Smooth crack-free targets for nuclear applications produced by molecular plating

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1. Introduction

Targets are an essential component in experimental nuclear science as a source of stationary nuclei, e.g., for atomic and nuclear spectroscopy or nuclear reaction studies [1–3]. Generally, targets should be chemically pure, uniform, homogeneous, crack-free over the area of interest, and well adherent to the deposition substrate. Among the several techniques available to prepare thin layers to be used as nuclear targets [4–8], molecular plating (MP) is one of the most effective and most successfully applied ones [9,10]. This technique, based on the electrodeposition of the element of interest from an organic medium either in the constant current or in the constant voltage mode, is usually chosen because of the possibility to quickly produce deposits of acceptable uniformity and adherence with quantitative yield. Recently Garcia et al. [11,12] applied polymer assisted deposition (PAD), developed by Jia et al. [13] as an alternative method to prepare thin films of metal oxides, to produce nuclear targets. PAD has been claimed to yield targets with better uniformity and homogeneity than MP. In the present work, we demonstrate that thanks to an improved understanding of the electrochemistry of MP and to more extensive surface analysis of the produced surfaces, radiographic imaging for surface homogeneity inspection, atomic force microscopy for surface roughness evaluation, and scanning electron microscopy for surface morphology investigation. The results allowed identifying the optimum parameters for the production of smooth and crack-free targets by means of molecular plating. The smoothest layers, which had an average RMS roughness of ca. 20 nm and showed no cracks, were obtained using 0.22 mM [Nd(NO3)3·6H2O] plated from N,N-dimethylformamide at current densities in the range of 0.3–0.7 mA/cm² on the smoothest deposition substrate available.

Abstract

The production process of smooth and crack-free targets by means of constant current electrolysis in organic media, commonly known as molecular plating, was optimized. Using a Nd salt, i.e., [Nd(NO3)3·6H2O], as model electrolyte several constant current density electrolysis experiments were carried out to investigate the effects of different parameters, namely the plating solvent (isopropanol and isobutanol mixed together, pyridine, and N,N-dimethylformamide), the electrolyte concentration (0.11, 0.22, 0.44 mM), the applied current density (0.17, 0.3, 0.7, and 1.3 mA/cm²), and the surface roughness of the deposition substrates (12 and 24 nm). Different environments (air and Ar) were used to dry the samples and the effects on the produced layers were investigated. The obtained deposits were characterized using γ-ray spectroscopy for determining Nd deposition yields, X-ray photoelectron spectroscopy for chemical analysis of the produced surfaces, radiographic imaging for surface homogeneity inspection, atomic force microscopy for surface roughness evaluation, and scanning electron microscopy for surface morphology investigation. The results allowed identifying the optimum parameters for the production of smooth and crack-free targets by means of molecular plating. The smoothest layers, which had an average RMS roughness of ca. 20 nm and showed no cracks, were obtained using 0.22 mM [Nd(NO3)3·6H2O] plated from N,N-dimethylformamide at current densities in the range of 0.3–0.7 mA/cm² on the smoothest deposition substrate available.

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block covered with a circular Ti foil fixed with a PEEK ring. A constant temperature of 16 °C was maintained in the cell by water-cooling the Ti blocks, which were in physical contact with the electrodes.

2.2. Solvents used for molecular plating

Different solvents were used to perform the MPs: isopropanol, isobutanol, pyridine, and N,N-dimethylformamide (DMF). Isopropanol and isobutanol were always used together as a mixture (3% isopropanol, 97% isobutanol), hereafter referred to as IP+IB. Table 1 shows some physical properties of relevance of the used solvents [16].

The data reported in Table 1 show DMF to have the highest boiling point, conductivity, and relative permittivity; its viscosity is instead the lowest one. Pyridine has the lowest relative permittivity, a viscosity approaching that of DMF and a boiling point quite close to the one of the IP+IB mixture. The physical properties of the IP+IB mixture can be approximated to those of pure isobutanol, which makes up 97% by volume of the mixture. IP+IB has the highest viscosity and the lowest conductivity.

2.3. Deposition substrates

The deposition substrates used in the present work are listed in Table 2. Their surface roughnesses, evaluated by AFM (see Section 2.5 in [14]) after the cleaning procedure, are reported as average root mean square values (RMS). To facilitate comparison of the results reported here and previously in [14], we keep the labelling of our substrate types as in [14].

T13 was a circular 50 μm-thick Ti foil cut from a bigger foil (Goodfellow). T15 was produced by coating a 300-μm thick, mechanically polished Si wafer with 100 nm metallic Ti. The substrates used for deposition in the cell had different surface roughnesses: larger, corresponding to a more defective surface (T13,Fig. 2a), and smaller, corresponding to a less defective surface (T15,Fig. 2b).

2.4. Molecular plating at constant current density

Before any deposition, both the anode (Pd foil) and the cathode (Ti foil) were cleaned with 6 M HCl (1×3 mL washing), water (3×3 mL washings), and isopropanol (1×3 mL washing). Only T15 substrates were cleaned using HCl with a lower concentration (i.e., 0.6 M) in order to avoid corrosion-induced increase in surface roughness, as verified in [14]. For the MPs, 0.338, 0.675, and 1.350 g of [Nd(NO3)3·6H2O] were dissolved in 20 mL 0.1 M HNO3 to give final solutions of three different concentrations. From each solution an aliquot of 100 μL was added to a mixture of 1 mL isopropanol (Fisher Chemical) and 34 mL isobutanol (Applichem), or to 35 mL of pyridine (Merck) or DMF (Merck), finally yielding three different solutions of Nd3+ in the cell; these are referred to as low concentration (LC, 0.11 mM), medium concentration (MC, 0.22 mM), and high concentration (HC, 0.44 mM). In the cell the concentration of HNO3 was 0.3 mM. The plating solutions were stirred by means of a quartz tip ultrasonic stirrer (Bandelin Sonopuls HD 2070) operated at 30% power pulse. MP experiments were always carried out for 3 h by applying a constant current of 6 mA, corresponding to a current density of 0.7 mA/cm². Three-hours long MPs were also performed by testing different constant currents, i.e., 1.5, 3, or 12 mA. The corresponding current densities were 0.17, 0.3, and 1.3 mA/cm² (Table 3).

2.5. Atomic force microscopy (AFM) studies

2.5.1. Experiments

The surface roughness of the deposited layers, i.e., the measure of the texture of a surface quantified by the vertical deviations of a real surface from its ideal planar form, was investigated by using atomic force microscopy (AFM) (MFP 3D Asylum Research). AFM investigations were performed in air and in liquid.

The tapping-mode AFM analysis in air was the same for all the produced targets: ten (5×5) μm² and one (25×25) μm² images were recorded for each target. From each of the ten (5×5) μm² images two root mean square (RMS) values were evaluated using a (1.5×1.5) μm² mask in two different positions inside the image. From each of the (25×25) μm² images, twenty RMS values were evaluated using a (1.5×1.5) μm² mask in twenty different positions of the image. In the case of samples presenting surfaces with cracks, the masks were used only in the crack-free areas, as the cracks do not belong to the surface and hence do not define the real roughness of the deposited layers. The final surface roughness value of each of the image sizes was evaluated for each target as the arithmetic mean of all the RMS values measured for that specific image size (i.e., RMS).

2.5.2. Results

2.5.2.1. Layer-growth studies from DMF solution. The growth mechanism of the layers was studied as a function of plating time by measuring the RMS of air-dried targets. Twelve targets were produced using different deposition times: the first target was produced after 15 min plating and the subsequent ones after increments of 15 min up to 180 min. The targets were prepared using a 0.22 mM Nd salt (i.e., MC), T15 deposition substrates, at a current density of 0.7 mA/cm², and DMF as plating solvent. The growth study was restricted to DMF because it showed very promising results, as we briefly reported in [17] where some preliminary tests with this solvent were described.

In Fig. 3 the RMS trends of the growing layer show an almost constant behaviour (average value ca. 14 nm) during the whole period of deposition (3 h); the surface roughness does not undergo major variations. This is in contrast to similar studies performed...
The solvents used for the platings were IP + IB, pyridine, and DMF; the deposition substrates were Ti3 foils. The RMS values of the analysis are reported in Table 3.

Table 3 clearly demonstrates that, independently of the chosen drying environment, the roughest samples are those obtained using pyridine, whereas those produced in DMF are the smoothest. No differences in RMS values within experimental uncertainties were found for each kind of sample dried in air or in Ar. The analysis of the recorded AFM images showed that samples characterized by cracking of the layers present surface troughs regardless of whether they were dried in air or Ar.

Fig. 5 shows a (25×25) μm² image of the DMF sample after drying in the inert environment.

Unlike the pyridine and IP + IB layers, characterized by severe surface cracking (see, e.g., Fig. 3a in [19] for targets made from IP + IB on Ti3 substrates), the DMF sample shows almost no cracks, except for those mostly arising from the surface defects (i.e., white outgrowths in Fig. 5).

2.5.2.2. Influence of the plating solvent and of the drying environment. AFM studies in air were performed on targets produced from MC solution, after drying for one month either in air or in a glove-box where an Ar inert environment was maintained.

2.5.2.3. Influence of the electrolyte concentration. Air-dried HC, MC, and LC targets produced using Ti5 as deposition substrates and DMF as plating solvent were investigated in air. The results of the analysis reported in Table 3 show that the HC target is much rougher than the MC and LC targets, which have identical roughness. Moreover, Table 3 shows that DMF MC targets prepared on Ti5 substrates are smoother than the same targets prepared on Ti3 foils.

We notice that the recorded AFM images revealed no cracks in the surfaces of the HC, MC, and LC targets plated from DMF.

2.5.2.4. Influence of the current density. AFM analyses in air were performed on air-dried MC targets produced using different constant current densities, namely 0.17, 0.3, and 1.3 mA/cm². Ti5 was used as deposition substrate, and DMF as plating solvent. The RMS values of the analysis are reported in Table 3, whereas the corresponding (25×25) μm² images of the deposits are shown in Fig. 6.

Table 3 shows that only the intermediate current density of 0.3 mA/cm² gives RMS values similar to those recorded for the 0.7 mA/cm² experiments. Both higher (i.e., 1.3 mA/cm²) and lower (i.e., 0.17 mA/cm²) constant current densities result in much rougher final deposits. In particular, the roughest deposit is obtained with 0.17 mA/cm². These results are well evidenced by the (25×25) μm² images of the 0.17 and 1.3 mA/cm² samples shown in Fig. 6, where both surfaces present evident outgrowths (i.e., white spots in the figures), which are bigger in the case of the targets made at 1.3 mA/cm², but not covering the entire analyzed area (c), and smaller but spread over the entire surface in the case of the targets made at 0.17 mA/cm² (a). No surface cracks were noted for any of the investigated samples prepared with DMF.

2.5.2.5. Liquid AFM study from IP + IB solution. Fig. 7 shows the results of a liquid AFM study (already briefly reported in [17]), which was performed on a MC target produced using Ti3 as deposition substrate, and IP + IB as plating solvent. At the end of the deposition, the cell was top-sealed, the anode was removed, and the solvent was extracted till only the cathode surface was covered by liquid. Then, the PEEK body of the cell was removed, leaving the cathode deposit covered in liquid: an AFM analysis in contact mode was performed recording one (25×25) μm² image (Fig. 7a). After the liquid was removed from the cathode assembly and the target dried, a tapping-mode AFM analysis was performed and a (25×25) μm² image was recorded (Fig. 7b). Both images were elaborated using the WSxM 5.0 version developed by Nanotech Electronica [18].
The 3D AFM images of Fig. 7 indicate that the cracking of the surfaces develops after the removal of the liquid and the drying of the deposited layer.

2.6. Scanning electron microscopy (SEM) studies

2.6.1. Experiments

The morphology of the targets was studied using scanning electron microscopy (SEM) (Philips XL30). The microscope was operated at 20 kV. The pressure inside the vacuum chamber was 5 mPa. Backscattered and secondary electrons were detected. A magnification factor of 1000× was used.

2.6.2. Results

Fig. 8 shows 1000× SEM images of three MC samples produced using the following deposition substrates and plating solvents: (a) Ti3 and pyridine, (b) Ti3 and DMF, (c) Ti5 and DMF. All samples were air-dried. Among the several SEM images recorded, these particular ones were chosen for display here because of the important surface morphology information they convey.

Fig. 8a shows a large-area image of an air-dried pyridine MC target. The surface of the sample suffers from severe cracking and it is visibly rough.

Figs. 8b and c show an important effect of the deposition substrate (Ti3 vs. Ti5) on the surface morphology of the deposits. These images indicate that MC targets prepared using DMF and a Ti3 substrate (Fig. 8b) exhibit cracks in the layer, caused mainly by the grown surface defects, as already shown in Fig. 5. If, instead, a smoother deposition substrate is used, i.e., Ti5, MC targets produced using DMF (Fig. 8c) show no surface cracks (see also Fig. 4).

In contrast, MC experiments using IP+IB (see Fig. 2b in [14])

![RMS vs. time trend of the layer-growth study of the air-dried targets produced using 0.22 mM Nd salt with DMF and Ti5 deposition substrates.](image)

![Layer-growth studies from DMF solution. (5×5 μm² topography AFM image of the sample plated for 180 min using Ti5 substrate and MC plating solution in DMF.](image)

### Table 3

Overview of relevant molecular plating parameters. Average root mean square (RMS) values obtained from the AFM analyses of the samples investigating the influence of plating solvent, drying environment, electrolyte concentration, and current density on the surface roughness of the produced layers.

<table>
<thead>
<tr>
<th>Substrate type</th>
<th>Solvent</th>
<th>Drying environment</th>
<th>Concentration</th>
<th>Current density (mA/cm²)</th>
<th>Scan size (μm²)</th>
<th>RMS (nm)</th>
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<tr>
<td>Ti3</td>
<td>Pyridine</td>
<td>Air</td>
<td>MC</td>
<td>0.7</td>
<td>(5×5)</td>
<td>242±140</td>
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<td>(25×25)</td>
<td>372±190</td>
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<tr>
<td>Ti3</td>
<td>Pyridine</td>
<td>Ar</td>
<td>MC</td>
<td>0.7</td>
<td>(5×5)</td>
<td>320±160</td>
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<tr>
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<td>(25×25)</td>
<td>350±180</td>
</tr>
<tr>
<td>Ti3</td>
<td>IP+IB</td>
<td>Air</td>
<td>MC</td>
<td>0.7</td>
<td>(5×5)</td>
<td>150±40</td>
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<td></td>
<td></td>
<td>(25×25)</td>
<td>190±70</td>
</tr>
<tr>
<td>Ti3</td>
<td>IP+IB</td>
<td>Ar</td>
<td>MC</td>
<td>0.7</td>
<td>(5×5)</td>
<td>140±60</td>
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<td></td>
<td>(25×25)</td>
<td>180±80</td>
</tr>
<tr>
<td>Ti3</td>
<td>DMF</td>
<td>Air</td>
<td>MC</td>
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<td>(5×5)</td>
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<td>(25×25)</td>
<td>130±40</td>
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<tr>
<td>Ti3</td>
<td>DMF</td>
<td>Ar</td>
<td>MC</td>
<td>0.7</td>
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<td>110±50</td>
</tr>
<tr>
<td>Ti3</td>
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<td>Air</td>
<td>HC</td>
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<td>79±14</td>
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<tr>
<td>Ti3</td>
<td>DMF</td>
<td>Air</td>
<td>MC</td>
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<td>(5×5)</td>
<td>17±5</td>
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<td>(25×25)</td>
<td>18±9</td>
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<tr>
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<td>DMF</td>
<td>Air</td>
<td>LC</td>
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<td>18±8</td>
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<td>Ti3</td>
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<td>Air</td>
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<td>(25×25)</td>
<td>130±25</td>
</tr>
<tr>
<td>Ti3</td>
<td>DMF</td>
<td>Air</td>
<td>MC</td>
<td>0.3</td>
<td>(5×5)</td>
<td>10±4</td>
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<tr>
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<td></td>
<td></td>
<td>(25×25)</td>
<td>12±5</td>
</tr>
<tr>
<td>Ti3</td>
<td>DMF</td>
<td>Air</td>
<td>MC</td>
<td>1.3</td>
<td>(5×5)</td>
<td>70±30</td>
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<td></td>
<td></td>
<td></td>
<td>(25×25)</td>
<td>80±22</td>
</tr>
</tbody>
</table>
or pyridine showed the presence of cracks in the deposited layers also when Ti5 substrates were used.

2.7. Radiographic imaging (RI) studies

2.7.1. Experiments

Radioactive targets were produced by performing MPs of Nd solutions containing a radioactive tracer of the element to be plated (Nd: $^{147}$Nd, $\beta^-$, $t_{1/2} = 10.98$ d, $E_\gamma = 91$ keV). The tracer was produced in a neutron capture reaction ($n,\gamma$) on $^{146}$Nd present in the used Nd of natural isotopic composition (natNd) by irradiating an aliquot of 100 $\mu$L of the stock solution with thermal neutrons in the TRIGA Mainz research reactor. The homogeneity of such targets was inspected by using radiographic imaging (RI), employing a FUJIFILM FLA 7000 equipped with reusable imaging plates and a 650 nm laser for the reading process. To verify the homogeneity of the target, the image was overlaid by a grid of square spots with an area of 0.25 mm$^2$ each. The spatial resolution for $\gamma$-emitting nuclides was <200 $\mu$m [19].

2.7.2. Results

Fig. 9a shows the 3D graph of a sample obtained using IP + IB as plating solvent and Ti3 as deposition substrate. Fig. 9b displays the graph of a sample obtained using DMF and Ti5.

The images show a definite improvement in surface homogeneity of the deposited layers once DMF is chosen as plating solvent and Ti5 as deposition substrate. Experiments showed that better homogeneity of the surfaces is achieved also when DMF is used instead of pyridine in the preparation of MC targets on Ti3 foils.

2.8. X-ray photoelectron spectroscopy (XPS) studies

2.8.1. Experiments

X-ray photoelectron spectroscopy (XPS) was used to perform the chemical surface characterization of the samples by applying Al K$_\alpha$ and Mg K$_\alpha$ radiations. The X-ray source was operated at 10 kV anode voltage and a current of 10 mA. The pass-energy of the hemispherical energy analyzer was 50 eV. The pressure inside the ultra-high vacuum chamber was in the range of $(1.2 - 2.0) \times 10^{-7}$ Pa. The analyzed samples were six MC targets produced using IP + IB, pyridine, and DMF as plating solvents, and Ti3 foils as deposition substrates. Two targets were produced in each solvent: one was dried in air, while the other was dried inside a glove-box, where an inert environment was assured by a constant Ar pressure.

![Fig. 5](image1.png)

Fig. 5. Influence of the plating solvent and of the drying environment. (25x25) $\mu$m$^2$ topography AFM image of the sample produced using MC plating solution, Ti3 substrate, DMF, and dried in an Ar inert environment.

![Fig. 6](image2.png)

Fig. 6. Influence of the current density. (25x25) $\mu$m$^2$ topography AFM images of samples produced using MC plating solution in DMF and a Ti5 substrate; deposits obtained after galvanostatic electrolysis at current densities of 0.17 mA/cm$^2$ (a), 0.3 mA/cm$^2$ (b), or 1.3 mA/cm$^2$ (c).
(50 Pa below atmospheric pressure, $O_2<1$ ppm, $H_2O<1$ ppm). We refer to [14] for details about the drying procedure in an inert environment and the subsequent XPS analysis. Drying in different environments was investigated to probe whether reactive or inert environments like, respectively, air or Ar cause different surface chemical reactions to take place, the products of which might be identified by XPS.

After the first surface investigation, the samples underwent 10 min sputtering using 4 kV $Ar^+$ ions, followed by further XPS analysis.

The binding energies of the XPS lines of interest were determined by fitting the recorded XPS spectra. The analyses were performed using the CasaXPS software (version 2.3.15), with which the background was subtracted adopting the Shirley algorithm, and the peaks were fitted using the Gaussian/Lorentzian product formula. As the aim of the present work was mainly to investigate the surface chemistry of the samples after drying in different environments, a detailed chemical surface analysis of the
deposits was performed only for C, which was known from [14]
to be the main element composing the outer layer of the produced
samples. The C 1s spectra are therefore the only ones to be shown
with multicomponent fitting analysis, which was always modelled
by fixing the separations in binding energy, the intensity ratios,
and the full widths at half maxima (FWHM) of the different C 1s
components of IP + IB, DMF, and pyridine. For the IP + IB and DMF
samples dried both in air and in Ar environment, the analysis
before sputtering required correcting the binding energies for the
surface charging using one of the components of the C 1s lines: the
284.9 eV component [20] for IP + IB, and the 285.8 eV component
[21] for DMF. After sputtering, the imprecise fittings of the
reference C 1s peaks required the use of a higher binding energy
peak (ca. 289 eV) present in the same C 1s spectra to correct for
the surface charge effect. For the air- and Ar-dried pyridine
samples, instead, the correction of the binding energies was
performed both before and after sputtering using the intense
N 1s line of pyridine at 400.2 eV [22]. The experimental uncer-
tainty of the binding energies is ±0.1 eV.

2.8.2. Results

The XPS analyses of the six MC targets never showed the presence
of Ti signals, indicating that all the deposited layers are thick.
The Nd analysis of the samples required the use of Mg Kα
radiation to avoid an interference with the Auger lines of oxygen in
the binding energy region of Nd 3d. The Nd 3d spectra of all the
samples dried in Ar environment (not shown) were identical to
those of the air-dried targets, except for smaller intensities of the
recorded peaks.

Differences among the recorded XPS signals of the IP + IB, DMF
and pyridine spectra were noted only for the C 1s lines of the air-
and Ar-dried samples.

2.8.2.1. Samples prepared from IP + IB. The XPS analyses of the air-
and Ar-dried IP + IB samples showed the presence of the following
elements: C, O, Pd, and Nd. The corresponding XPS signals were:
C 1s, O 1s, Pd 3d, Nd 3d, and Nd 4d. Fig. 10 shows the spectra of Nd
3d (a) and C 1s (b) of the sample dried in air and Fig. 10c the C 1s
spectra of the sample dried in Ar environment. The model applied
for the multicomponent fitting analysis of the C 1s peaks took into
account only the C 1s components of isobutanol, as isopropanol
corresponds to just 3% of the total volume of the solvent.

According to literature [20], the C 1s components of isobutanol
have binding energies at 284.9 eV, 285.0 eV, and 286.5 eV, with an
intensity ratio of 2:1:1. Table 4 shows the ratios of the total peak
areas C 1s/Nd 3d, O 1s/C 1s, and the ratios between the sum of the
peaks fitting a certain spectral component and a total peak area,
like the ratios reported as C 1s (ca. 285 eV)/Nd 3d, C 1s (ca.
286 eV)/Nd 3d, and C 1s (ca. 289 eV)/Nd 3d. The ratios were
evaluated before and after 10 min sputtering.

The measured binding energy of the Nd 3d5/2 peak is 982.8 eV
(Fig. 10a) and can be compared to the literature value of 983.1 eV
[23] for Nd3+. This indicates that Nd is present as Nd3+ in the
deposited layer (possibly as carboxylate, oxide, or hydroxide) [14].

The C 1s spectra reported in Figs. 10b and c show the presence
of two components: the C 1s peaks at ca. 285.0 eV, which
remarkably decrease upon Ar + sputtering, are due to physisorbed
aliphatic carbon of the mixed isopropanol/isobutanol solvent. The
peaks at the higher binding energy (i.e., 289.9 and 289.6 eV for air-
and Ar-dried samples, respectively) can be attributed, according to
literature, to COOM – metal carboxylate – (289.0 eV [23]) groups
formed during the plating process, which are chemisorbed, as
follows from the observation that they are not easily removed by
the sputtering. This shows that the solvent undergoes electrolysis
during the MP process and becomes oxidized.

For a more detailed analysis of the Nd 3d and C 1s spectra of the
IP + IB samples we refer to [14].

The spectra of Fig. 10, as well as the C 1s/Nd 3d 3d ratios reported
in Table 4, show that for both air- and Ar-dried samples the
intensity of the Nd 3d signal increases, while that of C 1s decreases
when the surface is sputtered with Ar + for 10 min. This result
shows that the Nd deposit is covered with a layer of solvent.

A consequence of drying the samples in air can be inferred from
the different intensity ratios of the two C 1s components at 0 min
sputtering in Figs. 10b and c: the 1:1 ratio of the Ar-dried sample
becomes almost 1:2 after drying in air. From the analysis of the
C 1s (ca. 285 eV)/Nd 3d and C 1s (ca. 289 eV)/Nd 3d ratios it is
evident that while the intensity of the chemisorbed C 1s (ca.
289 eV) component is not affected by the method of drying, the
physisorbed one (ca. 285.0 eV) decreases when the sample is dried
in air. As a consequence more intense Nd 3d signals, hence smaller
C/Nd ratios (see Table 4), are recorded for the air-dried sample.

2.8.2.2. Samples prepared from DMF. As regards the XPS analyses of
the targets prepared in DMF, the spectra of both air- and Ar-dried
samples showed the presence of the following elements: C, N, O,
and Nd. The corresponding XPS signals were: C 1s, N 1s, O 1s, Nd
3d, and Nd 4d. No signals of Pd were detected. Fig. 11 shows the
spectra of Nd 3d (a) and C 1s (b) of the sample dried in air, and
Fig. 11c the C 1s spectra of the sample dried in Ar environment.
According to literature [21], the C 1s components of DMF have binding energies at 285.8 eV, and 288.8 eV with an intensity ratio of 2:1.

The XPS spectra of the DMF samples (Fig. 11) are quite similar to those of the IP⁺IB samples (Fig. 10). The measured binding energy of the Nd 3d₃/₂ peak (i.e., 983.6 eV, Fig. 11a) confirms that Nd is deposited as Nd³⁺ also in DMF (possibly as carboxylate, oxide, or hydroxide). The C 1s spectra reported in Figs. 11b and c show again the presence of two components: the components of the physisorbed solvent at ca. 286 eV and 289 eV, and the components of chemisorbed species at 290.2 eV in air, and at 289.4 eV in Ar environment. As in the case of IP⁺IB, the C 1s peak of the chemisorbed species can be attributed to possible COOM⁻–metal carboxylate– groups formed during the plating process. The produced samples are again characterized by an outer layer of solvent which covers the Nd deposit, as indicated by the spectra of Fig. 11, as well as by the ratios C 1s/Nd 3d reported in Table 4. These ratios show that, for both air- and Ar-dried samples, the intensity of the Nd 3d signal increases with sputtering time, while that of C 1s decreases.

As in the case of MPs in IP⁺IB, the difference between drying in air or Ar is reflected by the smaller C 1s/Nd 3d ratio in air than in Ar, i.e., 0.05 instead of 0.10 (Table 4). This difference is also confirmed by the larger value of the C 1s (ca. 286 eV)/Nd 3d ratio in Ar than in air. Therefore, also in the case of DMF, the 1:1 ratio of the C 1s signals of the Ar-dried sample becomes almost 1:2 after drying in air. The C 1s (ca. 289 eV)/Nd 3d ratios are very similar both in air and in Ar, indicating that the chemisorbed products are not considerably affected by drying in different environments.

### Table 4

<table>
<thead>
<tr>
<th>Sample</th>
<th>Ratios of XPS peak areas</th>
<th>Air 0 min sputtering</th>
<th>Ar 0 min sputtering</th>
<th>Air 10 min sputtering</th>
<th>Ar 10 min sputtering</th>
</tr>
</thead>
<tbody>
<tr>
<td>IP⁺IB</td>
<td>C 1s/Nd 3d</td>
<td>0.04</td>
<td>0.015</td>
<td>0.11</td>
<td>0.020</td>
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<tr>
<td></td>
<td>O 1s/C 1s</td>
<td>6</td>
<td>10</td>
<td>4</td>
<td>9</td>
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<tr>
<td></td>
<td>C 1s (ca. 285 eV)/Nd 3d</td>
<td>0.014</td>
<td>0.0026</td>
<td>0.05</td>
<td>0.008</td>
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<tr>
<td></td>
<td>C 1s (ca. 289 eV)/Nd 3d</td>
<td>0.026</td>
<td>0.012</td>
<td>0.04</td>
<td>0.011</td>
</tr>
<tr>
<td>DMF</td>
<td>C 1s/Nd 3d</td>
<td>0.05</td>
<td>0.011</td>
<td>0.10</td>
<td>0.014</td>
</tr>
<tr>
<td></td>
<td>O 1s/C 1s</td>
<td>6</td>
<td>15</td>
<td>5</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>C 1s (ca. 286 eV)/Nd 3d</td>
<td>0.022</td>
<td>0.0018</td>
<td>0.07</td>
<td>0.005</td>
</tr>
<tr>
<td></td>
<td>C 1s (ca. 289 eV)/Nd 3d</td>
<td>0.03</td>
<td>0.009</td>
<td>0.028</td>
<td>0.009</td>
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<tr>
<td>Pyridine</td>
<td>C 1s/Nd 3d</td>
<td>0.9</td>
<td>0.14</td>
<td>1.1</td>
<td>0.20</td>
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<tr>
<td></td>
<td>O 1s/C 1s</td>
<td>1.3</td>
<td>1.4</td>
<td>1.2</td>
<td>1.4</td>
</tr>
<tr>
<td></td>
<td>C 1s (ca. 286 eV)/Nd 3d</td>
<td>0.8</td>
<td>0.13</td>
<td>1.0</td>
<td>0.17</td>
</tr>
<tr>
<td></td>
<td>C 1s (ca. 289 eV)/Nd 3d</td>
<td>0.10</td>
<td>0.015</td>
<td>0.12</td>
<td>0.021</td>
</tr>
</tbody>
</table>

The XPS analyses of the DMF samples (Fig. 11) are quite similar to those of the IP⁺IB samples (Fig. 10). The measured binding energy of the Nd 3d₃/₂ peak (i.e., 983.6 eV, Fig. 11a) confirms that Nd is deposited as Nd³⁺ also in DMF (possibly as carboxylate, oxide, or hydroxide). The C 1s spectra reported in Figs. 11b and c show again the presence of two components: the components of the physisorbed solvent at ca. 286 eV and 289 eV, and the components of chemisorbed species at 290.2 eV in air, and at 289.4 eV in Ar environment. As in the case of IP⁺IB, the C 1s peak of the chemisorbed species can be attributed to possible COOM⁻–metal carboxylate– groups formed during the plating process. The produced samples are again characterized by an outer layer of solvent which covers the Nd deposit, as indicated by the spectra of Fig. 11, as well as by the ratios C 1s/Nd 3d reported in Table 4. These ratios show that, for both air- and Ar-dried samples, the intensity of the Nd 3d signal increases with sputtering time, while that of C 1s decreases.

As in the case of MPs in IP⁺IB, the difference between drying in air or Ar is reflected by the smaller C 1s/Nd 3d ratio in air than in Ar, i.e., 0.05 instead of 0.10 (Table 4). This difference is also confirmed by the larger value of the C 1s (ca. 286 eV)/Nd 3d ratio in Ar than in air. Therefore, also in the case of DMF, the 1:1 ratio of the C 1s signals of the Ar-dried sample becomes almost 1:2 after drying in air. The C 1s (ca. 289 eV)/Nd 3d ratios are very similar both in air and in Ar, indicating that the chemisorbed products are not considerably affected by drying in different environments.

#### 2.8.2.3. Samples prepared from pyridine

The XPS analyses of the air- and Ar-dried pyridine samples showed the presence of the following elements: C, N, O, Pd, and Nd. The corresponding XPS signals were: C 1s, N 1s, O 1s, Pd 3d, and Nd 3d. Fig. 12 shows the spectra of Nd 3d (a) and C 1s (b) of the sample dried in air, and after 10 min sputtering with Ar⁺ ions. Offsets were applied for clarity.

According to literature [21], the C 1s components of DMF have binding energies at 285.8 eV, and 288.8 eV with an intensity ratio of 2:1.
Fig. 11. XPS spectra of samples obtained after plating on Ti3 substrates using MC plating solution in DMF: (a) Nd 3d (air-dried), (b) C 1s (air-dried), and (c) C 1s (Ar-dried) before sputtering (indicated as 0 min sputtering in the figure), and after 10 min sputtering with Ar$^+$ ions. Offsets were applied for clarity.

Fig. 12. XPS spectra of samples obtained after plating on Ti3 substrates using MC plating solution in pyridine: (a) Nd 3d (air-dried), (b) C 1s (air-dried), and (c) C 1s (Ar-dried) before sputtering (indicated as 0 min sputtering in the figure), and after 10 min sputtering with Ar$^+$ ions. Offsets were applied for clarity.
Fig. 12a). This could indicate the presence of different Nd\(^{3+}\) ions. Therefore, the produced sample is characterized by a thicker solvent layer covering the deposit compared to IP\(^+\)IB and DMF. The measured binding energy of the Nd \(3d_{5/2}\) peak (i.e., 983.4 eV, Fig. 12a) confirms that Nd is deposited as Nd\(^{3+}\) as previously found in the case of IP\(^+\)IB and DMF (possibly as carboxylate, oxide, or hydroxide). The C 1s spectra reported in Figs. 12b and c show the presence of two components, which are visible only after multicomponent fitting analysis: the components of physisorbed pyridine at ca. 286 eV, and the components of chemisorbed species at ca. 289 eV (i.e., air: 289.3 eV, Ar: 289.4 eV). These higher binding energy components can be attributed to possible COOM\(–\) metal carboxylate \(-\) groups formed during the plating process. This indicates that the solvent undergoes electrolysis during the MP process and becomes oxidized. Nonetheless, the very small intensities of the chemisorbed components suggest that pyridine undergoes electrolysis only to a smaller degree than IP\(^+\)IB or DMF.

Slight differences before drying in air or Ar are shown by the smaller C 1s/Nd 3d ratio in air (i.e., 0.9 instead of 1.1), and by the larger value of the C 1s (ca. 286 eV)/Nd 3d ratio in Ar than in air. These differences are smaller than in the case of IP\(^+\)IB and DMF. The C 1s (ca. 289 eV)/Nd 3d ratios are very similar both in air and in the Ar environment.

It has to be noticed that all the values of the ratios reported in Table 4 for the deposits prepared in pyridine are quite different from the corresponding values determined for IP\(^+\)IB or DMF deposits, irrespective of the methods of sample drying. This follows from the large intensities of the C 1s signals in pyridine, as evidenced by the C 1s spectra reported in Figs. 12b and c, where even after 10 min sputtering the intensities of the physisorbed components are only slightly decreased.

The comparisons between the C 1s spectra in Figs. 10b, c, 11b, c and 12b, c show no chemical shifts before and after sputtering. Small shifts of ca. \(-0.5\) eV after sputtering were observed for the Nd 3d spectra of the IP\(^+\)IB and DMF samples (see Fig. 10a for IP\(^+\)IB, and Fig. 11a for DMF). Shifts of ca. \(+0.5\) eV after sputtering were observed for the Nd 3d spectra of the pyridine samples (see Fig. 12a). This could indicate the presence of different Nd\(^{3+}\) species in the topmost surface layers of the samples produced from the different solvents.

From the analysis of all the obtained XPS results, the main difference which appears to occur between samples dried in air and in a controlled inert environment, such as Ar in a glove-box, is related to the higher intensities of the peaks of the physisorbed solvent components when the drying is performed in Ar. Besides this, a slightly larger amount of O adsorbed on the surfaces of all the samples dried in air can be noticed (see O 1s/C 1s ratios in Table 4). Nonetheless, no strong indication for the presence of surface chemical reaction products promoted by the drying in air instead of Ar was found.

### 2.9. Molecular plating from DMF solution

#### 2.9.1. Cell potential trends

Fig. 13 shows trends of cell potential vs. time recorded during preparation of targets, using 0.22 mM [Nd(NO\(_3\)]\(_3\)-6H\(_2\)O] solution (MC), Ti3 as deposition foil, and DMF (a) or IP\(^+\)IB (b) as plating solvent. The applied constant current was 6 mA for 3 h (current density 0.7 mA/cm\(^2\)). IP\(^+\)IB cell potential curves were already discussed in detail in [14].

From the comparison between the two cell potential curves, it is evident that after an increase at the beginning of the plating process, both curves decrease to a minimum value, then they increase again to reach a maximum. After this maximum, both cell potential curves tend to decrease slightly with time. The initial DMF cell potential increase is much bigger (ca. 100 V) and lasts longer than that of IP\(^+\)IB, and forms a wide maximum after ca. 30 min. In the case of IP\(^+\)IB the increase is only a few tens of volts and it lasts only for a few minutes at the very beginning of the deposition. The cell potential drop after the initial increase is much bigger for IP\(^+\)IB (ca. 350 V) than for DMF (ca. 50 V) and the minimum is reached earlier (45 min vs. ca. 60 min). The minimum area is wider for IP\(^+\)IB than for DMF. The second maximum is wide in both cases but it is achieved somewhat earlier in DMF (ca. 90 min) than in IP\(^+\)IB (ca. 110 min). The cell potentials in DMF are much smaller than the values recorded in IP\(^+\)IB.

#### 2.9.2. Kinetics derived from \(\gamma\)-ray spectroscopy

The kinetics of the MP with DMF was studied by determining the Nd deposition yield values with \(\gamma\)-ray spectroscopy. A high-purity germanium detector (GEM series HPGe Detector Model No. GEM 23158 P-Plus, ORTEC Company), was used after MPs of MC Nd solutions containing radioactive \(^{147}\)Nd tracer. To obtain quantita-

tive data, reference sources with known amounts of the tracer were prepared. They consisted of filter papers with the same geometry as the targets, soaked with the tracer-containing solution. Twelve Ti3 targets were produced by electrolysis of aliquots of 100 \(\mu\)L of this solution at different deposition times: the first target after 15 min and the subsequent ones after increments of 15 min up to 180 min. The \(\gamma\)-measurements of the twelve targets gave the deposition yields reported in Fig. 14, where the dashed curve fitting the open circles shows the Nd\(^{3+}\) depletion trend obtained by subtracting the deposition yield from 100%.

Nd\(^{3+}\) depletion shows an exponential decrease which can be fitted by a first-order rate law \(y = y_0 \exp(-kt)\), i.e., \([\text{Nd}] = [\text{Nd}]_0 e^{-kt}\), where [Nd] and [Nd] are the initial and instantaneous concentrations of Nd\(^{3+}\) in solution, respectively, and \(k\) is the rate constant of the process.

For comparison Fig. 14 shows also the Nd\(^{2+}\) depletion curve obtained during MP in IP\(^+\)IB (open triangles) [14]. Both MPs follow a first-order rate law, but it is evident that the depletion kinetics is faster (i.e., \(k\) is larger) in DMF than in IP\(^+\)IB. In
particular, it is important to notice that, as in the case of IP+IB [14], the MP in DMF reaches an almost complete depletion of the Nd ions at a time corresponding to the recorded minimum (i.e., ca. 60 min) of the cell potential curve.

2.9.3. Conductivity measurements

The conductivity was measured using an inoLab Cond 720 conductometer connected to a TetraCon 325 standard conductivity cell. These measurements were performed only in a stirred DMF solution containing 0.22 mM [Nd(NO₃)₃·6H₂O] and 0.3 mM HNO₃. The conductivity (\(\sigma\)) was measured to be 46.5±0.1 \(\mu\)S/cm. This value was used to evaluate the solution resistance, \(R\), in the cell. The expression used for this evaluation is \(R=\rho S/l\) [24], where \(\rho\) is the resistivity of the system (i.e., the inverse of the conductivity, \(\rho=21.50±0.05\) kΩ/cm), \(l\) is the distance between the electrodes (2.80±0.01 cm), and \(S\) is the surface area of the electrodes (9.0 ±0.1 cm²). The obtained cell resistance is 6.691±0.011 kΩ. Using Ohm’s first law, \(V=RI\), where \(R\) is the cell resistance and \(I\) is the applied current (\(I=6.00±0.01\) mA), the contribution of the ohmic drop to the overall cell potential was estimated to 40.15±0.09 V. This value is in perfect agreement with the initial experimental value in Fig. 13a. The cell potential during the MP process is, however, at least twice as high as the initial value. The voltage increase during the first 15 min of the electroplating is probably due to an increase of the overall resistance of the cell. Indeed, during MP the concentration of the electrolyte decreases as Nd\(^{3+}\) is deposited on the cathode, resulting in a decrease of conductivity. Also the formation and growth of the deposit, which is principally composed of Nd salts and complexes, presents an additional resistance. It appears reasonable that the cell potential observed during MPs is dominated by the ohmic drop.

2.9.4. Temperature measurements

Temperature measurements of the DMF plating solutions were performed using a TFK 150 thermometer connected to an inoLab pH 720 pH meter. The temperature was measured immediately before and after 3 h constant current density MP experiments at 0.17, 0.3 and 1.3 mA/cm². In all cases the MP experiments started after the solutions were cooled down to 16 °C. The measurements showed temperature increases of 0.4±0.1 °C, 0.7±0.1 °C, and 2.0 ±0.1 °C, respectively. These results indicate that despite constant cooling during the experiments, a higher current density leads to a more pronounced temperature increase inside the system.

3. Discussion

Among various experimental parameters that may affect constant current density MP, two were found to be of crucial importance for the preparation of smooth and crack-free layers to be used as nuclear targets. These are the roughness of the deposition substrate and the solvent used for the platings. The different RMS values of the DMF MC targets grown on Ti3 and Ti5 substrates confirm that even small differences in surface roughness between the deposition substrates can influence the RMS of the growing layers [14]. Indeed, if the deposition substrate is rough (i.e., defective), the surface mobility of the deposited atoms will be reduced by trapping in the defects, and the produced layer will be of very low quality. Therefore, to produce smooth targets, deposition substrates as smooth as possible are required.

As regards the solvent, its importance does not only derive from being the medium in which the process is performed and hence the major factor influencing the electrochemistry of the MP, but also from the fact that the produced layers are mainly composed of physisorbed as well as chemisorbed solvent molecules. Hence, the role of the solvent in MP is twofold: it influences the electrochemistry of the process, but also the properties of the layer to be produced. Reasonable physical properties to be taken into account when choosing a solvent for MP are boiling point, viscosity, conductivity, and relative permittivity [16]. The boiling point determines the volatility of the solvent. The viscosity influences the rate of mass transfer in the solvent and is important in mass-transport controlled electrolytic processes like MP [14]. In turn, the rate of mass transfer influences the conductivity of the electrolyte solutions, which is fundamental for constant current density electrolysis experiments. Finally, the relative permittivity of a solvent has a decisive influence on the electrostatic solute-solute and solute-solvent interactions as well as on the dissolution and dissociation of electrolytes. In this study, pyridine and DMF were preferred over the more commonly used IP+IB mixture [9,10], after a careful analysis of these physical and chemical properties. All the samples produced using IP+IB, pyridine, and DMF as plating solvents and Ti3 foils as deposition substrates showed the presence of surface cracks. The liquid AFM analysis of the IP+IB sample suggested that the drying of the layer is at the basis of the surface cracking. To rule out the possibility that cracking may be connected to eventual layer-destructive chemical surface reactions occurring during the drying in air of the samples, XPS investigations were performed. The results of these investigations give no indications for the occurrence of such reactions, as air- and Ar-dried samples presented the same XPS signals. The absence of reactivity made then possible the AFM analysis in air of all these samples. These analyses confirmed the hypothesis of the drying-related nature of cracking. Indeed, the layers presenting surface cracks after drying in air also underwent cracking in Ar environment. More important than the drying environment is the boiling point of the solvent. Samples prepared in solvents with low boiling point (i.e., IP+IB and pyridine) showed the most severe surface cracking and the highest roughness. In contrast, films produced in DMF on Ti3 foils showed much smaller surface cracks and roughness. The reason why solvents with lower boiling point produce layers with severe surface cracking is very likely connected to the drying process. During drying in air, removal of solvent by evaporation implies solvent diffusion to the surface, conversion at the interface from the fluid phase to the vapour phase, and finally diffusion through the boundary layer into the distant air surrounding [25]. Because solvent evaporation diminishes the volume of the stress-free state of the coating, the current state shrinks to the maximum possible extent. Ahead of the solidification front, i.e., the front indicating the transition from liquid to solid, the shrinkage produces flow in the still liquid
coating; behind, it produces in-plane stress in the solid coating because its in-plane shrinkage is frustrated by adhesion to the substrate [26,27]. The difference between the current state and the stress-free state of the coating is elastic strain to which stress is proportional via the elastic modulus (i.e., $\lambda = \text{stress/strain}$, where $\lambda$ is the elastic modulus). Experimental and theoretical investigations [27,28] have shown that high elastic modulus and high yield-stress raise the level of stress and in-plane stress gradients, making the coatings susceptible to cracking and delamination. It is then reasonable to argue that because the high volatility IP-IB and pyridine solvents evaporate faster than the less volatile DMF (as we experimentally observed), the big volume decrease of the stress-free state of the coatings produced with such solvents will bring to a considerable difference between the current state and the stress-free one. The consequent large elastic modulus will then make these layers more likely to crack. In the “slow evaporation” regime achieved with DMF, instead, the difference between the states during the drying and the consequent elastic modulus will unlikely be as high as to create cracks. Cracking will mainly occur where a larger fragility of the layer is dictated by higher surface stress, i.e., where the surface defects have grown from the deposition foil. Once such defects are eliminated by using smoother deposition substrates, like Ti5, no cracks are observed in the surfaces of the produced DMF layers anymore. In particular, the layer growing on the Ti5 substrate during a MC plating experiment performed in DMF is assigned to follow a Frank-van der Merwe mechanism (layer-by-layer growth mechanism). In this growth mode a new layer is nucleated only after completion of the layer below and can spread over macroscopic distances creating very homogeneous surfaces, as demonstrated also by the RI studies.

The surface roughness and the morphology of the layers produced using DMF as plating solvent and Ti5 as deposition substrate can be considerably varied by changing some basic variables of the MP, like the concentration of the electrolyte to be plated and the applied current density. In the first case, the higher concentration of the electrolyte increases the flux of Nd$^{3+}$ ions to the cathode, and hence the deposition rate onto the substrate. Because a high deposition rate causes the system to grow disorderly, the final surface roughness of the produced layer will be large [14]. In the case of the layers obtained applying different constant current densities, this simple interpretation is no longer sufficient to explain the changes observed in both roughness and morphology of the surfaces. In fact, if on the one hand, as expected, the current density of 1.3 mA/cm$^2$ increases the surface roughness of the layer produced with 0.3 mA/cm$^2$, on the other hand, the decrease to 0.17 mA/cm$^2$ increases the roughness even more. A possible explanation for these results comes from the fact that good epitaxial growths require not only low deposition rates, but also high surface mobility of the deposited species. In the case of the deposit obtained with the highest current density, the big outgrowths not covering the entire analyzed area may in fact suggest that the deposited species had a high surface mobility, which allowed them to condensate. In contrast, as regards the lowest current density sample, the rather small outgrowths spread over the entire surface area may indicate low surface mobility. The different mobilities could be related to the different temperatures of the substrates, which influence the ability of the deposited species to pass the activation barrier of diffusion. As the MP performed in DMF is controlled by Ohm’s first law, $V=RI$, it is easy to understand that the range of cell potential values must be smaller in DMF, because this solvent, having a larger conductivity, implies a smaller resistance. It is important to notice that in both solvents the deposition yield at the minimum of the cell potential curves is already high, with the depletion kinetics being faster in DMF than in IP-IB.

Finally, we like to stress that in none of the MPs performed from any of the used solvents metallic Nd was deposited. Rather, Nd is deposited on the cathodic surfaces as Nd$^{3+}$ derivatives, and this is reasonable on the basis of the highly negative value of the Nd$^{3+}$/Nd standard redox potential ($E = -2.323$ V [29]), typical of lanthanide cations.

4. Conclusion

To optimize the production of smooth and crack-free layers to be used as nuclear targets, several constant current density MPs were performed using Nd as a model element. The variables chosen for these investigations were: plating solvent, electrolyte concentration, applied current density, and surface roughness of the deposition substrate. Different environments were also used to dry the produced samples. The obtained deposits were analyzed using γ-ray spectroscopy, XPS, RI, AFM, and SEM. The results obtained from the characterization were used to identify the key parameters controlling the molecular plating of smooth and crack-free layers. The roughness of the deposition substrate and the solvent used for the platings proved to be two core factors playing a decisive role for the production of high quality layers. Surface cracking in fact occurred by means of drying only when volatile solvents and deposition substrates that were not smooth enough, were used. Different drying environments like air and Ar did not show to have any effect on the deposited layers.

Molecular platings from DMF proved to be superior over the widely performed IP-IB platings [9,10]. Indeed, despite being also governed by Ohm’s first law and by similar depletion kinetics, DMF MPs performed on Ti5 substrates produced targets showing complete elimination of the surface cracks as well as high smoothness and homogeneity. Moreover, no Pd impurities from the oxidation of the Pd anode were found on the cathode deposit when MPs were performed in DMF. The absence of such impurities may be of importance in nuclear science investigations as they could give rise to unwanted background in accelerator-based experiments.

Finally, the surface roughness and morphology of the layers produced using DMF and Ti5 substrates proved to be very sensitive to changes in, e.g., electrolyte concentration and current density. The present work demonstrates that molecular plating can be applied to produce layers that are as smooth and crack-free as those produced by PAD, and suitable as nuclear targets. An important
advantage of MP over PAD is that it is a faster deposition process, as it needs just one plating cycle to reach almost 100% deposition yield. With PAD, in contrast, coating efficiencies for one single application of ca. 33% are reported [11]; hence several deposition cycles need to be performed to obtain quantitative yields.

The results obtained in this study may also be applied for the production of smooth and crack-free lanthanide targets other than Nd, and also for the preparation of heavier actinide targets with an oxidation state of 3+ (e.g., americium and curium), for which Nd is a good model element.

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References