly detrimental effect on the electrocatalytic ORR activity. Cyclic voltammetry (CV) (Fig. S3b and S3e, ESI†) of cathode electrode films in a degassed electrolyte yielded time-resolved ECSA trajectories shown in Fig. 3a and d. The ECSA loss is a macroscopic descriptor for the total catalytically active surface area and is affected by a large number of different factors, such as particle dispersion, support structure, and the accessibility of particles on the support surface. With initial ECSA values in the 50–60 m² g⁻¹ range,5,6 both catalysts showed similar ECSA losses over the 2400 potential cycles of 33% for the uncoated CNT and somewhat reduced 27% for the N-CNT supported particles.

To test our hypothesis of decreased particle coarsening on N-CNT supports, we performed in situ SAXS measurements monitoring particle size changes during voltage cycling. Background-corrected scattering I(q) curves of Pt-NPs on CNTs and N-CNTs are given in Fig. S3c and S3f (ESI†), respectively. Shifts of the decreasing portions of the scattering profiles towards smaller q values over time, seen in Fig. S3c (ESI†), evidence particle coarsening.17,35 Using the I(q) data, time-resolved particle size distributions, time trajectories of correlation length particle diameters (2Rg) and normalized number densities of scattering centers (N0) were extracted and compared for the CNT and N-CNT supports (see Fig. 3b, c, e, f, respectively). Pt-NP number density normalized diameters (see ESI†) on unmodified CNTs rapidly increased from 3.8 nm to about 5.5 nm during the voltage cycling test. Simultaneously the normalized number density of particles decreased dramatically by about 50%, suggesting a reduction in the number of particles due to coarsening.

The time trajectories of the N-CNT supported nanoparticles offer a stark contrast to those supported on the unmodified CNT. The mean particle size increased from 3.2 nm to only 3.9 nm, a mere 22% growth compared to the 45% growth of the CNT-supported particles. The size distribution function remains nearly unchanged during the potential protocol (Fig. 3f). Also, rather than a steep exponential drop, the particle number density exhibited a gradual decrease by only 30%.

The observed evolution of the SAXS-derived structural parameters of the Pt nanoparticles strongly suggests that, owing to its IL pyrolysis coating, the N-CNT support can provide improved structural stability of a Pt particle ensemble and reduce Pt particle coarsening during potential cycling. Based on the measured ECSA values, a mere increase in the surface area of the N-CNT support can be excluded as the origin of the stabilization. For the given initial particle diameters, the observed ECSA losses of about 30% appear to be poorly correlated with particle growth, and rather controlled by structure of the catalyst and the support as well as the interaction with the electrolyte, respectively. Carbon corrosion, e.g., is a process which contributes to the ECSA loss, but not to the particle size observed with SAXS. The pyrolysis of ILs is known to form pyridinic, pyrrolic, and other nitrogen containing functional surface sites, often at graphene sheet edges.36–40 A stabilizing interaction between such nitrogen-containing surface regions and small Pt nanoparticles could be the explanation for reduced dissolution/Ostwald-type ripening and/or reduced migration and coalescence. Surface nitrogen thereby raises the
dissolution potential of Pt making the formation of metal ions, which are migrating intermediates in the Ostwald ripening process, more difficult.

Conclusions

In summary, we have prepared a N-doped carbon nanotube material (N-CNT) by pyrolyzing an IL on the surface on a commercial CNT material. The pyrolysis products formed a thin N–C coating on the carbon nanotubes. We have used the N-CNT as support for Pt nanoparticles, the ORR performance of which was identical to that on unmodified CNT supports. In situ surface area measurements and particle size trajectories suggested a significantly reduced particle coarsening and degradation behavior. The surface nitrogen thus mediates the enhancement of platinum adsorption on the surface of carbon materials. Our results suggest that, more generally, N-doped carbons offer a promising family of porous support materials for electrodes, which operate at corrosive potentials and hence are prone to dissolution–reprecipitation of a catalytically active metallic component.

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Notes and references