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Fluorine-free organic electrolytes for the stable electrodeposition of neodymium metal†

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Electrowinning is regarded as a clean process to recover neodymium metal from secondary sources such as spent Nd-Fe-B permanent magnets, but the current methods are severely limited by a high energy consumption (molten salts), or by the high costs and environmental impact of the electrolyte components (ionic liquids). Therefore, there is a demand for more sustainable electrowinning methods for the recovery of neodymium metal. Inspired by our own previous work and the work of others, we developed new fluorine-free organic electrolytes that enable the electrodeposition of neodymium metal at room temperature. The electrolytes consist of solvated neodymium borohydride, Nd(BH₄)₃, dissolved in the ether solvents tetrahydrofuran (THF), 2-methyltetrahydrofuran (2-MeTHF), 1,2-dimethoxyethane (DME) and diethylene glycol dimethyl ether (diglyme, G2), and these complexes can be prepared entirely from non-fluorinated precursors such as neodymium(III) chloride (NdCl₃) and sodium borohydride (NaBH₄). In contrast to our previous bis(trifluoromethylsulfonyl)imide-containing electrolytes, electrodeposition of neodymium proceeds over time without significant loss of current density, indicating a higher stability against unwanted side-reactions that lead to passivation of the deposit on the electrode. Characterization of the deposits by scanning electron microscopy (SEM), energydispersive X-ray fluorescence (EDX), and X-ray photoelectron spectroscopy (XPS) unambiguously indicated the presence of neodymium metal.

Introduction

Neodymium is economically the most important rare-earth element (REE) because it is one of the main components in Nd-Fe-B permanent magnets, which are used extensively in household electric appliances, hard disk drives, electric motors and generators. In order to meet the goals of the green energy

transition, the demand of Nd-Fe-B magnets for (hybrid) electric vehicles and wind turbines is expected to increase sharply in the near future.² Therefore, the development of sustainable methods for the recovery of neodymium metal from secondary sources such as end-of-life products is urgently needed.3,4 Electrowinning is widely regarded as a reliable, low-cost and clean method for the recovery of metals because an electric current is used to reduce the desired metal cations from solution, as opposed to a chemical reducing agent which generates additional waste.⁵ This process is also used for REEs, including neodymium, but additional challenges regarding the choice of electrolyte must be addressed. Due to the high chemical reactivity and very negative standard reduction potentials of these elements (ranging between -2.3 to -2.4 V vs. SHE), aqueous electrolytes are not suitable due to the hydrogen evolution reaction, and research has been mainly focussed on nonaqueous electrolytes such as molten salts and ionic liquids. In the industry, molten salt electrolysis is currently the most attractive recovery method because the electrolyte salts (e.g. LiCl-KCl eutectic) are inexpensive and high deposition current densities can be achieved. 6-8 However, high temperatures (>500 °C) are required to operate these electrowinning baths, and the molten salts are corrosive to the infrastructure. This imposes a high energy requirement and important

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[†] Electronic supplementary information (ESI) available: Powder XRD diffractograms of $Na[Nd(BH_4)_4][(DME)_4$ and $Nd(BH_4)_3(THF)_3$. CVs of $Li(Tf_2N) + 4$ molar equivalents of LiBH4, and NaTf2N + saturated NaBH4 in THF. CVs of Na[Nd(BH₄)₄](DME)₄ in THF with varying cathodic vertex potential. CVs of Na[Nd(BH₄)₄](DME)₄ in MeTHF, DME, and G2. CVs of Nd(BH₄)₃(THF)₃ + LiBH₄ in THF, and Nd(BH₄)₃(THF)₃ + TBABH₄ in THF. CV of Nd(BH₄)₃(THF)₃ in THF. Chronoamperograms for a potentiostatic deposition in Nd(BH₄)₃(THF)₃ + one molar equivalent of LiBH4 or TBABH4 in THF. SEM micrograph of a neodymium deposit at 500× magnification + corresponding EDX analysis in several spots. XPS survey scan in the range 0-1100 eV of a neodymium deposit obtained from potentiostatic deposition in 0.1 mol L⁻¹ Na[Nd(BH₄)₄] in THF electrolyte. See DOI: https://doi.org/10.1039/d3cp01262j

considerations on the maintenance and safety of the process. Therefore, there is a high demand for alternative electrolytes that are operable at moderate temperatures (≤150 °C) while still allowing high deposition current densities. Currently, ionic liquids (ILs), which are low-melting organic molten salts (mp \leq 100 °C) and are non-volatile, are being investigated to fulfill this role. 9-11 For electrowinning of neodymium metal, research has been focused mainly on ILs with phosphonium and ammonium cations and the bis(trifluoromethylsulfonyl)imide (bistriflimide, TFSI-, Tf2N-) anion because they have many desirable properties for electrodeposition, i.e. a very negative cathodic decomposition limit, good thermal stability, hydrophobicity, a relatively low viscosity, and a high conductivity. 12-15 While these reports demonstrate the proof-ofconcept for the achievability of low-temperature electrowinning of neodymium and other REEs, upscaling of these processes is not sustainable due to the fluorinated nature of the IL anions. Numerous studies have shown that strongly fluorinated organic compounds are very poorly (bio)degradable, and have a high degree of (eco)toxicity. 16-18 Furthermore, the fluorinated bistriflimide anion is expensive because of its multi-step synthesis procedure and its niche application fields. 19 Hence, its use should be avoided or at least minimized, especially for largescale applications such as electrowinning.

Recently, research on electrodeposition of REEs from ILs has shifted to more cost-effective and non-fluorinated ILs, in particular N-butyl-N-methylpyrrolidinium dicyanamide ([BMPyrr]-[DCA]).²⁰⁻²² In an earlier report, it was found that electrolytes consisting of Ln(Tf₂N)₃ (Ln = La, Nd, Sm) in [BMPyrr][DCA] were not suitable for electrodeposition because the cathodic decomposition of the IL occurred simultaneously with the electrodeposition of metal.20 However, more recently it was discovered that using 0.5 mol kg⁻¹ of the hydrated salt Nd(NO₃)₃·6H₂O as the source of Nd³⁺ cations in this IL, resulted in a significant positive shift of the deposition onset potential ($-1.75 \text{ V} \text{ vs. } \text{Fc}^+\text{/Fc}$) and an increase in the deposition current density (-40 mA cm⁻²).²¹ The obtained deposits consisted of a mixture of metallic neodymium and Nd₂O₃. The presence of water in the coordination sphere around Nd3+ was proven to be essential to achieve neodymium electrodeposition.21,22 However, while water might improve the thermodynamics and kinetics of the deposition process and represent a "more realistic" (i.e. not rigorously dried) environment, these wet electrolytes also provide an inherently unstable environment for the deposits, leading to contamination of the metallic REEs with oxides and hydroxides, and a decreased current efficiency.²³

Organic electrolytes are composed of salts which are dissolved in an organic solvent and they could be a less expensive and sustainable alternative to ILs, provided that they are sufficiently electrochemically stable to support REE electrodeposition. This category of electrolytes has been much less explored than ILs. "Neutral-ligand" ionic liquids (NLILs), composed of RE(Tf₂N)₃ dissolved in neutral ligands (L) such as trimethyl phosphate or dihexyl octanamide, resulting in a cationic complex [RE(L)₃][Tf₂N]₃, were briefly investigated by Bagri et al. and Krishna et al. 24,25 However, the authors did not characterize the obtained deposits in detail. Similarly, Zhang and co-workers have recently proposed two different systems, NdCl₃-AlCl₃ and Nd(SO₃CF₃)₃-LiNO₃ dissolved in the green aprotic organic solvent 1,3-dimethyl-2-imidazolidinone, for the electrodeposition of neodymium at ambient temperature. 26,27 Evidence for the presence of metallic neodymium in the deposits was provided by X-ray photoelectron spectroscopy (XPS) and X-ray diffraction (XRD) analysis.

In our previous work, we have reported the successful electrodeposition of neodymium- and dysprosium-containing layers from organic electrolytes consisting of Ln(Tf₂N)₃ (Ln = Nd, Dy) and a commercially available borohydride salt, dissolved in 1,2-dimethoxyethane (DME, G1).28 Due to the presence of strongly electron-withdrawing CF₃SO₂-functionalities, the bistriflimide anion is prone to reductive side-reactions, leading to the formation of passivating surface films. 29-31 This was evidenced by a fast decrease in deposition current density during chronoamperometry (CA) experiments, and by the presence of fluorineand sulfur- containing species in the EDX spectra of the deposits. Therefore, in order to achieve electrodeposition of pure metallic REE at a constant and high current density, it is necessary to avoid the use of this anion completely.

In this Paper, we describe a new strategy to prepare organic electrolytes for neodymium electrodeposition at ambient temperatures, without relying on fluorinated salts. Commercially available neodymium(III) chloride and sodium, lithium and tetrabutylammonium borohydride are used as the precursors. Alternatively, neodymium(III) methanesulfonate, which is conveniently synthesized from Nd2O3 and methanesulfonic acid can also be used as a starting salt. Moreover, because the electrolytes do not contain any reactive impurities, electrodeposition proceeds at a constant current density, as no passivation occurs.

Experimental

Products

Sodium borohydride (NaBH₄, 99%, powder), lithium borohydride (LiBH₄, 95%), tetrabutylammonium borohydride (TBABH₄, 98%), 1,2-dimethoxyethane (ethylene glycol dimethyl ether, DME, 99+%, extra dry over molecular sieves), bis(2-methoxyethyl) ether (diethylene glycol dimethyl ether, diglyme, 99+%, extra dry over molecular sieves), tetrahydrofuran (THF, 99.5%, extra dry over molecular sieves), 2-methyltetrahydrofuran (2-MeTHF, 99+%, extra dry over molecular sieves), ferrocene (Fc, 98%), and neodymium(III) chloride (NdCl₃, 99.9% trace metal base, anhydrous) were purchased from Acros Organics (Geel, Belgium). Ferrocenium hexafluorophosphate (FcPF6, 97%) was purchased from Sigma Aldrich (Diegem, Belgium). Hydrogen bis(trifluoromethylsulfonyl)imide (HTf₂N, 98%, 80 wt% aqueous solution), lithium bis(trifluoromethylsulfonyl)imide (LiTf₂N, 99%), and 1-butyl-1-methylpyrrolidinium bis(trifluoromethylsulfonyl)imide ([BMP][Tf₂N], 99%) were purchased from IoLiTec (Heilbronn, Germany). Methanesulfonic acid (HOMs, ≥99.5%, for synthesis) was purchased from Carl

Roth (Karlsruhe, Germany). Neodymium(III) oxide (Nd₂O₃, 99.99%) was purchased from Rare Earth Products ltd. (Beverly, USA). Neodymium(III) bis(trifluoromethylsulfonyl)imide (Nd(Tf₂N)₃),³² neodymium(III) methanesulfonate (Nd(OMs)₃),³³ sodium bis-(trifluoromethylsulfonyl)imide (NaTf2N)34 were synthesized according to previously reported procedures.

All chemicals were used as received without any further purification, with the exception of the bis(trifluoromethylsulfonyl)imide and methanesulfonate salts, BMPTf₂N, and neodymium(III) borohydride complexes which were dried prior to use on a Schlenk line for 12 hours at 220 °C, 120 °C, and ambient temperature, respectively. All the products used in the electrolytes were stored and manipulated inside an argon-filled glovebox with water and oxygen concentrations below 1 ppm.

Characterization

Powder XRD diffractograms were recorded on a Bruker D2 phaser in a 2θ range of 5° – 85° , using a Cu K α X-ray source. UV-Vis absorption spectra were recorded on a Varian Cary 6000i UV-VIS-NIR spectrophotometer at a scan rate of 10 nm s⁻¹ with 1 nm intervals. The instrument was used in double-beam mode. Samples were measured in quartz glass cuvettes with a path length of 5 mm. Blanks consisted of the same solvent used in the samples, and were measured in a matched pair of cuvettes. Scanning electron microscopy (SEM) was performed on an XL30 FEG scanning electron microscope. Electrodepositions were made on platinum-coated silicon wafers and were thoroughly rinsed with anhydrous THF, and were allowed to dry at ambient glovebox conditions before they were attached to the SEM sample holder with conductive carbon tape. Energy dispersive X-ray fluorescence (EDX) analysis was performed in the same setup, using an octane elite super silicon drift detector (Ametek EDAX), and using TEAM software. Scans were recorded over a total time of 200 s to achieve a sufficient signal-to-noise ratio. The acceleration voltage was 20 kV or 30 kV for both imaging and EDX. During the sample preparation, exposure of the deposits to air was unavoidable, causing oxidation. Electrodepositions for XPS analysis were made and treated identically to the SEM samples, but were transported in air-tight vials to another glovebox, where they were transferred to a vessel which was directly attached to the ultra-high-vacuum system to avoid exposure to air. The XPS measurement and etching was conducted by a PHI5000 Versaprobe-II, UlVAC-PHI with an Al target working at 1486.8 kV. The surface layer of the deposits was etched with Ar⁺ beams. The etching area was 2 mm imes 2 mm and the etching time was 30 s for a total of 10 times. The XPS measurement was conducted during the interval between two etchings. The XPS spectra were calibrated by shifting the low binding energy component of the C 1s peak to 284.8 eV. The peak fitting and quantitative analysis of the spectra was performed with CasaXPS software. The spectra were fitted with Doniach-Sunjic functions convoluted with a Gaussian/Lorentzian distribution and a Shirley background.

Electrochemical experiments

All the electrochemical measurements were performed inside an argon-filled glovebox with water and oxygen concentrations

below 1 ppm, at ambient temperature (28 °C). The electrolyte solutions were prepared by mixing the appropriate amounts of products and solvents in 20 mL glass vials and stirring for 5 minutes at ambient temperature. In many cases, precipitation was observed in the electrolyte solutions, which was allowed to settle to the bottom of the vessel before proceeding with the measurement. All the electrochemical measurements were performed in three-electrode setup, using an Autolab PGSTAT302N potentiostat and Nova 2.1 software. The working electrodes for cyclic voltammetry (CV) were pieces of silicon wafer, coated with 500 nm of silica, 10 nm of titanium and 100 nm of platinum (Imec, Belgium) (surface area of 0.3 cm²), the counter electrode a larger piece of platinum-coated silicon wafer (surface area of approx. 1.0 cm²) and the scan rate was 10 mV s⁻¹, unless stated otherwise. The reference electrode consisted of a platinum wire, submerged in a mixture of ferrocene (0.005 mol L⁻¹) and ferrocenium hexafluorophosphate $(0.005 \text{ mol L}^{-1})$ dissolved in the ionic liquid 1-butyl-1methylpyrrolidinium bis(trifluoromethylsulfonyl)imide, contained inside a fritted glass tube (referred to as Fc⁺/Fc). The CVs were started at open circuit potential (OCP) and are uncorrected for Ohmic drop. Electrodepositions were performed at constant potential, using pieces of platinum-coated silicon wafers (surface area of approx. 0.3 cm²) as substrates.

Synthesis of salts/complexes

Synthesis of neodymium(III) methanesulfonate, Nd(OMs)3. Nd₂O₃ (7.06 g, 1 equiv., 20.98 mmol) was added to a stirring solution of methanesulfonic acid (9.49 g, 4.7 equiv., 98.75 mmol) in demineralized water (30 mL) and the mixture was stirred at 90 °C for 30 min. Excess Nd₂O₃ was removed by filtration to give a clear pale pink solution. Water was removed on a rotary evaporator to give a highly viscous purple liquid which slowly crystallized to Nd(OMs)3.2H2O. The product was heated on a Schlenk line at 250 °C for 48 h to drive off any crystal water to give anhydrous neodymium(III) methanesulfonate, Nd(OMs)3 (12.72 g, 29.63 mmol, 90% yield) as a pink/ violet crystalline solid. IR (ATR, $\nu_{\text{max}}/\text{cm}^{-1}$): 3029 (ν C-H), 2940 $(\nu C-H)$, 1429 ($\delta C-H$), 1412, 1331 ($\nu S=O$), 1301, 1199 ($\nu S=O$), 1150, 1123, 1055, 974, 797, 782, 553, 534, 517, 506, 409.

Synthesis of Na[Nd(BH₄)₄](DME)₄ from NdCl₃. A suspension of NdCl₃ (2.000 g, 1 equiv., 7.981 mmol) and NaBH₄ (1.210 g, 4.0 equiv., 31.985 mmol) in anhydrous DME (50 mL) was stirred for 24 h inside an oven-dried 100 mL round bottom flask under argon. The color of the solution changed from colorless to violet and a fine white precipitate (NaCl) was observed. The reaction mixture was filtered under argon through a Schlenk filter to give a clear violet solution. The solution was placed inside a NaCl/ice bath ($T = \text{approx.} -15 \,^{\circ}\text{C}$) and large violet crystals precipitated from solution. The supernatant was carefully removed with a syringe and the crystals were briefly dried on the Schlenk line at room temperature to give $Na[Nd(BH_4)_4](DME)_4$ (2.23 g, 3.798 mmol, 48% yield) as a violet crystalline solid. IR (ATR, $\nu_{\text{max}}/\text{cm}^{-1}$): 2937 (ν C-H), 2844 $(\nu C-H)$, 2828 $(\nu C-H)$, 2418 $(\nu B-H)$, 2298 $(\nu B-H)$, 2223 $(\nu B-H)$, 1452 (δ C-H), 1401, 1368, 1295, 1281, 1255, 1243, 1206, 1186,

1152 (ν C-O), 1123 (ν C-O), 1082 (ν C-O), 1035, 1013, 978, 855, 818, 707, 557, 469.

Synthesis of Na[Nd(BH₄)₄](DME)₄ from Nd(OMs)₃. A suspension of Nd(OMs)₃ (1.508 g, 1 equiv., 3.511 mmol) and NaBH₄ (0.676 g, 4.2 equiv., 17.869 mmol) in anhydrous DME (40 mL) was stirred for 24 h inside an oven-dried capped glass vial under argon. The color of the solution changed from colorless to violet and a fine white precipitate (NaOMs) was observed. The reaction mixture was filtered under argon through a Schlenk filter to give a clear violet solution. The solution was placed inside a NaCl/ice bath ($T = \text{approx.} -15 \,^{\circ}\text{C}$) and large violet crystals precipitated from solution. The supernatant was carefully removed with a syringe and the crystals were briefly dried on the Schlenk line at room temperature to give Na[Nd(BH₄)₄](DME)₄ (1.07 g, 1.823 mmol, 42% yield) as a violet crystalline solid. IR (ATR, $\nu_{\text{max}}/\text{cm}^{-1}$): 2937 (ν C–H), 2844 $(\nu C-H)$, 2828 $(\nu C-H)$, 2418 $(\nu B-H)$, 2298 $(\nu B-H)$, 2223 $(\nu B-H)$, 1452 (δ C-H), 1401, 1368, 1295, 1281, 1255, 1243, 1206, 1186, 1152 (ν C-O), 1123 (ν C-O), 1082 (ν C-O), 1035, 1013, 978, 855, 818, 707, 557, 469.

Synthesis of $Nd(BH_4)_3(THF)_3$. In an oven-dried 100 mL twoneck round bottom flask, charged with a stirring bar, was added NdCl₃ (2.51 g, 1 equiv., 10.02 mmol), NaBH₄ (1.26 g, 3.3 equiv., 33.31 mmol), and anhydrous THF (35 mL). The flask was attached to a reflux condenser, the setup was flushed with argon and sealed airtight with an argon balloon and septum, and the mixture was stirred at 60 $^{\circ}\text{C}$ for 48 hours. The color of the supernatant solution was changed from colorless to deep violet, and a colorless solid was formed (NaCl). The solid was removed by vacuum filtration under argon to give a clear deep violet solution. The solution was subsequently concentrated to dryness on a rotary evaporator at 40 °C to give a pale violet solid. The product was dried for 30 minutes on the Schlenk line at room temperature to give Nd(BH₄)₃(THF)₃ (3.23 g, 7.97 mmol, 80% yield) as a pale violet solid. IR (ATR, $\nu_{\rm max}/{\rm cm}^{-1}$): 2984 $(\nu C-H)$, 2890 $(\nu C-H)$, 2437 $(\nu B-H)$, 2324 $(\nu B-H)$, 2217 $(\nu B-H)$, 2163 (ν B-H), 1457 (δ C-H), 1366, 1344, 1316, 1295, 1249, 1150 $(\nu C-O)$, 1091 $(\nu C-O)$, 1037, 1012, 959, 918, 854, 840, 669, 579, 532, 508, 493.

Results and discussion

Synthesis and characterization of complexes

In our previous work, an ether-soluble neodymium(III) salt, Nd(Tf₂N)₃, was combined with a borohydride salt, NaBH₄, LiBH₄, or TBABH₄ to give a soluble anionic neodymium borohydride complex, according to eqn (1):²⁸

$$Nd(Tf_2N)_3 + 4Q(BH_4) \rightarrow Q[Nd(BH_4)_4] + 3Q(Tf_2N)$$
 (1)

where $Q = Na^+$, Li^+ , TBA^+ . It is evident that in this reaction, three equivalents of a soluble bistrifimide salt are formed as a byproduct. This bistriflimide salt only contributes to the electrical conductivity of the electrolyte but has no role in the actual neodymium electrodeposition process, where $[Nd(BH_4)_4]^{-1}$ is the active species. However, it was observed that the bistriflimide anion reacted with the strongly reducing metallic rare-earth deposits to give fluorine- and sulfur-containing species on the surface, as evidenced by EDX analysis. This reactive nature of the bistriflimide anion is well reported in the literature and it is therefore recommended to not use it in the design of electrolytes for electrodeposition of neodymium and other rare-earth elements. 29-31 Makhaev and coworkers have synthesized and characterized a large variety of REE-borohydride complexes. 35-39 Anhydrous neodymium(III) chloride is readily converted to solvated borohydride complexes by means of a metathesis reaction with NaBH4 in an ether solvent, using the precipitation of NaCl as a driving force. The reaction can be performed using three or four molar equivalents of NaBH₄ to give a neutral (eqn 2) or an anionic (eqn 3) solvated complex, respectively.

$$NdCl_3 + 3NaBH_4 + 3THF \rightarrow Nd(BH_4)_3 (THF)_3 + 3NaCl$$
(2)

$$NdCl_3 + 4NaBH_4 + 4DME \rightarrow Na[Nd(BH_4)_4](DME)_4 + 3NaCl$$
(3)

After removal of NaCl by filtration, the solvated neodymium(III) complex can be isolated by crystallization (cooling) or solvent removal. These reactions afford a convenient strategy to isolate the redox-active complexes in a pure form without relying on a bistriflimide salt. We also found that anhydrous neodymium(III) methanesulfonate (Nd(OMs)₃) can be used as a substitute for the chloride in reaction (3). This salt is prepared conveniently via the reaction of neodymium(III) oxide with environmentally benign methanesulfonic acid (MSA, HOMs) (egn 4), followed by a thermal dehydration reaction (egn 5).³³

$$Nd_2O_3 + 6HOMs \xrightarrow{H_2O} 2Nd(OMs)_3(H_2O)_2 \tag{4} \label{eq:4}$$

$$Nd(OMs)_3(H_2O)_2 \xrightarrow{\Delta T} Nd(OMs)_3 + 2H_2O$$
 (5)

The neutral complex Nd(BH₄)₃(THF)₃ (eqn (2)) can be used as a precursor for the synthesis of anionic complexes with different counter cations by the addition of one equivalent of a borohydride salt (eqn (6)).

$$Nd(BH_4)_3 (THF)_3 + Q(BH_4) + (x - 3)THF \rightarrow Q[Nd(BH_4)_4](THF)_x$$
(6)

where $Q = Na^+$, Li^+ , TBA^+ . The detailed synthesis procedures of the salts and the complexes can be found in the experimental section. A powder XRD diffractogram was recorded for Na[Nd- $(BH_4)_4$ $(DME)_4$ and $Nd(BH_4)_3(THF)_3$ and these are presented in the ESI† (Fig. S1). The speciation of the neodymium(III) borohydride complexes in THF was studied by UV-Vis absorption spectroscopy (Fig. 1).

The most intense transition in the absorption spectra is the ${}^{4}G_{5/2} \leftarrow {}^{4}I_{9/2}$ transition with an absorption maximum at about 586 nm. This is a so-called "hypersensitive" transition, which means that its intensity and crystal-field fine structure are very dependent on the ligand environment around the Nd³⁺ ion, not only on the nature of the atoms in the first coordination sphere,

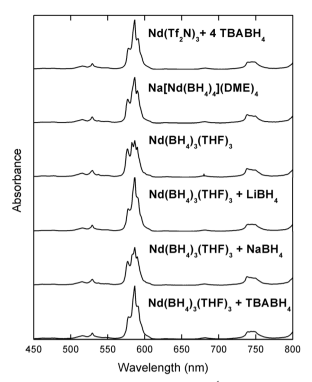


Fig. 1 UV-Vis absorption spectra of 0.1 mol L^{-1} Nd(Tf₂N)₃ + 4 molar equivalents of TBABH₄ in THF, 0.1 mol L⁻¹ isolated Na[Nd(BH₄)₄](DME)₄ in THF, 0.1 mol L⁻¹ isolated Nd(BH₄)₃(THF)₃ in THF, and 0.1 mol L⁻¹ isolated Nd(BH₄)₃(THF)₃ + 1 molar equivalent of LiBH₄, NaBH₄ and TBABH₄ in THF.

but also on the symmetry of the coordination polyhedron.⁴⁰ Since this transition reflects small changes in the coordination sphere, it can be used as a "spectroscopic probe" to monitor complex formation upon addition of a ligand. Alternatively, if two neodymium(III) complexes have a ${}^4G_{5/2} \leftarrow {}^4I_{9/2}$ transition with the same intensity and crystal-field fine structure, one can conclude that the local environment of Nd³⁺ in the 2 complexes is very similar. When Nd(Tf2N)3 is combined with 4 molar equivalents of TBABH4 in THF, the violet-colored anionic complex [Nd(BH₄)₄]⁻ is formed, which gives a very characteristic hypersensitive transition with a maximum at 586 nm.²⁸ The spectrum is very similar to those of the isolated Na[Nd(BH₄)₄](DME)₄ in THF, and the spectra of isolated Nd(BH₄)₃(THF)₃ + 1 molar equivalent of LiBH₄, NaBH₄ and TBABH₄ in THF. This indicates that the same anionic complex is present in these solutions. Only in the spectrum of neat $Nd(BH_4)_3(THF)_3$ in THF, the ${}^4G_{5/2} \leftarrow {}^4I_{9/2}$ transition has a different intensity and fine structure, indicating that the speciation of the neutral complex is indeed different from that of the anionic complex. As expected, the other absorption bands are very similar for all the complexes, and this can be expected for the non-hypersensitive transitions of Nd³⁺.

Cyclic voltammetry

An often overlooked property of electrolytes for the electrodeposition of reactive metals is how the current density of electrodeposition evolves over multiple scans in a cyclic

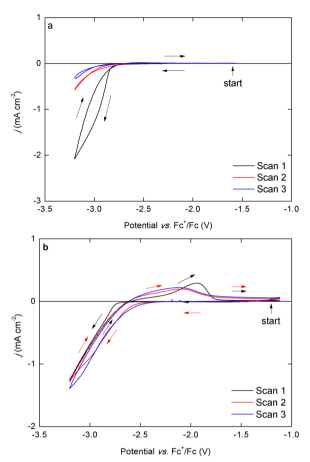


Fig. 2 Cyclic voltammograms (three scans) of (a) $0.1 \text{ mol L}^{-1} \text{ Nd}(\text{Tf}_2\text{N})_3 +$ 4 molar equivalents of NaBH $_4$ in THF, and (b) 0.1 mol L $^{-1}$ Na[Nd(BH₄)₄](DME)₄ in THF, recorded at ambient temperature at a scan rate of 10 mV s⁻¹. The scan direction is indicated with arrows. The working and counter electrodes were pieces of platinum-coated silicon wafers with a surface area of 0.3 cm² and 1.0 cm², respectively. The reference electrode was Fc⁺/Fc (0.005 mol L⁻¹ each) dissolved in [BMP][Tf₂N].

voltammogram (CV). Due to the reactivity of the deposits towards electrolyte components, non-conductive surface films can be formed which passivate the electrode, resulting in a significantly decreased current density over subsequent scans. Fig. 2 shows the CVs (three subsequent scans) of a $Nd(Tf_2N)_3$ + NaBH₄-containing electrolyte and an electrolyte based on the pure isolated Na[Nd(BH₄)₄](DME)₄ complex.

In the first scan of the CV recorded in the bistriflimidecontaining electrolyte, a high current density cathodic wave was observed with an onset potential around $-2.8 \text{ V} \text{ vs. Fc}^+/\text{Fc.}$ In the backwards scan, nucleation hysteresis was observed indicating an electrodeposition process, but a stripping peak was absent. After the experiment, a black deposit was observed on the working electrode. This cathodic wave corresponds to the electrodeposition of neodymium, because the deposition of sodium metal is only observed at a lower potential of $-3.2 \text{ V} \nu s$. Fc⁺/Fc (Fig. S2, ESI†).²⁸ In the second and third scan, the electrodeposition wave is shifted to significantly more negative potentials and the current density is significantly lower. On the other hand, for the bistriflimide-free electrolyte,

the electrodeposition onset potential was also observed around $-2.8 \text{ V} \text{ vs. Fc}^+\text{/Fc}$, but shifted to less negative potentials in subsequent cycles and the current density is not decreased. In the backwards scan, a distinct stripping peak was observed but the electrodeposition process is clearly not completely reversible, as residual deposits were also observed postexperiment on the working electrode in this case. This stripping peak definitely corresponds to the electrodeposition wave, as its size scales with the cathodic vertex potential in a broad range (Fig. S3, ESI†). The shift of the electrodeposition onset to less negative potentials is probably also caused by the incomplete stripping, as in the second and third cycle the electrodeposition occurred on the residual deposit from the first cycle, as opposed to a clean platinum substrate. These results indicate that passivation of the deposit occurs in the bistriflimidecontaining electrolyte, whereas it occurs to at least a much lesser degree in the bistriflimide-free electrolyte. This is also supported by the scan rate-dependent behavior of the current density in both electrolytes (Fig. 3).

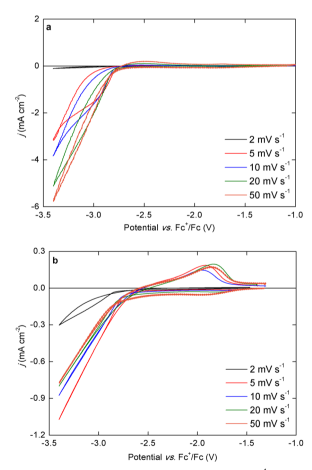


Fig. 3 Cyclic voltammograms (first scan) of (a) $0.1 \text{ mol } L^{-1} \text{ Nd}(Tf_2N)_3 +$ 4 molar equivalents of NaBH $_{4}$ in THF, and (b) 0.1 mol L^{-1} Na[Nd(BH₄)₄](DME)₄ in THF, recorded at ambient temperature at varying scan rate. The working and counter electrodes were pieces of platinumcoated silicon wafers with a surface area of 0.3 cm² and 1.0 cm², respectively. The reference electrode was Fc⁺/Fc (0.005 mol L⁻¹ each) dissolved in [BMP][Tf2N].

For the bistriflimide-containing electrolyte, the electrodeposition current density was relatively high (approx. -6.0 mA cm^{-2}) at 50 mV s⁻¹, but a negative trend was observed as the scan rate was decreased, and the current density almost reached a limiting value at 5 mV s^{-1} . At the lowest scan rate (2 mV s⁻¹), only a very small current could flow in the electrolyte as the deposited metal was fully passivated before the negative vertex potential was reached. For the bistriflimidefree electrolyte, the current density was initially lower because the total concentration of charge carriers was lower than for the bistriflimide-containing electrolyte. However, from 50 mV s⁻¹ to 5 mV s⁻¹, there was no observed negative trend for the current density. Only at 2 mV s⁻¹ the current density was lower, but still significantly higher than for the analogue CV in the bistriflimide-containing electrolyte, indicating a smaller degree of passivation. For the 50 mV s⁻¹ CV, a small pre-wave is observed starting around $-1.9 \text{ V} \text{ vs. Fc}^+/\text{Fc.}$ It is not entirely clear what the origin of this feature is and possibly, it's related to adsorption processes of the borohydride anion on the working electrode surface. The electrodeposition of neodymium from the isolated Na[Nd(BH₄)₄](DME)₄ complex was also studied in other ether solvents, such as 2-methyltetrahydrofuran (2-MeTHF), 1,2-dimethoxyethane (DME), and diglyme (G2). In each solvent, the overall electrodeposition behavior was very similar to that observed in THF, with an onset at approx. -2.8 Vvs. Fc⁺/Fc in the first cycle, which shifted to less negative potentials in the later cycles (Fig. S4, ESI†). However, the stripping behavior was notably different. Compared to the CV recorded in THF, the stripping peak in the 2-MeTHF electrolyte was larger and broader, indicating a more complete stripping of the deposit. On the other hand, in the DME and G2 electrolytes, the stripping peak was smaller and seemed to consist of two successive separate broad peaks. Because Na⁺ cations are also present in these electrolytes, and the cathodic vertex potential is sufficiently negative (-3.4 V vs. Fc^+/Fc), it is possible that co-deposition of sodium metal occurred. The peak at more negative potentials probably corresponds to the stripping of sodium while the second peak corresponds to the stripping of neodymium.

Neodymium electrodeposition was also investigated in electrolytes consisting of the neutral complex Nd(BH₄)₃(THF)₃ and one molar equivalent of LiBH₄ or TBABH₄ (Fig. S5, ESI†). This allowed for the introduction of cations other than Na⁺ in the electrolytes, thereby avoiding possible co-deposition of sodium. Similarly to the Na[Nd(BH₄)₄](DME)₄-containing electrolytes, the onset for neodymium electrodeposition in the first scan is observed at approx. $-2.8 \text{ V} \text{ vs. Fc}^+/\text{Fc}$ for the LiBH₄ electrolyte, whereas it is observed at a significantly more negative -3.1 V vs.Fc⁺/Fc for the TBABH₄ electrolyte. In the second and third scan, the onset is shifted to less negative potentials for both electrolytes. Interestingly, the electrodeposition wave is also observed at $-2.8 \text{ V} \text{ vs. } \text{Fc}^+/\text{Fc}$ for an electrolyte consisting of neat Nd(BH₄)₃(THF)₃ dissolved in THF but the current density was lower due to the neutral character of this complex, resulting in a lower conductivity (Fig. S6, ESI†). However, for both the neat Nd(BH₄)₃(THF)₃ electrolyte and the one with added TBABH₄,

a distinct stripping peak is observed. Because these electrolytes do not contain any Na+ or Li+ cations, this is evidence that this anodic process corresponds to the stripping of deposited neodymium, and not just stripping of co-deposited sodium or lithium metal. The occurrence of neodymium stripping peaks in the CVs is in contrast with the findings in our previous work, where the electrodeposition of neodymium was claimed to be completely irreversible.²⁸ However, in those bistriflimidecontaining electrolytes, it is possible that the formation of passivating surface layers inhibits the stripping of the deposits, while in these bistriflimide-free electrolytes, passivation is much less problematic.

An additional advantage of the LiBH₄- and TBABH₄electrolytes is that higher neodymium concentrations can be achieved compared to the Na[Nd(BH₄)₄](DME)₄-containing electrolytes which seem to saturate at approx. 0.1 mol L⁻¹ in DME, THF or 2-MeTHF. Because Li⁺ is more strongly solvated than Na⁺, it is possible to prepare electrolytes with a Nd³⁺ concentration of 0.5 mol L-1, and this is also the case for the organic TBA+ cation (Fig. 4). The electrolytes seem to be

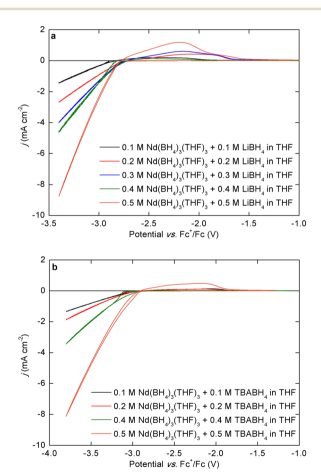


Fig. 4 Cyclic voltammograms (first scan) of (a) Nd(BH₄)₃(THF)₃ + one molar equivalent of LiBH₄ in THF at different concentrations, and (b) Nd(BH₄)₃(THF)₃ + one molar equivalent of TBABH4 in THF at different concentrations, recorded at ambient temperature at a scan rate of 10 mV $\rm s^{-1}$. The working and counter electrodes were pieces of platinum-coated silicon wafers with a surface area of 0.3 cm² and 1.0 cm², respectively. The reference electrode was Fc⁺/Fc (0.005 mol L⁻¹ each) dissolved in [BMP][Tf₂N].

saturated at this concentration, as no more of the complex would dissolve.

In both electrolytes, a positive trend was observed for the current density as the concentration of the neodymium complex is increased, with a value of almost -10.0 mA cm⁻² reached for the most concentrated LiBH4-electrolyte at a cathodic vertex potential of $-3.4 \text{ V} \text{ vs. Fc}^+/\text{Fc.}$ This is a very promising result for reaching sufficiently high current densities during potentiostatic electrodeposition without relying on supporting electrolyte salts as in the case of the bistriflimide-containing electrolytes. As the amount of electroactive species is increased, the current should increase linearly with the concentration, whether the polarization is steady or unsteady. It is evident from Fig. 4 that this is not the case. This implies that the neodymium metal that is being electrodeposited, reacts with the electrolyte, reducing the electroactive part of the electrode. As the concentration of electro-active species in solution is increased, neodymium metal is depositing faster than it can react with the electrolyte and the current is increasing faster than the increase in concentration. This also explains why the anodic currents are larger after crossing the zero-current line. It also explains why in the CVs in Fig. 3, the cathodic current decreases with increasing scan rates for the same concentration of electroactive species. As the scan rate decreases, more time is available for the deposited neodymium metal to react with the electrolyte, which results in the partial passivation of the working electrode.

Potentiostatic electrodeposition

Electrodeposition of neodymium was performed potentiostatically at -3.0 V or -3.1 V vs. Fc⁺/Fc for the Na⁺- and Li⁺containing electrolytes, respectively in order to avoid excessive co-deposition of sodium or lithium metal (Fig. 5). For the TBABH₄ electrolytes, the potential was $-3.3 \text{ V } \nu s. \text{ Fc}^+/\text{Fc}$ to compensate for the higher deposition overpotential.

Similarly to the cyclic voltammetry experiments, the electrodeposition current density was initially higher for the bistriflimide-containing electrolytes at approx. -2.6 mA cm⁻², but quickly decreased during the first 500 s of the experiment to less than 4% of this initial value. For the rest of the duration of the experiment, a plateau was observed at a negligibly small current density (<-0.1 mA cm⁻²). This behavior is again indicative of the formation of passivating surface films due to side reactions of the neodymium deposit with the bistriflimide anion. On the other hand, for the bistriflimide-free electrolyte, the current density was initially lower at approx. -1.0 mA cm^{-2} but remained nearly constant over the duration of the experiment, indicating that the deposit was not passivated during electrodeposition. As a result, it was possible to deposit much more charge in the bistriflimide-free electrolyte (-1.03 C, as)opposed to -0.14 C for the bistrifimide-containing electrolyte), and to achieve thicker deposits. More concentrated electrolytes consisting of 0.5 M of the neutral complex Nd(BH₄)₃(THF)₃ + one molar equivalent of LiBH4 or TBABH4 in THF were studied as well (Fig. S7, ESI†). Due to the higher concentration, the electrodeposition current density was initially higher for the

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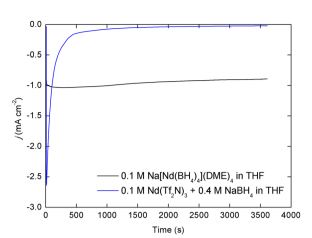
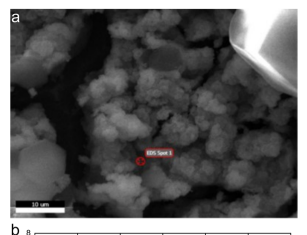


Fig. 5 Chronoamperograms recorded at ambient temperature for a potentiostatic deposition at -3.0 V vs. Fc⁺/Fc for 3600 s in a static solution of 0.1 mol L⁻¹ Na[Nd(BH₄)₄](DME)₄ in THF (black curve), and 0.1 mol L⁻¹ Nd(Tf₂N)₃ + 4 molar equivalents of NaBH₄ in THF (blue curve). The working and counter electrodes were pieces of platinum-coated silicon wafers with a surface area of 0.3 cm² and 1.0 cm², respectively. The reference electrode was Fc⁺/Fc (0.005 mol L⁻¹ each) dissolved in [BMP][Tf₂N].

LiBH $_4$ electrolyte (approx. -6.4 mA cm $^{-2}$), but a gradual decrease was observed over the duration of the experiment, to 20% of the initial value after one hour. For the TBABH $_4$ electrolyte, the current density was initially lower (approx. -2.5 mA cm $^{-2}$), but remained much more constant over time. A scanning electron micrograph and corresponding EDX analysis of a neodymium deposit is shown in Fig. 6.

A spongy, nodular morphology was observed, with deep cracks dividing the deposit. This is in contrast to the smooth shard-like morphology observed in previous work.²⁸ Some crystalline particles are observed on top of the cracked layer, which are most likely crystals of electrolyte salts (NaBH₄) that were not rinsed off completely, judging by EDX analysis (Fig. S8, ESI†). According to the EDX analysis of the deposit itself, neodymium (L α = 5.23 keV, M = 0.98 keV) is present as the major constituent at an estimated 75 atomic %, while oxygen ($K\alpha = 0.53 \text{ keV}$) is only present at 3.5 at%. The main impurity of the deposit is co-deposited sodium ($K\alpha = 1.04 \text{ keV}$) at an estimated 21 at%. It should also be noted that typical impurities such as fluorine ($K\alpha = 0.68 \text{ keV}$), sulfur ($K\alpha = 2.31 \text{ keV}$), chlorine ($K\alpha = 2.62 \text{ keV}$) and carbon ($K\alpha = 0.28 \text{ keV}$) are not detected in deposits originating from the bistriflimide-free electrolytes, providing further evidence that no passivating films are formed. However, this elemental composition is not representative for the entire deposit, as EDX analysis of other spots on the surface showed a significantly higher oxygen content, up to 67 at% (Fig. S8, ESI†). During transfer to the SEM vacuum chamber it was not possible to avoid short exposure of the deposit to air, therefore resulting in partial oxidation, although the increased layer thickness seems to have somewhat compensated for this. Nevertheless, this is a very promising result, giving evidence that the electrodeposition of pure metallic neodymium layers is possible.

Motivated by this positive result, the black deposits resulting from 0.1 mol $\rm L^{-1}$ Na[Nd(BH₄)₄](DME)₄ in THF were further analyzed by X-ray photoelectron spectroscopy (XPS) surface and



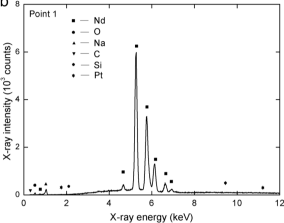


Fig. 6 (a) SEM micrograph of a neodymium deposit (theoretical thickness = $2.8~\mu m$) at $2000\times$ magnification, obtained after potentiostatic electrodeposition from $0.1~mol~L^{-1}~Na[Nd(BH_4)_4]$ in THF at -3.0~V vs. Fc⁺/Fc for 2.5 hours at ambient temperature. The substrate was a piece of platinum-coated silicon wafer with an approximate surface area of $0.3~cm^2$. (b) EDX spectrum in the range 0-12~keV corresponding to the spot indicated on the SEM micrograph, recorded at an acceleration voltage of 30~kV.

depth profiling to investigate the composition changes in the inner layers of the deposits. The deposits were transported under argon and placed inside an air-tight vessel which was directly attached to the ultra-high vacuum line of the XPS instrument in order to avoid exposure to air. The Nd $3d_{5/2}$ spectra for the sample at various depths are shown in Fig. 7.

The Nd $3d_{5/2}$ region of the XPS spectra was deconvoluted into three peaks: Nd_2O_3 (>982 eV), neodymium metal (979–980 eV), and the O $KL_{23}L_{23}$ auger electron line. 13,21,41 At the surface, the lines of Nd_2O_3 and neodymium metal are observed, giving evidence for the presence of both species. As the etching progressed from the surface (*i.e.* at 0 s etching) to the inner layers, an increase in the Nd $3d_{5/2}$ peak intensity was observed, indicating an increased amount of both species present. The relative atomic concentrations of Nd_2O_3 and neodymium metal determined from the spectra at each etching time are summarized in Fig. 7(e). At the surface, neodymium metal is present at 51% of the total neodymium content and Nd_2O_3 at 49%. Surprisingly, as the etching progresses, the content of neodymium metal is slightly decreased to approx.

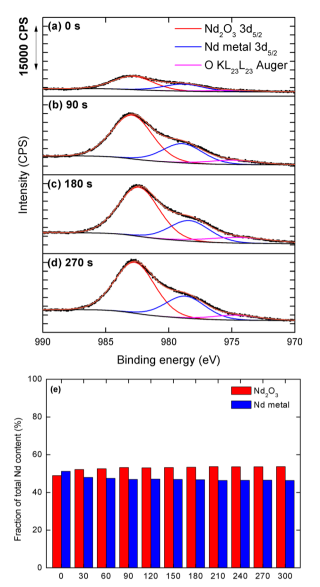


Fig. 7 (a)-(d) Curve-fitted Nd 3d_{5/2} region XPS spectra after 0 s, 90 s, 180 s and 270 s of etching time from a neodymium deposit obtained from potentiostatic deposition in 0.1 mol L⁻¹ Na[Nd(BH₄)₄] in THF electrolyte at -3.0 V vs. Fc $^+$ /Fc for 2.5 hours at ambient temperature. (e) Comparison of neodymium metal and Nd₂O₃ content after different etching times.

47%, whereas the content of Nd_2O_3 is increased to approx. 53%, which is opposite to the expected trend in case of surface oxidation. Furthermore, the relative content of Nd₂O₃ and neodymium metal remains constant over the entire depth of the deposit. However, this could be a result of the rough, spongy morphology of the deposit. This was also observed by Periyapperuma et al., who hypothesized that the presence of water in their ionic liquid electrolytes was responsible for constant side-reactions during electrodeposition, resulting in oxide formation throughout the entire deposit.²¹ However, the organic electrolytes in this work are completely water-free, due to the reactivity of the borohydride anions. Therefore, the origin of Nd₂O₃ throughout the entire thickness of the deposit is unclear, other than maybe a side-reaction with the THF solvent. It is also surprising that the presence of oxides does not result in passivation during electrodeposition, as was observed during the CV and CA experiments. Nevertheless, both Nd₂O₃ and Nd metal are considered valuable materials to recover due to their many applications. The survey spectrum of the surface of the deposit is shown in Fig. S9 (ESI†). Other than the peaks corresponding to neodymium, the most prominent other peaks correspond to oxygen (O 1s) and carbon (C 1s), whereas peaks of typical bistriflimide-derived elements such as fluorine or sulfur are completely absent.

Conclusions

A new class of fluorine-free organic electrolytes was developed for the electrodeposition of neodymium metal. The electrolytes consist of an isolated solvated neodymium borohydride active complex, Na[Nd(BH₄)₄](DME)₄ or Nd(BH₄)₃(THF)₃, dissolved in an ether solvent. These complexes can be prepared from commercially available NdCl3 or the conveniently prepared Nd(OMs)3, and the borohydride salt NaBH4, without relying on any salts with fluorinated anions such as bis(trifluoromethylsulfonyl)imide (bistriflimide). Because the electrolytes do not contain this reactive anion, they are much more stable against the metallic neodymium deposits, resulting in the possibility to perform potentiostatic electrodeposition at constant current density, without the interference of passivating surface films. The deposits were characterized by different analytical techniques, giving evidence for the presence of both Nd_2O_3 and Nd metal throughout the entire deposit. Despite these promising results, there are some obvious limitations associated with this system that must be addressed in the context of electrowinning of neodymium. Firstly, the borohydride-based organic electrolytes are highly moisture-sensitive, which requires handling in a rigorously controlled dry and inert atmosphere. Secondly, the electrical conductivity of organic electrolytes is always lower than that of any molten salt electrolyte, resulting in lower deposition current densities. Thirdly, the reaction at the counter electrode must be considered as well. The anode reaction is probably oxidation of the borohydride anion in this case, which results in gradual contamination of the electrolyte by decomposition products. Nevertheless, we are convinced that this system is a leap towards low-temperature electrowinning of neodymium and other rare earths in fluorine-free electrolytes and that it can be significantly improved by further research.

Conflicts of interest

There are no conflicts to declare.

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