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Surface transfer doping can mediate both colloidal stability and self-assembly of nanodiamonds

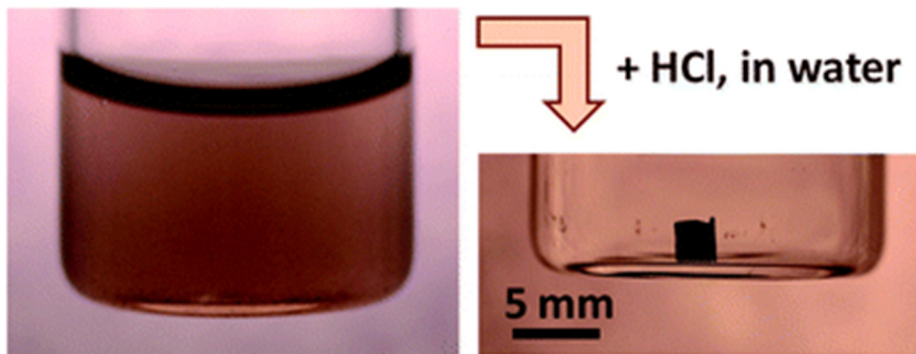
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Abstract

Although undoped diamond is insulating, hydrogenated bulk diamond surfaces exhibit surface conductivity under air and are electrochemically active in aqueous solutions. Due to their large surface/volume ratio, similar surface effects may exhibit a dramatic impact on the properties of nanodiamonds. Here we show that plasma-hydrogenated detonation nanodiamonds (NDs-H) display a positive zeta potential in water due to charge transfer with a redox couple involving oxygen in water. The transfer doping of NDs-H in water can be modulated by pH. Surprisingly, after acid addition, strong Coulomb coupling between NDs-H and adsorbed counterions induces the self-assembly of NDs-H into organized macro-structures reaching millimeter scale.



Nanomaterials often exhibit fundamentally new properties when compared to their bulk counterparts. Quantum confinement effects, high surface/volume ratio or new material crystal phases are some of the reasons for these new properties.¹ In the case of nanodiamonds (NDs) produced by detonation synthesis,² astonishing properties such as exceptional colloidal stability,^{3,4} giant permittivity⁵ or unusual charge exchange properties⁶ are hardly explained by the properties of their diamond core only. In contrast, peculiar surface phenomena observed on hydrogenated bulk diamond material have not been reported for NDs yet, such as the surface transfer doping occurring after air exposure^{7,8} or in water⁹ and their unique chemical reactivity.^{10,11} Similar surface effects may strongly influence ND properties due to their large surface to volume ratio. Here we show that surface transfer doping exists for NDs, and, beyond an electrostatic stabilization of the hydrogenated NDs (NDs-H) in water (zeta potential in water is +40 mV), it also mediates their self-assembly into millimeter scale structures in solution when brought to acidic pH values. NDs of larger size, synthesized by the High Pressure High Temperature (HPHT) method, could not be dispersed in water after similar hydrogenation treatment therefore the effects related here are specific to detonation NDs-H. NDs are generally considered as inert nanoparticles however our result demonstrates that surface

treatments such as hydrogenation can enhance electronic properties of the nanodiamond core. Driving NDs-H electrostatic interactions by modulation of their electrochemical environment may open new perspectives for self-assembly and also for gene delivery.

NDs-H were prepared by microwave-enhanced plasma treatments as described elsewhere.¹² Hydrogen plasma treatments have been shown to be the most adapted method to efficiently passivate the ND surface with hydrogen atoms while etching non-diamond carbon and oxygen species.¹³ In a previous study, we demonstrated that vanishing of the oxygenated terminations and the presence of C–H bonds after such plasma treatments occur, evidenced by FTIR and XPS analysis.¹² Furthermore, hydrogenation of NDs was also confirmed by the validation on our NDs-H of grafting routes specific to hydrogen-terminated diamond surfaces.^{10,14,15}

First, interactions between NDs and water molecules were investigated. The colloidal stability of NDs-H is unexpected since hydrogen-terminated bulk diamond surfaces are known to be hydrophobic.¹⁶ Water adsorption isotherms on NDs-H and oxidized NDs (NDs-COOH), used as a reference for hydrophilic NDs, are plotted and shown in Fig. 1. Surprisingly, a stronger water adsorption is observed on NDs-H compared to NDs-COOH. By taking into account their specific surface area, the density of hydrophilic sites can be estimated to be 1.4 and 1.7 sites per nm² on NDs-COOH and NDs-H, respectively (see ESI†).¹⁷ Water molecules interact strongly with the NDs-H surface as shown by the large mass increase at low relative water pressure, unlike with NDs-COOH ($P/P_0 < 0.05$, inset of Fig. 1A). In this pressure range, the mass increase is most probably related to electrostatic or hydrogen bonding of water molecules as well as water dissociation on the NDs-H surface. Water dissociation has already been observed on polycrystalline diamond and leads to the formation of hydroxyl groups.¹⁸ These hydrophilic sites on NDs-H are likely to facilitate the dispersion of NDs, initially thought to be hydrophobic, in water.

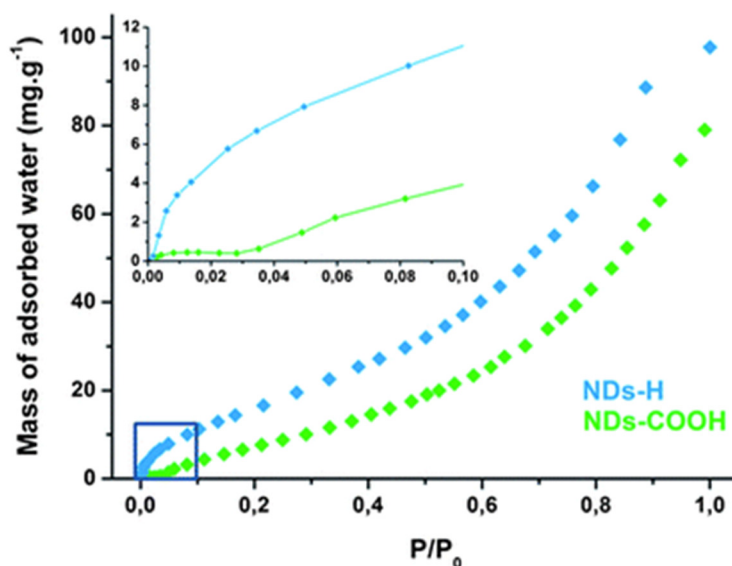


Fig. 1 Water adsorption on surface-modified nanodiamonds. Water vapor adsorption isotherms on NDs-H (blue) and NDs-COOH (green). Prior to water vapor exposure, the surface of NDs was annealed at 300 °C under vacuum to remove physisorbed and weakly bound species. The inset shows a magnified view of adsorption at low water relative pressure ($P/P_0 < 0.1$).

† Electronic supplementary information (ESI) available: Experimental procedures, specific surface area and hydrophilic site density calculations and the assembly experiment using sulfuric acid. See DOI: 10.1039/c3nr02492j

Colloidal properties of NDs-H were then evaluated. After 2 h ultrasonication in water, stable suspensions of NDs-H are formed (Fig. S2 and S3†). NDs-H display a highly positive ZP, stabilized around a value of +45–50 mV for pH values in the range of 4 to 9 (Fig. 2A). From pH = 9, the ZP decreases progressively up to an isoelectrical point (IEP), *i.e.* the point of zero charge, at pH = 12.4. Chakrapani *et al.* have shown that in water, the hydrogenated diamond surface can exchange charges with an electrochemical redox couple involving oxygen.⁹ On diamond microparticles, the authors demonstrated that hole accumulation on the diamond surface leads to a positive ZP at pH below 6. Considering that the positive ZP has the same origin on NDs-H, with the acidic character coming from atmospheric CO₂, the following reaction takes place on the NDs-H surface:

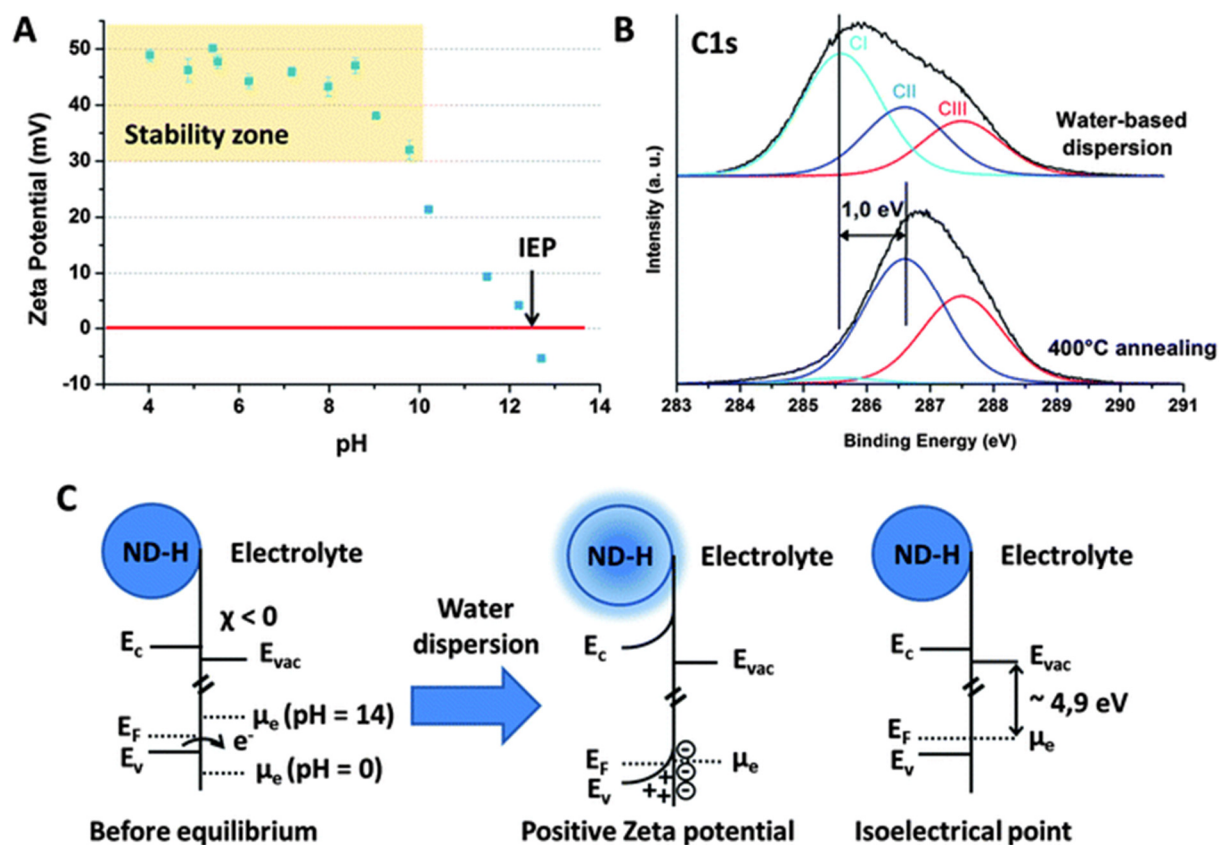
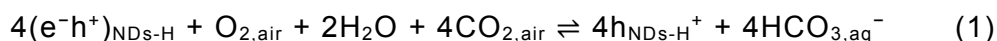


Fig. 2 Surface transfer doping on NDs-H in water. (A) Evolution of the Zeta potential of NDs-H in water. (B) Evolution of the XPS C1s spectrum of NDs-H after dispersion in water (top) and after *in situ* annealing at 400 °C (bottom). (C) Schematic view of the transfer doping model on NDs-H in water. The electron density in and around the ND-H is illustrated in blue color.

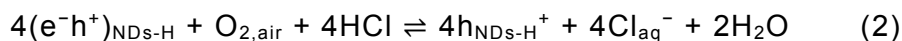
In this model, the charge transfer direction depends on the equilibrium between the electrochemical potential of the oxygen redox couple, driven by pH, and the Fermi level of the NDs-H. The key role of oxygen in the electrochemical process was validated by flushing the solution with N₂, which resulted in a rapid flocculation of the NDs-H, probably related to the loss of positive charges on their surface. The IEP thus corresponds to the pH at which, out of the equilibrium, the electrochemical potential of the redox couple equals to the Fermi level of the NDs-H. Thus, when the equilibrium is reached, no band bending occurs, and there is no accumulation layer. A schematic band model of transfer doping on NDs-H in water is proposed in Fig. 2C. By using the Nernst equation and considering a temperature of 298 K and a partial pressure of O₂ of 0.21 bar, an electrochemical potential of -4.9 eV is estimated for $\text{pH}_{\text{IEP}} =$

12.4. This value is higher than the one obtained on microparticles (−5.3 eV) by Chakrapani *et al.*, which may be related to nanosize effects. As a consequence, NDs-H exhibit a large band bending, leading to a high ZP up to pH = 9, which is not the case for microparticles.

The presence of a hole accumulation layer on the NDs-H after air exposure and water dispersion was validated by X-ray Photoelectron Spectroscopy (XPS). Oxygen and nitrogen atomic contents extracted from the O1s and N1s peaks are estimated as 3.0 and 5.7 at%, respectively. *In situ* experiments have previously shown that oxygen is fully desorbed after hydrogen plasma treatment under similar conditions.¹³ Therefore detected oxygen mostly results from adsorption of oxygenated molecules, which is in agreement with water adsorption measurements (Fig. 1). Note that this point is also in agreement with previous FTIR analysis of such hydrogenated NDs, which revealed the necessity to anneal the samples under vacuum to desorb these oxygen-related species from their surface.¹² The C1s spectrum of NDs-H was fitted with 3 components CI, CII and CIII with full widths at half-maximum (FWHM) of 1.5 eV at 285.6, 286.6 and 287.5 eV, respectively (Fig. 2B). Attributions of these different components are difficult at this stage, because adsorbates influence the XPS signature of the NDs-H. Indeed, adsorbates are known to induce the formation of a surface conductive layer on hydrogenated bulk diamond by transfer doping.⁸ As a consequence, downshifts of 0.7–0.8 eV of the C1s core level of hydrogenated bulk diamond have already been reported after air exposure,¹⁹ or adsorption of electron acceptors.^{20,21} An *in situ* annealing at 400 °C was thus performed to remove adsorbed moieties followed by a second XPS analysis. Surprisingly, the CI component almost disappears after adsorbate removal (from 50 to 3% of the total carbon area after 2 h annealing at 400 °C), while the oxygen content drops to 1.0 at%. The two other components CII and CIII are not affected by the annealing treatment. CII is thus attributed to undoped sp³ carbon bonds from the diamond core. CIII is mostly related to structural defects and C–N bonds as discussed elsewhere.^{22,23} CH_x (x = 1 to 3) bonds induced by the hydrogen plasma treatment may also contribute to the CIII component.²⁴ The binding energy difference between CI and CII is of 1.0 eV, which is comparable to the C1s downshift observed on bulk diamond after transfer doping. The CI component is therefore related to a surface conductive layer on the NDs-H, induced by transfer doping after air exposure and dispersion in water. Considering that the whole ND volume is probed by XPS, transfer doping thus affects half of the carbon atoms in the NDs-H.

Note that for surface-graphitized NDs, a totally different XPS C1s peak evolution after annealing has been observed.²² Transfer doping is therefore specific to NDs-H and the positive ZP of surface-graphitized NDs has another origin which has been discussed elsewhere.²²

Since transfer doping on diamond depends on the pH in water, it can be increased by acid addition. After addition of HCl, the presence of Cl[−] counterions in the solution should be considered and the electrochemical equilibrium at the NDs-H surface described in eqn (1) is replaced by:



The addition of HCl results in an increase of the hole density on the NDs-H surface, which is compensated by Cl[−] counterions adsorption. When equilibrium (2) dominates, addition of HCl should not result in a pH decrease since protons are directly converted into holes in the NDs-H and the only end-product would be water. In Fig. 3, the ZP of NDs-H is plotted after HCl addition to a water-based dispersion containing 6 ml of NDs-H at a concentration of 0.8 mg ml^{−1}. The initial pH of the NDs-H dispersion is slightly acidic (5.9) due to the dissolved CO₂ as discussed earlier. Although the surface charge of NDs-H increases upon acid addition, the ZP progressively decreases. ZP depends not only on the surface charge of NDs-H but also on the counterions in the Helmholtz diffuse layer surrounding the NDs-H,²⁵ therefore the strong

adsorption of Cl^- counterions on the NDs-H surface reduces the ZP. For a HCl concentration of 20 mM, the measured pH is 3.7 while it should be 1.7 considering the proton concentration related to HCl addition. Therefore NDs-H behave like proton sponges at acidic pH, which is in agreement with the transfer doping model. For 20 mM HCl, the hole density in the NDs-H is estimated to be 10^{17} to 10^{18} holes per cm^3 considering that the whole NDs-H volume is doped (which corresponds to underestimation of the hole density since only the surface volume should be doped according to XPS measurements).

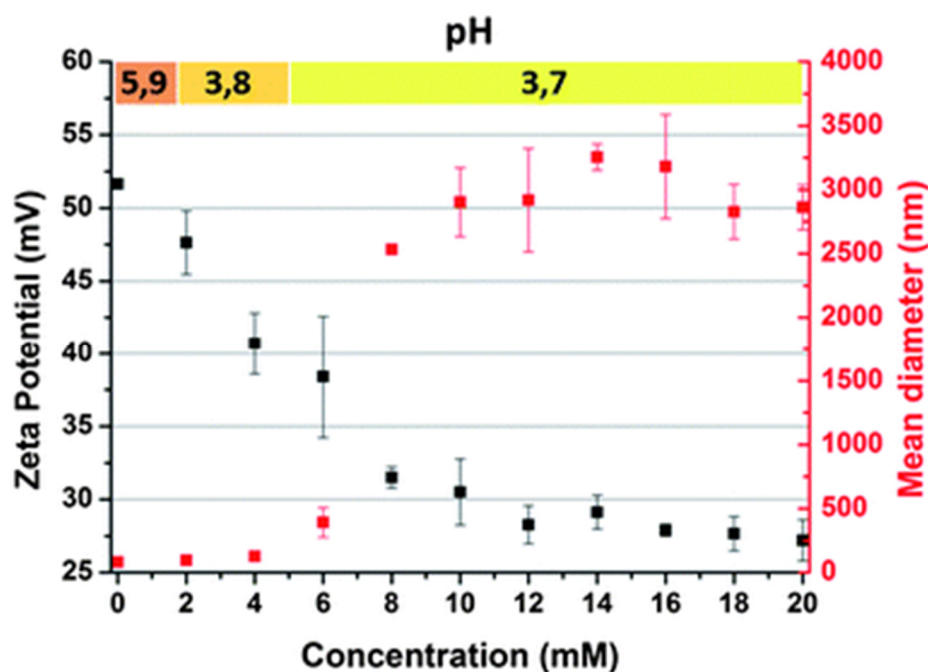


Fig. 3 Effects of acid addition in a water-based dispersion of NDs-H. The Zeta potential (black) and mean diameter of NDs-H/ Cl^- aggregates (red) are plotted after HCl addition. The initial NDs-H concentration is 0.8 mg ml^{-1} . pH values measured at each point are indicated on top.

In parallel to hole accumulation, the size increase of NDs-H aggregates is observed (Fig. 3). Up to 6 mM HCl, the size increase is limited below 500 nm while a sharp increase up to the micrometer scale is measured at 8 mM. The aggregation cannot be attributed to a low colloidal stability since the ZP of NDs-H is still higher than +30 mV, normally ensuring a good electrostatic stabilization. Aggregation is most probably related to electrostatic interactions between Cl^- counterions and NDs-H. Allahyarov *et al.* have shown that strong Coulomb coupling between nanoparticles and counterions can induce an attractive force between like-charged nanoparticles through a Coulomb depletion effect.²⁶ Briefly, for large densities of counterions adsorbed on nanoparticles by a strong Coulomb coupling, repulsive forces between counterions dominate over repulsive forces between the nanoparticles and create a depletion zone between two nearby nanoparticles. This Coulomb depletion results in a counterion-induced attractive force between the two like-charged nanoparticles, leading to aggregation. Here, Coulomb depletion forces are likely to appear at high HCl concentrations due to the high surface charge of NDs-H and the strong Coulomb coupling between NDs-H and Cl^- counterions.

Aggregates formed by HCl addition were observed by Scanning Electron Microscopy (SEM) (Fig. 4A–C). At low concentrations, aggregates have spherical shapes (Fig. 4A). Surprisingly, cubic structures with smooth surfaces appear at higher concentrations (Fig. 4B and C), with a high reproducibility on several samples. Organized structures with sizes ranging up to the millimeter scale resulting from NDs-H/ Cl^- ion coupling were even observed in solution at a HCl

concentration of 80 mM (Fig. 4D). Similar experiments performed with NDs-COOH did not result in any organized structure therefore transfer doping certainly plays an important role in the self-assembly process. Other reports suggested that Coulomb interactions between facets of detonation NDs, presenting different electrostatic potentials, may impact their self-assembly.^{27,28} Although few HRTEM investigations reported facets on detonation NDs,²⁹ their presence on our NDs-H has not been demonstrated.

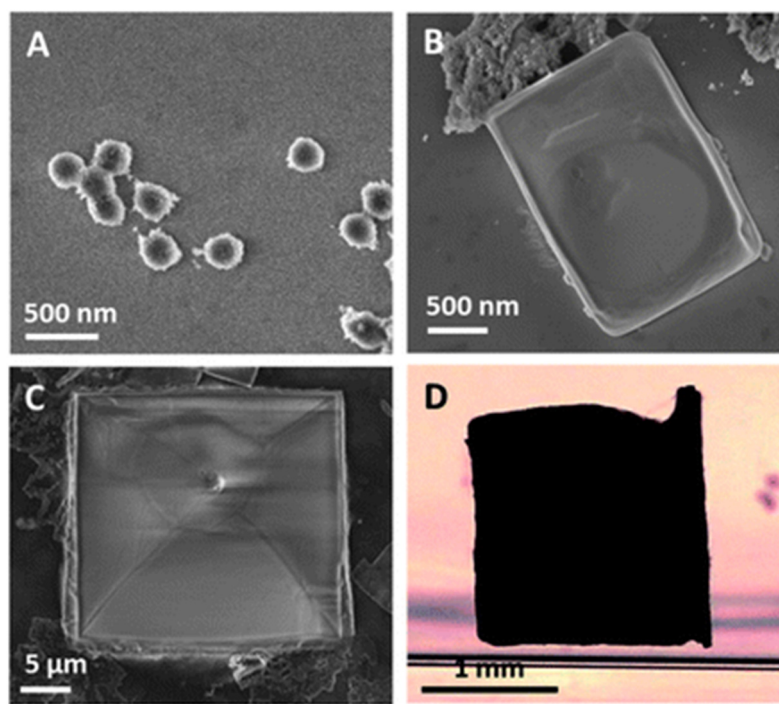


Fig. 4 Self-assembled NDs-H/Cl⁻ complexes. (A–C) SEM pictures of self-assembled structures for increasing HCl concentrations. (D) Optical image of a self-assembled structure with dimensions $1.7 \times 1.7 \times 0.8 \text{ mm}^3$.

Furthermore, the valence of counterions in water also plays a critical role in the shape formation of the self-assembled structures. Indeed, similar experiments were performed with sulfuric acid (H₂SO₄), resulting in the adsorption of SO₄²⁻ counterions (see ESI†). In this case, large scale self-assembly was also observed at lower acid concentrations and resulted in rod-like structures (Fig. S4†). The effect of the counterions on the shape of the macrostructures is an intriguing question which remains to be explained.

To conclude, we anticipate that self-assembly induced by transfer doping may be applied to other nanomaterials based on graphene and organic semiconductors,³⁰ silicon³¹ or other semiconducting materials³² which are also sensitive to transfer doping. By using various acids and electron acceptors, more complex shapes might be obtained directly in solution. The proton sponge effect observed on NDs-H at acidic pH also suggests that they are good candidates as vectors for gene delivery since cationic nanoparticles with buffering properties already showed enhanced gene transfection capabilities.³³

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