Nanoscale



PAPER View Journal | View Issue



Cite this: Nanoscale, 2020, 12, 21757

Controlling palladium morphology in electrodeposition from nanoparticles to dendrites *via* the use of mixed solvents†

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By changing the mole fraction of water (χ_{water}) in the solvent acetonitrile (MeCN), we report a simple procedure to control nanostructure morphology during electrodeposition. We focus on the electrodeposition of palladium (Pd) on electron beam transparent boron-doped diamond (BDD) electrodes. Three solutions are employed, MeCN rich (90% v/v MeCN, χ_{water} = 0.246), equal volumes (50% v/v MeCN, χ_{water} = 0.743) and water rich (10% v/v MeCN, χ_{water} = 0.963), with electrodeposition carried out under a constant, and high overpotential (–1.0 V), for fixed time periods (50, 150 and 300 s). Scanning transmission electron microscopy (STEM) reveals that in MeCN rich solution, Pd atoms, amorphous atom clusters and (majority) nanoparticles (NPs) result. As water content is increased, NPs are again evident but also elongated and defected nanostructures which grow in prominence with time. In the water rich environment, NPs and branched, concave and star-like Pd nanostructures are now seen, which with time translate to aggregated porous structures and ultimately dendrites. We attribute these observations to the role MeCN adsorption on Pd surfaces plays in retarding metal nucleation and growth.

Received 29th July 2020, Accepted 6th October 2020 DOI: 10.1039/d0nr05630h

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Introduction

In electrochemical devices used for energy conversion, nanostructures such as nanoparticles (NPs), made of metals, for example the platinum group metals, control the electrocatalysis process. $^{1-3}$ The electrocatalytic reaction can be optimised by careful control of the metal catalyst crystal structure, size, loading, presence of defects $\it etc.^{4-9}$ The need to produce effective electrocatalysts has led to the exploration of many different chemical and physical NP and nanostructure synthesis strategies. 2,10,11 These include chemical vapour deposition, atomic layer deposition, colloidal synthesis, sputtering $\it etc.^{12-16}$ Palladium (Pd) is an interesting metal due to its use in

Electrodeposition offers an interesting route for nanomaterial production, as it is effectively bottom-up synthesis, lends itself to scalability, and does not require sophisticated instruments or a high technical level of training. 18–22 Electrodeposition also offers a wide variety of tuneable parameters for controlling nanostructure morphology, these include *e.g.* electrode deposition potential and time of deposition, temperature, and electrolyte composition. 23–26 It is thus not surprising that electrodeposition has been used to make a variety of controlled Pd nanostructures such as nanowires, 17 cubes, nanoflowers, nanoframes, and high index faceted Pd nanocrystals. 11,27–29

Interestingly, despite the vast number of electrodeposition studies, the majority are carried out in aqueous solutions, as is the case for Pd, using salts that contain anions such as Cl⁻, ClO₄⁻, SO₄⁻², NO₃^{-,30} Few studies use different salt and solvent systems, and even less consider the role mixed solvent systems can play in controlling deposition morphology.³¹ Changing solvent can offer advantages to electrodeposition such as changing the (i) double layer structure, which includes the impact of solvent adsorption, (ii) solvation shell structure, (iii) redox potential, (iv) metal reduction electron transfer

the production of hydrogen from the electrochemical water splitting reaction and hydrogen storage. ^{1,17} To move from laboratory based research to commercial applications, low cost, scalable methods of nanostructure fabrication are essential. ¹⁷

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[†] Electronic supplementary information (ESI) available: Assignment of the Raman spectra recorded in the three different MeCN–water solutions. ESI2: Cyclic voltammetry of Pd-acetate in the three different MeCN–water solutions. ESI3: High resolution ADF-STEM of Pd nanostructure electrodeposition. ESI4: Statistical analysis for Pd nanostructures electrodeposited for $t_{\rm dep}$ = 50 s and 15 s. (PDF). See DOI: 10.1039/d0nr05630h

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kinetics, (v) altering the diffusional properties of ions in the solvent, (vi) increasing the potential window, enabling metal reduction outside the aqueous solvent window, and (vii) increasing the variety of electrodeposition salts which can be dissolved etc.32

Mixing two or more (miscible) solvents together offers a further control parameter in the study of electrodeposition.23,31,33 Whilst each solvent possesses its own chemical and physical characteristics, 33 when mixed the structure and physical properties of each individual solvent component, the solvation state of the metal ion, the electrical double layer, along with the other factors highlighted above, will change depending on the mole fraction of the individual solvent in the solution.³³ For example, in batteries it has already been shown that by moving to mixed-solvents, metal dendrite formation can be suppressed³⁴ and in electrocatalysis, water molecule reactivity can be increased through watersolvent co-ordination bonding.35 Furthermore, many of the studies that work in a non-aqueous solvent system, be it organic, ionic liquid or deep eutectic solvents are in reality highly likely to be mixed solvent due to the uptake of trace water, which can be difficult to exclude completely.36

In this work we investigate the electrodeposition of Pd, from its metallo-organic salt (Pd-acetate) in miscible organicaqueous solvents; here water and acetonitrile (MeCN), under organic rich (90%:10% MeCN:water v/v), water rich (10%:90% MeCN: water v/v) and equal contributions of both solvent by volume (50%:50% v/v), conditions. We provide insights into how MeCN and water bulk solvent structures are modified by the presence of the second solvent and explore the role the water-MeCN mixed solvent systems play in controlling atom level and nanoscale electrodeposition of Pd.

Experimental

Chemicals and solution preparation

Experiments were carried out using mixed-solvent systems, prepared using variable ratios of MeCN, water, tetrabutyl ammonium tetrafluoroborate (Bu₄N·BF₄), and KCl, while keeping the concentration of Pd-acetate fixed at 1×10^{-3} M. Solution 1 (organic rich) is made of 1×10^{-3} M Pd-acetate + 0.1 M Bu₄N·BF₄ in 90%: 10% MeCN: water v/v; corresponds to a mole fraction (χ) of $\chi_{\text{MeCN}} = 0.757 + \chi_{\text{water}} = 0.243$. Solution 2 is made of 1×10^{-3} M Pd-acetate + 0.05 M $Bu_4N \cdot BF_4 + 0.05 \text{ M KCl in } 50\% : 50\% \text{ MeCN : water } v/v \left(\chi_{MeCN} = 1.000 \text{ MeCN : water } v/v \right)$ $0.257 + \chi_{\text{water}} = 0.743$). Solution 3 (water rich) is made of 1×10^{-3} M Pd-acetate + 0.08 M KCl in 10% : 90% MeCN : water $v/v (\chi_{MeCN} = 0.037 + \chi_{water} = 0.963 (10\% : 90\% v/v)$. The solutions were prepared with trace element grade chemicals, and all chemicals were used as received without further purification. 99.99% HPLC far UV-Vis grade MeCN and deionised water from a Purite Select HP system were used for solution preparation (with a resistivity of 18.2 M Ω cm at 25 °C). Pure 99.999% Ar and N2 (BOC industries, UK) were used to deaerate the solutions.

Electrochemical studies

Electrochemical measurements were performed using a CHI potentiostat (750C) and IVIUM Compactstat potentiostat in a three-electrode configuration with a Ag|Ag⁺ double junction reference electrode. The reference electrode was constructed from Ag wire immersed in a MeCN solution containing 0.1 M Bu₄N·BF₄ and separated from a compartment containing the same solution by a porous Vycor tip (BASI). The working electrodes were produced from a freestanding microcrystalline BDD wafer (boron dopant level of $\sim 3 \times 10^{20}$ B atoms per cm³) grown using microwave chemical vapor deposition (Element Six, Harwell, Oxford, UK). For all electrochemical experiments, BDD transmission electron microscopy (TEM) supports, with a (110) crystalline surface texture, which also function as electrodes, were employed.³⁷ These were made from the BDD wafer thinned down to $\sim 50 \, \mu m$ thickness by mechanical polishing, with both sides showing a surface roughness of ca. 0.17 nm (measured using atomic force microscopy) on the surface of a grain (grain size ca. 2-8 µm). The thinned wafers were then cut into 3 mm diameter disks using laser micromachining (355 nm Nd:YAG laser micromachiner, E-355-ATHI-O system, Oxford Lasers Ltd UK, with a nominal pulse-length of 34 ns). The supports were acid cleaned in KNO₃/H₂SO₄ solution to remove debris after laser micro-machining. To produce areas of electron transparency the supports were Ar⁺ ion milled at an accelerating voltage of 6 kV and an angle of incidence of ~4°; the preparation procedure is described in detail in ref. 37. After ion milling, the BDD was acid cleaned using 0.1 M HNO3 followed by 0.1 M H₂SO₄ and then washed carefully in deionised water. To make an ohmic contact to the TEM electrode, the upper quarter of one of the edges was sputtered (Moorfield MiniLab 060 Platform) with Ti (20 nm)/Au (400 nm) and annealed in a tube furnace for 5 h at 450 °C. The electrochemical set-up showing how to use the BDD TEM support/ electrode is described in ref. 37. Electrodeposition takes place on both sides of the support over an area of ~ 0.06 cm².

Microscopy characterisation

Microscopic studies were carried out ex situ after electrodeposition on the TEM BDD electrode. Before imaging, the electrode was gently rinsed with de-oxygenated water, dried under inert atmosphere and kept in a desiccator to avoid any contamination during transfer. The morphology of the deposited Pd material on the TEM electrodes were characterised by ex situ techniques using field emission scanning electron microscopy (FE-SEM) for low magnification and annular dark-field scanning transmission electron microscopy (ADF-STEM) for higher resolution. FE-SEM images were recorded using the in-lens, EsB, SE2 and STEM detectors on a Zeiss Gemini or Zeiss Supra 55-VP operating at 5 kV, 15 kV and 25 kV, respectively. For each deposition, at least three images (n = 3) were recorded in different areas of the surface for both FE-SEM and ADF-STEM measurements and an average of 150 Pd nanostructures analysed. ADF-STEM images were obtained using a double aberration corrected JEOL JEM-ARM200F at 200 kV using an annular field detector with a fine-imaging probe of FWHM ~0.085 nm,

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and probe current of 23 pA with a convergence semi-angle of \sim 25 mrad and an inner angle of 50 mrad.

X-ray energy dispersive spectrometry (XEDS) spectra were recorded using the XEDS unit (Oxford Instruments) with the FE-SEM, at a working distance of 10 mm and accelerating voltage of 15 kV. Raman was conducted on a Horiba LabRam HR Evolution Raman microscope at room temperature, with laser wavelengths of 660 nm (100 mW), 660 nm edge laser filter, 1800 mm $^{-1}$ gratings, laser power 100% (50 mW), and 150 μm iris size. LabRam was equipped with a UV-Vis-NIR macro multipass cuvette cell holder and the samples were in a Quartz Cuvette.

Results and discussion

Solvent-solvent interactions in MeCN-water systems

Raman microscopy was used to provide information on changes in intra- and inter-solvent–solvent interactions as the water content in MeCN was increased. The Raman spectra of the three solutions containing 1×10^{-3} M Pd-acetate in (1) organic rich, 90% MeCN: 10% water v/v ($\chi_{\text{water}} = 0.243$); (2) equal by volume MeCN and water *i.e.* 50% MeCN: 50% water

v/v ($\chi_{\text{water}} = 0.743$), and (3) water rich, 10% MeCN: 90% water v/v ($\chi_{\text{water}} = 0.963$) solutions, are shown in Fig. 1, and ESI1, Fig. S1.† The vibrational stretching mode band assignments for C=N and C-H (MeCN) and O-H (water) are listed in ESI1, Table S1† and assist with peak labelling in Fig. 1.

Fig. 1(a) shows the Raman spectral region associated with the C \equiv N stretch of MeCN whilst Fig. 1(b) shows the Raman spectral region of both the C-H and O-H³⁸⁻⁴⁰ stretches, associated with MeCN and water, respectively. The emergence of the O-H band is associated with a tetrahedrally hydrogen-bonded network of water molecules *i.e.* bulk water,³⁸ whilst the C-H and C \equiv N bands are associated with the MeCN molecules. The different coloured lines represent the different solutions as water content is increased (expressed in terms of water mole fraction, χ_{water} , in the solution). Grey is $\chi_{\text{water}} = 0.243$ (solution 1, organic rich), red is $\chi_{\text{water}} = 0.743$ (solution 2, equal volume), and green is $\chi_{\text{water}} = 0.963$ (solution 3, water rich).

Monitoring changes in the position and intensity of bands in the different Raman regions as a function of χ_{water} can be used as a diagnostic for how inter- and intra-solvent bonding interactions change. ^{38–40} Qualitatively, as water content increases both the C \equiv N and C–H bands shift to a higher wavenumber and the intensity decreases. Only in solution 3 was an

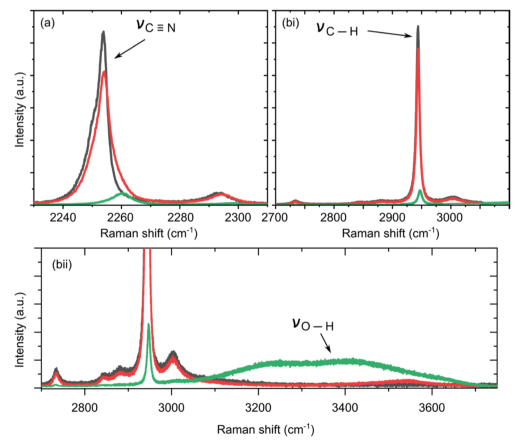


Fig. 1 Raman spectra of (a) $C \equiv N$ (MeCN), and (bi) C - H (MeCN) and O - H (water) stretches as function of increasing water mole fraction χ_{water} , (bii) is a zoom-in to the spectra shown in bi. Grey line represents solution 1 = 90% MeCN: 10% water 10%

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O-H band recorded. This behaviour can be interpreted from a microheterogeneity viewpoint. 38,41,42 In solution 1 (MeCN/ organic rich), the absence of an O-H band indicates that the minority water molecules exist as discrete molecules and are unable to form a hydrogen bonded water network. In solution 2, as the water content increases, the percentage of MeCN molecules coordinated to water through H-bond interactions increases as evidenced by the positive shift of the C=N Raman bands. In solution 3 (water rich), there is enough water to form a H-bonded water network and the characteristic O-H band is observed.

The C≡N Raman spectral region in Fig. 1(a), and ESI1, Fig. S2 and S3,† can be fit with two components, one that is associated with bulk MeCN (recorded at low Raman shift), and a second associated with MeCN bound to water molecules (recorded at higher Raman shift). Using the Shurvell method⁴³ it is possible to determine the fraction of MeCN molecules bound to MeCN molecules, as water content increases. In solution 1, this fraction is determined as 0.83 i.e. the majority of MeCN are binding to other MeCN, whilst in solutions 2 and 3 this fraction decreases to 0.67 and 0, respectively i.e. in the presence of 90% water, all the MeCN molecules exist bound to water.

Electrodeposition

To explore the impact of χ_{water} in MeCN on the resulting Pd morphology, nucleation and growth was monitored as a function of time in the three different MeCN: water mixtures containing the Pd-acetate salt. The corresponding CVs for electrodeposition in each solution are shown in ESI2, Fig. S4.† The CVs show that in the MeCN rich solution, electrodeposition initiates at a potential E_{dep} vs. $Ag|Ag^+ \sim -1.0$ V, with a maximum current occurring at -1.26 V vs. Ag|Ag⁺. As water content increases, $E_{\rm dep}$ shifts more positive, up to ~0.5 V in the water rich solution, indicating that the process is becoming more kinetically favourable in the presence of water. Note, in MeCN only (no water present) it has been previously shown using IR, NMR, XRD, and UV-vis that electrodeposition of Pd is not possible as the acetate ligands (bound trimerically, Pd₃(acetate)₆) remain co-ordinated during reduction of Pd²⁺ to Pd(0).⁴⁴ In the organic rich solution, adding a small amount of water (10% v/v, ($\chi_{\text{water}} = 0.243$) results in partial destabilisation of the Pd-acetate bond due the strength of the interaction between water and acetate overcoming that between Pd and acetate.44 This results in water preferentially binding to the acetate ligands and freeing up the Pd cation for solvation by MeCN, Pd can now electrodeposit.

To investigate how water content affects the morphology of the resulting Pd nanostructures at atom level resolution, ADF-STEM images were recorded of the Pd deposits formed directly on an electron transparent BDD electrode. Using a combined TEM substrate-electrode allows direct interrogation of electrodeposition structures on the electrode without the need for material transfer (and possible loss or modification of deposited material). The same three solutions were used, as in Fig. 1 which progressively increase in water

content. An overpotential, $\eta = -1.0$ V, with an initial deposition time, $t_{dep} = 50$ s was employed for these studies. The large overpotential means that for all three systems electrodeposition is likely to be under mass-transport (diffusion) limited conditions, for a sufficient period of time to form nanostructures. Note electrodeposition occurs on both faces of the BDD TEM electrode, as shown by the nanostructures labelled by the green (front face) and blue (back face) arrows in Fig. 2.

After $t_{dep} = 50$ s, in organic rich media (solution 1), Fig. 2(a) and 3(a), demonstrate that electrodeposition yields Pd atoms (Fig. 3(ai)), amorphous atom clusters (Fig. 3(aii)), and crystalline Pd NPs (Fig. 3(aiii-av)). Note we use the term NP here to describe Pd structures which are largely spherical (circular from a 2D viewpoint), all other morphologies are encompassed by the term nanostructure. Pd NPs are the dominating morphology (95%) as shown in Fig. 2(ai-aii), with diameters in the range 1 nm-7 nm. Looking more closely at the highest resolution image in Fig. 2(aiii), shown in more detail in ESI3, Fig. S5,† there is evidence of epitaxial alignment of the Pd NPs with the underlying BDD TEM electrode. In a few isolated cases, NP coalescence events are also observed, as indicated by the red arrows in Fig. 2(ai).

Statistical analysis of the NPs and nanostructures shown in Fig. 2 and 3, is given in terms of the circularity (a circularity of 1 indicates a perfect atom-resolution circle, ImageJ analysis)⁴⁵ and the perimeter, as presented in Fig. 4. Such descriptors are useful for assessing structures which contain heterogeneity i.e. where not all structures are the same shape. This is especially important for analysing and characterising structure morphological evolutions as both water content in the solution and time of electrodeposition increases. In the organic rich solution 1, structures with an average perimeter = 8 ± 4 nm (Fig. 4(b)) and circularity = 0.7 ± 0.05 (n = 100, counted on both faces of the electrode) were obtained.

For equal volumes of water and MeCN, (solution 2), Fig. 2(b) and 3(b) show that electrodeposition results in the formation of Pd atoms (Fig. 3(bi)), amorphous atom clusters (Fig. 2(biii)), monocrystalline NPs, defected NPs and more complex nanostructures (Fig. 2(b) and 3(bii-bv); the latter three comprise 96% of all morphologies observed on the surface. Fig. 3(bii-3bv) provide higher resolution examples of representative NPs and nanostructures on the BDD TEM electrode. The NP in Fig. 3bii is an example of a single crystal, and almost spherical, NP representing 40% of the total Pd structures formed as a result of electrodeposition. The nanostructure in Fig. 3(biii) is also crystalline, but is hexagonal in shape. It is projected along the [110] direction with different planes exposed on the surface and missing surface atoms, as demonstrated by the green line.

Fig. 3(biv and bv) provide examples of an elongated and defected Pd nanostructure, respectively. Such Pd nanostructures, which can also be described as irregular, and result in a substantial deviation from circularity, are likely to have formed due to coalescence (aggregated growth) of two neighbouring NPs or an amorphous atom cluster and NP. 18,37

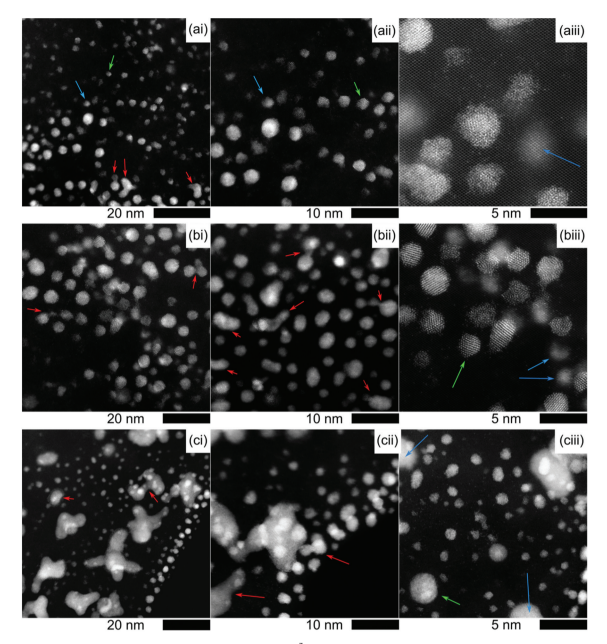


Fig. 2 ADF-STEM of Pd nanostructure electrodeposition from 1×10^{-3} M Pd-acetate in (a) 90% MeCN:10% water v/v ($\chi_{water} = 0.243$), (b) 50% MeCN:50% water v/v ($\chi_{water} = 0.743$), (c) 10% MeCN:90% water v/v ($\chi_{water} = 0.963$) at an E_{dep} of $\eta = -1.0$ V for $t_{dep} = 50$ s. Red arrow represents coalescence of nanostructures, green arrows provide examples of nanostructures electrodeposited at the front face of the BDD TEM electrode, whilst the blue arrows provide examples of nanostructures electrodeposited on the back face.

Whilst, such events have been seen before in electrodeposition they are often associated with much shorter timescales (early stage growth). $^{18,37,46-48}$ The observation of an increasing number of such structures demonstrates how increasing water content is influencing the growth process. The average perimeter of all assessed nanostructures (n = 200) for electrodeposition is now 11 ± 6 nm (Fig. 4(c)) with an average circularity value of 0.5 ± 0.1 (Fig. 4(a)). The drop of the circularity value accompanied with an increase in the perimeter value, highlights the increased prominence of the more elongated/defected nanostructures as seen in Fig. 2(bi and bii).

Electrodeposition of Pd nanostructures in water rich media ($\chi_{\text{water}} = 0.963$, solution 3) is shown in Fig. 2(c) and 3(c). After $t_{\text{dep}} = 50$ s, the following nanostructures exist; Pd atoms (Fig. 3 (ci)), amorphous atom clusters (Fig. 2(ciii)), both monocrystalline and defected Pd NPs with diameters in the range 3 nm to 7 nm (comprising 70% of all structures on the electrode surface), Fig. 2(ci-ciii) and 3(cii and ciii), elongated nanostructures but now also branched, concave²⁹ and star-like Pd nanostructures (Fig. 2(ci-ciii) and 3(civ and cv)). The atoms and amorphous atom clusters are in the minority (2%) in these images. In water rich solutions, NP coalescence events

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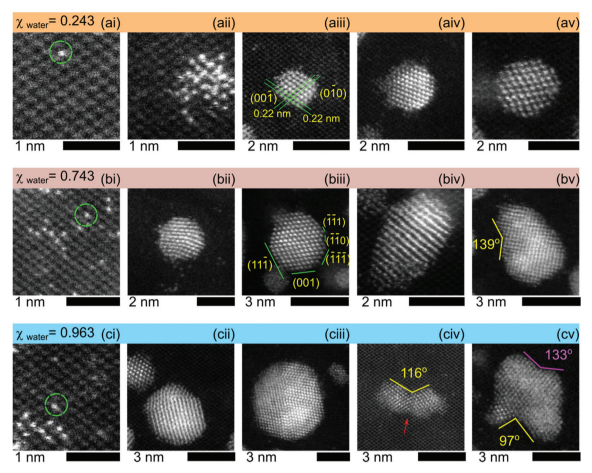


Fig. 3 High magnification ADF-STEM images of typical Pd structures observed on the BDD-TEM electrode after electrodeposition from 1×10^{-3} M Pd-acetate in (a) 90% MeCN:10% water v/v ($\chi_{water} = 0.243$), (b) 50% MeCN:50% water v/v ($\chi_{water} = 0.743$), (c) 10% MeCN:90% water v/v ($\chi_{water} = 0.963$) at an electrodeposition potential of $\eta = -1.0$ V for $t_{dep} = 50$ s. Single atoms are highlighted by the green dashed circle.

are more prominent. Such events are shown in the high magnification ADF-STEM images in Fig. 3(civ and cv). The former shows two Pd NPs that have merged laterally (coalesced) during electrodeposition, as indicated by the red arrow with an angle of 116°. The latter shows a resulting concave Pd nanostructure.

The resulting nanostructures suggest that during growth, when two or more of the NP coalesce, the newly formed structures are distorted containing defects and high index facets. 18,37,47,49,50 Interestingly, there is still no evidence of aggregation prior to coalescence, although this could be a result of capturing the data at $t_{dep} = 50$ s, shorter deposition timescales may reveal more information. Generally, the Pd nanostructures show a very low average circularity value of 0.2 ± 0.13 (Fig. 4(a)) indicating that most nanostructures in this case are not circular (highly distorted). Measurement of the perimeter values (Fig. 4(d)), indicates a bimodal distribution, highlighting the existence of Pd NPs that have an average perimeter = 15 ± 4 nm (68%; with diameters in the range 1-10 nm) and Pd nanostructures (elongated, branched, starlike and concave, 30%) that have an average perimeter = 65 \pm 33 nm, see Fig. 4(d) and 2(c). These more complex structures

are likely to contain higher index crystal facets and will have a higher density of atomic steps and kinks.

Fig. 5 shows ADF-STEM images highlighting how the morphologies evolve with longer deposition times for the three different solutions, $t_{\rm dep}=150~{\rm s}~(\eta=-1.0~{\rm V})$. Statistical analysis of the morphology of the resulting structures in terms of circularity and perimeter is given in Fig. 6. Electrodeposition from the organic rich solution, Fig. 5(a), 6(a) and (c), shows the formation of very similar features to those obtained at $t_{\rm dep}=50~{\rm s}$. The density of Pd NPs has doubled from $3\times10^5~{\rm Pd}$ NPs per cm² to $6\times10^5~{\rm Pd}$ NPs per cm². The average circularity has slightly decreased to $0.6\pm0.1~{\rm but}$ the average perimeter = $8\pm3~{\rm nm}$ is very similar (n=250). Thus, the nanostructures can still be described generally as spherical NPs. There is again little evidence of Pd NP coalescence.

For electrodeposition carried out in equal volumes, solution 2, for $t_{\rm dep} = 150$ s, Fig. 5(b), the growth of both Pd NPs and elongated Pd nanostructures (rod-like) are observed, with the latter again appearing to be the result of coalescence of two or more near neighbouring Pd NPs. These changes are evident as the average circularity drops to 0.3 ± 0.1 and the average perimeter increases to 13 ± 3 nm (n = 165), as shown

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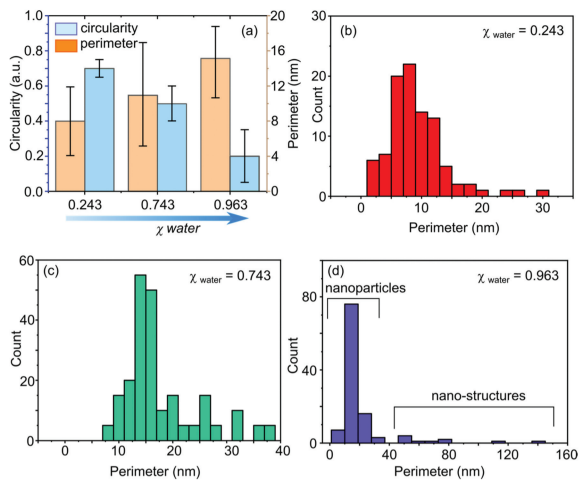


Fig. 4 Statistical analysis of Pd nanostructures obtained by electrodeposition, $E_{\rm dep} = \eta = -1.0 \text{ V}$ for $t_{\rm dep} = 50 \text{ s}$ in solution 1: 90% MeCN: 10% water v/v (\(\chi_{\text{water}} = 0.243\), solution 2: 50\% MeCN: 50\% water v/v (\(\chi_{\text{water}} = 0.743\)) and solution 3: 10\% MeCN: 90\% water v/v (\(\chi_{\text{water}} = 0.963\)). (a) Histogram displaying the change in Pd structure mean circularity and perimeter as function of \(\gamma_{water} \). (b-d) Histograms displaying a number count of perimeter lengths, extracted from the ADF-STEM images in Fig. 2, as a function of χ_{water}

in Fig. 6(a) and (d). Electrodeposition from the water rich solution 3, is shown in Fig. 5(c) after $t_{\rm dep}$ = 150 s. Under these conditions, the most prominent feature is the growth of a larger scale interconnected porous Pd structure, produced from agglomerated Pd nanostructures (fractal-like morphology⁵¹), which appear in some cases to have coalesced (Fig. 5(cii and iii)). Much smaller Pd NP and defected Pd nanostructures are also present on the surface of BDD TEM electrode (Fig. 5(ci)) accounting for the two different distributions as highlighted in Fig. 6(a), (b), (e) and (f). The circularity value for these smaller Pd NPs and nanostructures approaches an average of 0.2 ± 0.15 with an average perimeter of 16 \pm 4 nm (n = 130), i.e. the defected nanostructures are more prominent over the NPs. For those agglomerated nanostructures that could be measured using this approach, i.e. those that are still isolated from the larger interconnected macrostructure, e.g. as highlighted by the red arrow in Fig. 5 (ciii), an average circularity of 0.1 ± 0.05 is obtained and an average perimeter value of 50 \pm 5 nm (n = 80). These latter values should serve only as a guide as these structures are

three dimensionally (3D) complex and are likely to underestimate the exact geometry.

Violin plots comparing the evolution in circularity and perimeter of the electrodeposited nanostructures as the water content in the mixed solvent is increased (Fig. 2, 3 and 5) are presented in Fig. 7. Data was collated for all the analysed nanostructures for $t_{\text{dep}} = 50 \text{ s}$ and 150 s. ESI4 Fig. S6† shows histograms displaying a number count of perimeter lengths for all data collected. The black dots on Fig. 7 shows the average circularity or perimeter of the Pd nanostructures, for each χ_{water} , with the vertical line representing the mean \pm standard deviation. Fig. 7 clearly shows increasing water content results in both a deviation from circularity and a favouring of the growth of morphologically more complex Pd nanostructures, as seen by the increasing perimeter value and decreasing circularity values.

When electrodeposition was driven even longer, $t_{dep} = 300$ s, using the organic rich solution 1, Pd NP-like structures are still evident, as shown by the FE-SEM images in Fig. 8(a); although at this much lower resolution it is difficult to define the precise geometry. FE-SEM is now used in preference to

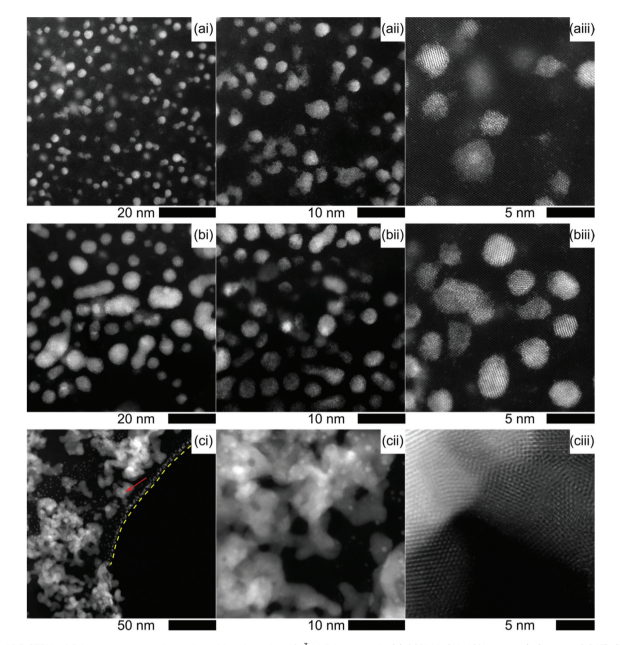


Fig. 5 ADF-STEM of Pd nanostructure electrodeposition from 1 \times 10⁻³ M Pd-acetate in (a) 90% MeCN : 10% water v/v (χ_{water} = 0.243), (b) 50% MeCN : 50% water v/v (χ_{water} = 0.743), (c) 10% MeCN : 90% water v/v (χ_{water} = 0.963) at an E_{dep} of η = -1.0 V for t_{dep} = 150 s. Note on (ci) the dotted line represents the edge of the BDD TEM electrode and the red arrow an indicative agglomerated nanostructure.

STEM due to the larger sized deposits that result. Electrodeposition from solution 2 reveals porous, almost "cauliflower-like" Pd micro-structures as displayed in Fig. 8(b). For water rich solution 3, two types of structures exist, one is large (greater than a $\mu m)$ Pd dendrites, whilst the second is much smaller Pd NPs and nanostructures, 20–500 nm diameter, as depicted in Fig. 8(c).

Discussion

Electrodeposition from the mixed solvent systems under high overpotential *i.e.* where the rate of electron transfer kinetics is sufficiently high that the process is likely to be diffusion-

limited, can be discussed qualitatively as follows. At the shortest deposition time ($t_{\rm dep}=50~{\rm s}$), in the organic rich solution, electrodeposition occurs according to $({\rm Pd}^{2+}({\rm MeCN})_x + 2{\rm e}^- \to {\rm Pd}^0)$, where x is indicative of the fact we do not know exactly how many MeCNs occupy the primary solvation shell. The Pd ions will be mostly liberated of acetate ligands, due to acetate interactions with the small volume of discrete water molecules present in the mixed solvent. The water molecules, if not associating with acetate are solvated by MeCN molecules. At Note, this process of acetate liberation from the Pd cation, is the reason why Pd electrodeposition, from the acetate salt is possible in MeCN rich solution but not a MeCN only solvent.

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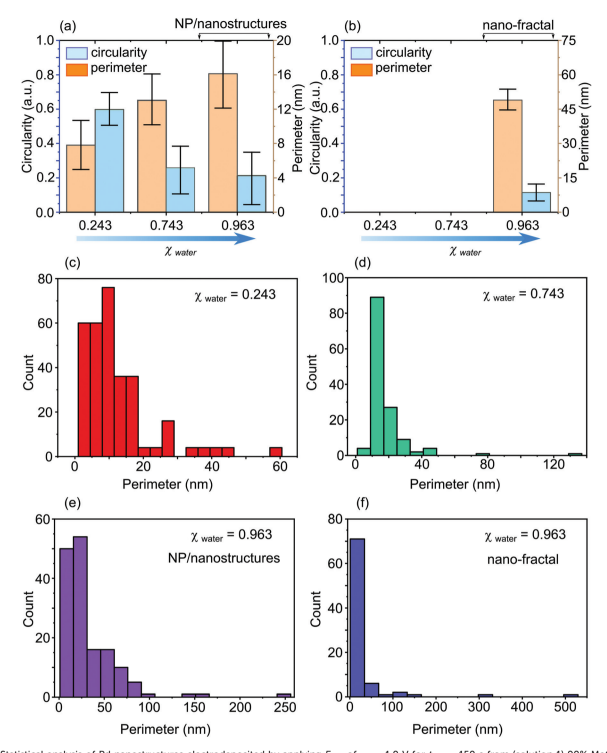


Fig. 6 Statistical analysis of Pd nanostructures electrodeposited by applying $E_{\rm dep}$ of $\eta=-1.0$ V for $t_{\rm dep}=150$ s from (solution 1) 90% MeCN:10% water v/v ($\chi_{\rm water}=0.243$), (solution 2) 50% MeCN:50% water v/v ($\chi_{\rm water}=0.743$), and (solution 3) 10% MeCN:90% water v/v ($\chi_{\rm water}=0.963$). (a and b) Histograms displaying the differences between Pd structure, mean circularity and perimeter as function of $\chi_{\rm water}$. For $\chi_{\rm water}=0.963$, both (a) NPs and nanostructures are represented, whilst (b) displays the agglomerated (fractal-like) structures. (c-f) Histograms displaying a number count of Pd structure perimeter lengths, extracted from the ADF-STEM images presented in Fig. 5 for (c) solution 1, (d) solution 2, and (e, and f) solution 3 for (e) NPs and defected nanostructures and (f) agglomerated (fractal-like) deposits.

After 50 s of nucleation and growth, single atoms, amorphous atom clusters and NPs are visible on the growth surface, with the latter in the majority at high density (3×10^5 Pd nano-

structures per cm²). Single atoms and amorphous atom clusters are the precursors to NP growth, as has been shown previously for Au electrodeposition at equally high overpotentials

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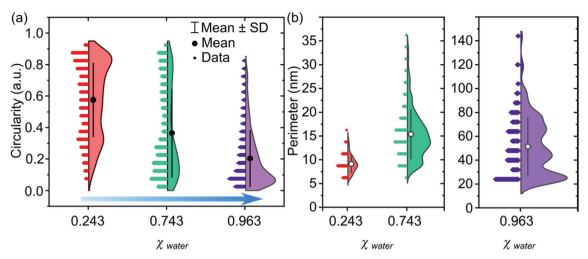


Fig. 7 A violin plot summary of the Pd nanostructures (a) circularity and (b) perimeter distributions as function of χ_{water} . $\chi_{\text{water}} = 0.963$ is represented on a different plot given the much larger perimeters recorded.

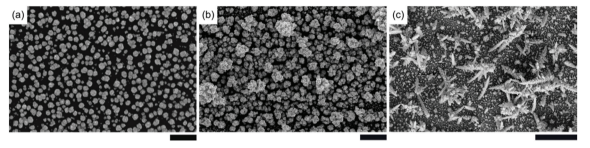


Fig. 8 FE-SEM images of Pd nano and micro-structure electrodeposition from 1×10^{-3} M Pd-acetate in((a) 90% MeCN: 10% water v/v ($\chi_{water} = 0.243$), (b) 50% MeCN: 50% water v/v ($\chi_{water} = 0.743$), (c) 10% MeCN: 90% water v/v ($\chi_{water} = 0.963$) at an E_{dep} of $\eta = -1.0$ V for $t_{dep} = 300$ s. (a and b) 400 nm and (c) 5 μ m.

in aqueous solution. Interestingly, the atoms, atom clusters and NPs for Au were observed at much shorter timescales (\sim 30 ms),³⁷ than seen here for Pd (50 s). Our results indicate that growth appears retarded in this system, mostly likely as a result of the majority solvent being MeCN. This is further confirmed by the negative increase in the Pd electrodeposition onset potential, by almost 0.5 V, by moving from a water rich solution to organic rich.

The formation of NPs at high density is not unsurprising given the high overpotential employed, resulting in many growth nuclei. Use of a low energy electrode surface, typical of carbon electrodes, favours NP (3D) rather than film (2D) formation at short times. ⁵² Electrodeposition for 150 s reveals a doubling in the number of Pd NPs, indicating that the growth of new nuclei is time-dependant "progressive in nature", and there is a concomitant small decrease in circularity, but the perimeters stay largely constant with time. Coalescence events between neighbouring NPs are limited, and direct attachment (Volmer–Weber growth mechanism) is favoured. ³⁷

We postulate the reason for the slow growth and predominate formation of NPs is due to MeCN adsorbing much more strongly on a Pd surface, than water, as a result of bonding interactions through both C and N atoms.⁵³ MeCN has been shown to adsorb on Pt in a similar fashion, with the C-C=N plane parallel to the surface. 54,55 Whilst MeCN coverage is not high enough to completely impede electrodeposition, we suggest it plays a significant role in slowing down the growth process, retarding (or blocking) both ad-atom mobility on the surface of growing nanostructures and coalescence events. On Pt, for example, MeCN adsorption has been shown to drastically decrease limiting current densities for the oxygen reduction reaction (ORR) and shift the onset potential to more negative values in aqueous (acidic) media.⁵⁴ Crystal orientation was also found to influence the strength of the MeCN-Pt interaction.54,55 As we expect similar behaviour for MeCN-Pd for future studies it would be of interest to simulate the strength of MeCN-Pd chemisorption on different crystal faces through density functional theory (DFT) simulations.

In solution 2, as the water content increases, the primary solvation shell of Pd is likely to contain both MeCN and water and electrodeposition proceeds according to $(Pd^{2+}(MeCN)_x(H_2O)_y + 2e^- \rightarrow Pd^0)$. As water content increases, MeCN-water H-bonding interactions also increase as seen from the Raman data. Under such circumstances, a

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competition between MeCN solvation by water molecules and MeCN adsorption on Pd is also likely. Coupled with the presence of water molecules within the solvation shell, the net result is a decrease in the number of Pd sites which are occupied by adsorbed MeCN and electrodeposition (after 50 s) showing not only NPs but now the presence of defected and elongated nanostructures. The latter likely result from NP–NP coalescence events as demonstrated by the high-resolution STEM images Fig. 3(b). Such structures also dominate at longer deposition times (150 s), whilst at 300 s agglomeration/aggregation of these nanostructures is evident.

Coalescence is due to either the growth (especially in the lateral direction) of closely spaced NPs or the movement of NPs over the electrode surface. Coalescence driven by Ostwald ripening is also a possibility, especially given the observation of high energy sites (atoms, atom clusters and small defected NPs) in the first 50 s of growth. Pd growth rates are increased due to the presence of water, as growth is no longer as highly constrained in the lower water content solutions. This results in larger sized NPs for a fixed growth time, which can grow into one another, aggregate, then coalesce. Whilst NP mobility over carbon surfaces is well documented during electrodeposition in aqueous solutions, the interaction strength between MeCN and an oxygen terminated BDD surface, and its impact on NP mobility, is unknown in the literature. DFT simulations are required for future studies.

In water rich solution, water molecules occupy the solvation shell of the Pd cations, due to hydrolysis of the acetate groups, hence electrodeposition proceeds via $Pd^{2+}(H_2O)_v + 2e^- \rightarrow Pd^0$. Raman indicates that the minority MeCN are H-bonded to the majority water molecules and thus minimal MeCN adsorption is also expected on the growing Pd nanostructures resulting in unrestricted growth of Pd on Pd in many directions. Under these conditions, and with a high driving overpotential, anisotropic growth is evident, especially at the longest times with the appearance of the Pd microscale dendrites (300 s). Dendrites are often associated with diffusion-limited growth and is dependent on Pd ad-atom mobility as well as the interplay between the size of the concentration gradient and the diffusion properties of the ion of interest. 56-58 Increasing solution viscosity has been shown to promote dendritic growth.⁵⁸ Here increasing water content in MeCN increases the viscosity of the resulting solution.⁵⁹ At the shorter times NPs, elongated and concave nanostructures are present but now also branched and star-like Pd nanostructures form. With further growth time, nanostructures agglomerate and coalesce, new growth centres continue to form on the Pd nanostructures, before transforming into dendritic-type structures.

Conclusions

The solvent plays a significant role in electrodeposition. Here we show how by using a mixed solvent system (MeCN-water) greater morphological control over electrodeposited (Pd) nanostructures can be exerted, switching from NPs to dendrites as

the water content increases. We also demonstrate the versatility of electron transparent BDD electrodes for making combined electrochemical and high resolution STEM measurements of metal electrodeposition.

In MeCN-rich solutions, under high overpotential deposition conditions, single atoms, amorphous atom clusters and spherical crystalline Pd NPs, the latter in the majority, were formed. MeCN is known to strongly adsorb to Pt and Pd surfaces and plays a critical role in retarding electrodeposition such that these structures were observed on fairly lengthy timescales (50 s), compared to seconds (and smaller) for aqueous solutions. Increasing water content resulted in an increasing growth rate and the observation of larger sized nanostructures and the formation of elongated structures arising from aggregation and coalescence events. Growth in a water rich environment, where the MeCN are now bound to majority water molecules as shown by Raman, resulted in the unconstrained growth of Pd nanostructures. At 50 s star-branched- and concave structures formed, which with further growth over 300 s, led to aggregated porous nanostructures (fractal-like) and dendritic structures.

The ability to control nanostructure morphology simply by modifying the properties of the mixed solvent is incredibly useful. For example, dendrite formation is a huge problem in metal battery (e.g. Li, Zn) chemistry^{60,61} but extremely useful for some applications, such as surface enhanced Raman scattering (SERS)⁶² and superhydrophobicity.⁶² By using a MeCN or water rich solution we have shown how dendrite formation can be suppressed or enhanced simply by changing mixed solvent composition.

Conflicts of interest

The authors declare no competing financial interest. There are no conflicts to declare.

Acknowledgements

JVM thanks the Royal Society for an Industry Fellowship (INF/R1/180026). HEMH thanks the ESPRC (EP/N035437/1), Astra-Zeneca, and the University of Warwick for funding. JVM and HEMH thank Element Six Ltd for the provision of high-quality BDD samples, Mareike Herrmann for lasering of BDD electrodes (Department of Chemistry, University of Warwick) and Prof. Mark Newton (Department of Physics, University of Warwick) for useful discussions. We are grateful to Steve York and Steve Hindmarsh from the Electron Microscopy Research Technology Platform, University of Warwick, for their initial assistance with the FE-SEM imaging experiments.

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