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1	Earth-abundant elements a sustainable solution for electrocatalytic
2	reduction of nitrate
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Abstract

21 Platinum group elements (PGEs) are widely-used electrocatalysts. However, the low 22 abundance of PGEs in the earth's crust and high environmental impacts to be acquired result in high 23 costs, limiting their use in drinking water treatment. Identifying sustainable alternatives to PGEs is a 24 major barrier in applying electrocatalysis for nitrate reduction. By moving up the periodic table, this 25 study provides a framework for selecting promising earth-abundant elements that can electrocatalytically degrade nitrate in water to innocuous by-products. We benchmarked platinum 26 (Pt) against less-endangered elements for electrodes by quantifying nitrate reduction rates, by-27 28 product selectivity, and energy efficiencies. Carbon (as boron-doped diamond) and tin had the 29 highest average selectivity towards nitrogen gas evolution (55% and 64%, respectively) 30 outperforming Pt, which only had 1% selectivity, and had comparable electrical energy per order 31 removal of nitrate. Thus, earth-abundant elements for electrocatalysis hold tremendous promise as 32 innovative, low-cost, and sustainable processes for the water treatment marketplace.

33

34 Keywords: water treatment; advanced reduction processes; electrochemical technologies; cathodic

35 materials; selectivity towards nitrogen

1. Introduction

37 Ensuring access to safe water and sanitation is essential to health and a human right[1]. Nitrate pollution is one of the top ten most common water quality violations reported worldwide[2– 38 39 5]. The World Health Organization (WHO) and United States Environmental Protection Agency set maximum concentration levels in drinking water of ~10 mg NO₃⁻-N L⁻¹.[6,7] Elevated NO₃⁻ levels 40 41 in drinking water can cause cancer, thyroid problems, and adverse respiratory effects[8–11]. As a consequence of anthropogenic nitrogen fertilizer inputs, nitrate concentrations in surface and 42 43 groundwater have dramatically increased during the last century[12]. There are ~45 million people 44 in the United States that rely on unregulated private groundwater wells, many of which have nitrate 45 above regulated limits[2,13]. In addition, many municipal water supplies are impacted by nitrate 46 and are commonly treated using costly ion exchange technology [14–19]. Thus, there is a pressing 47 need for efficient nitrate removal technologies suitable for large-scale water treatment systems as well as very small point-of-use (POU) treatment systems within homes[20,21]. In a comprehensive 48 49 review, we recently summarized how electrocatalytic reduction of nitrate (ERN) can selectively 50 reduce NO_3^- to innocuous dinitrogen (N₂) and serves as a viable drinking water treatment 51 technology[22]. Because electrode cost dominates the capital cost of electrochemical systems, a 52 thorough revision on the state of the art with detailed description of the electrodes applied to (ERN) was performed as well[22-24]. The majority of publications on ERN utilized platinoid materials 53 54 (i.e., Pt and Pd)[25–34]. Platinum electrodes are excellent electrocatalysts for nitrate reduction in 55 the water-energy nexus due to its corrosion resistance [35–37]. Unfortunately, platinum group elements (PGEs) are labeled as expensive and endangered elements due to their high cost and 56 57 limited availability as resources in our planet[38]. Lifecycle analysis pinpoints adverse environmental impacts related to extraction and purification of these endangered elements[39-42]. 58 59 Scientific and engineering challenges lie on uncovering alternative electrocatalysts based in high-60 efficient earth-abundant elements.

62 Earth-abundant alternatives to PGE electrodes likely exist but have been barely explored. Analogues for electrocatalytic responses using non-endangered substitutes to PGE might be 63 identified by moving up the periodic table. This research aimed to provide a framework for 64 selecting promising earth-abundant elements to electrocatalytically degrade nitrate in drinking 65 water. To the best of our knowledge, this is the first time a paper explored, under identical 66 experimental conditions, a direct comparison of PGE-based electrodes against electrodes fabricated 67 from earth-abundant elements of the first row transition metal d-block elements (e.g., Ti, Fe, Co, Ni, 68 69 Cu, Zn) plus carbon (C) and tin (Sn). Operating conditions were selected to minimize side reactions 70 with other electroactive species that may impact the fundamental understanding of the electrochemical transformations. Nitrate degradation rates, formation of aqueous intermediates 71 72 (nitrite and ammonium ion), and selectivity towards gaseous by-products were quantified. Nitrogen 73 gas is the preferred nitrate reduction by-product because (a) nitrite remains toxic to humans, (b) 74 ammonia has odor and potential human health concerns, (c) ammonia reacts with chlorine 75 disinfectants, and (d) nitrogen gases are permanently removed from solution. Engineering figures of 76 merit were used to evaluate operational requirements for electrical energy per order (E_{EO}) and 77 efficient use of electron delivered in terms of Faradaic efficiency. Results are discussed for relative 78 cost and efficiency of earth-abundant elements as alternatives to Pt electrocatalysts.

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- 80

2. Materials and methods

81 **2.1 Electrochemical experiments**

Electrochemical nitrate reduction was conducted using an open, undivided cylindrical glass batch reactor containing 200 mL of non-deaerated 100 mg NO₃⁻-N L⁻¹ solutions with 50 mM Na₂SO₄ (pH = 5.86 ± 0.07 and conductivity = 7.74 ± 0.08 mS cm⁻¹) at 25 °C. Reagent grade sodium nitrate, sodium nitrite, and ammonia sulfate (>99%) were purchased from Sigma-Aldrich. Analytical-grade sodium sulfate (99%, Sigma-Aldrich) was used as the supporting electrolyte. All

solutions were prepared with ultrapure water with resistivity >18.2 M Ω cm at 25 °C (Millipore Milli-Q system).

89 Batch reactor experiments were continuously mixed using magnetic stirring at 700 rpm to 90 reduce mass transport limitations between the bulk solution and the electrode surfaces. The electrochemical cell was equipped with commercial sheet parallel electrodes of 2.5 cm × 2.4 cm 91 92 with a defined geometrical area of 6.0 cm² (area delimited with Teflon, back part and sides). The 93 interelectrode gap distance was 1.5 cm. A Ti/IrO₂ (DeNora) anode, and different cathodes were 94 tested: titanium (99.9%, McMaster-Carr), stainless steel (as iron element, Victor Monteiro), cobalt 95 (99.9%, Fine Metals), nickel (99.6%, Trudsafe), copper (99.9%, CynKen), zinc (99.9%, Filzinc), carbon cloth (99.5% carbon content, Fuel Cell Store), carbon felt (99.0% carbon content, Fuel Cell 96 97 Store), synthetic boron-doped diamond (BDD; Adamant Technologies), tin (99.9%, McMaster-Carr), and a polycrystalline platinum (99.9 %, Stanford Advanced Materials). The electrocatalysts 98 99 were rinsed with ultrapure water before use. The carbon-based materials (see specifications in 100 Supplementary Information (SI) Table S1) were activated in a solution containing 50 mM of Na₂SO₄ at 100 mA cm⁻² for 120 min. 101

102 Experiments were conducted galvanostatically using a TENMA 72-13610 DC power supply under constant applied current density of 20 mA cm⁻². To evaluate the possible re-oxidation 103 104 of reduction by-products to nitrate, control experiments with Sn were performed using initial 105 solutions of 100 mg L^{-1} NO₂⁻-N or 100 mg L^{-1} NH₃-N solutions with 50 mM Na₂SO₄ the details are 106 shown in SI, Figure S1a and S1b, respectively. It was detected that a slight oxidation of nitrite to 107 nitrate may occur on the surface of the anode. However, that re-oxidation did not seem meaningful 108 when compared to nitrite's reduction to nitrogen gas (Figure S1a). No ammonia oxidation was 109 observed (Figure S1b). Because the oxidation reactions occur at the anode and the electrode 110 material was always the same, only one control was needed. Samples were collected over time and 111 analyzed for aqueous nitrogen species, conductivity, and pH. Experiments were run in triplicate, 112 and deviations between them were lower than 5% for all trials.

114 **2.3 Analytical techniques**

115 The pH and conductivity were measured over time using a 2 Hanna Instruments HI 322 and 116 VWR Scientific Products - EC Model 2052 meters, respectively. Nitrate and nitrite were quantified 117 using ionic chromatography. Sample aliquots of 20 µL were injected in Thermo Dionex ICS-118 5000DC equipment coupled to a conductivity detector AERS 500, fitted with a high capacity hydroxide-selective anion-exchange column Dionex Ionpac AS18 (2 mm × 250 mm) column at 30 119 °C. A 10-45 mM KOH gradient solution at 0.25 mL min⁻¹ was used as mobile phase per EPA 300 120 121 method[43]. Aqueous ammonia was measured according to Hach Method 10205, a salicylate-based 122 ammonia chemistry that is equivalent to EPA 350.1, EPA 351.1, and EPA 351.2. Measurements 123 used HACH DR6000 UV-vis equipment at 694 nm applying TNT 830 HACH kits. Reproducible 124 NO₃⁻-N, NO₂⁻-N, and NH₃-N concentrations were obtained with an accuracy of 1%. Nitrate removal 125 (NR) was calculated using Eq. (1)[44].

$$NR(\%) = \frac{C_{nitrate,i} - C_{nitrate,t}}{C_{nitrate,i}} \times 100$$
(1)

where $C_{nitrate,i}$ is the nitrate concentration in mg NO₃⁻-N L⁻¹ at time zero, and $C_{nitrate,t}$ is the nitrate concentration at time (*t*).

128 Some publications quantified specific N-volatile species (N2, NO, NO2 or N2O) that were 129 generated during the electrocatalytic reduction of nitrate. However, most of the works report all the obtained gas was N₂. Therefore, in this work, the evolution of gaseous nitrogen species determined 130 131 from mass balances on aqueous nitrogen species assumed noted as N-gas, but likely corresponds mostly to N₂. We do report ammonium ion concentrations, and near or above the pKa of 9.25 for 132 133 NH_4^+/NH_3 some ammonia volatilization can occur but the Henry's constant is not very large and 134 such losses from solution of ammonia gas would be highly depending upon mixing or gas-purging 135 conditions. The selectivity (S_X) towards N-gas evolution and ammonia was calculated using Eq. 136 (2)[45].

$$S_X(\%) = \frac{C_X}{C_{nitrate,i} - C_{nitrate,t}} \times 100$$
(2)

137 where C_X represents the concentration (mg N L⁻¹) of a species, X, produced over time.

The NO₃⁻-N decay followed pseudo-first order reaction kinetics and was fitted (R²>0.95) by a rate constant (k_1 , s⁻¹). Because the electrode surface area (6 cm²) was the same for all experiments, k_1 (s⁻¹) can be directly converted to a surface-normalized area ($k_1/6$ s⁻¹cm²). Nitrate reduction was evaluated in terms of Faradaic efficiency (FE, Eq. (3)), which determines the number of electrons consumed in an electrochemical reaction relative to the expected theoretical conversion ruled by Faraday's law[22].

$$Fe(\%) = \frac{n F N_i}{3600 I t} \times 100$$
(3)

where *n* is the amount of electrons required per mol of product (mol), *F* is the Faraday constant (96 487 C mol⁻¹), N_i is the amount (mol) of product generated during the electrolysis, *I* is the applied electric current (A), *t* is the electrolysis time (h), and 3600 is a unit conversion factor (3600 s h⁻¹).

147 Electrical energy per order (E_{EO}), was used as an engineering figure of merit to benchmark 148 the electric energy required to reduce NO₃⁻-N concentration by one order of magnitude in a unit 149 volume calculated from Eq. (4) for batch operation mode[46].

$$E_{EO}(kWh\,m^{-3}order^{-1}) = \frac{E_{cell}lt}{V_s log(C_0/C_t)}$$
(4)

where E_{cell} is the average of the cell potential (V), *I* is current intensity (A), *t* is time (h), *V_s* is solution volume (L), and C_0 and C_t are the initial and final concentration after one order of magnitude reduction.

153 Considering the relationship $\log(C_0/C_t) = 0.4343 \cdot t \cdot k_1$, the E_{EO} expression can be simplified 154 assuming first-order kinetics according to Eq. (5) where 6.39×10^{-4} is a conversion factor:

$$E_{EO}(kWh \, m^{-3} order^{-1}) = \frac{6.39 \times 10^{-4} E_{cell} I}{V_s k_1} \tag{5}$$

3. Results and discussion

157

3.1 Platinum electrode performance as reference electrocatalyst for nitrate reduction

158 Electrochemical reduction of nitrate has been focused mostly on the study of platinum as 159 standard cathodic material [47–49], therefore Pt is used herein to benchmark alternative electrodes 160 based on more sustainable elements. Figure 1 shows nitrate reduction and evolution of nitrite 161 intermediates as ammonium ion and nitrogen gas by-products. A pseudo-first order rate constant (k_1) fits well the observed nitrate degradation $(k_1 = 3.7 \times 10^{-5} \text{ s}^{-1}; \text{ R}^2 = 0.996)$. The sluggish reduction 162 163 kinetics on Pt is controlled by the first charge transfer reaction as limiting step (Eq. (6)). Bimetallic 164 electrodes on PGEs are often required to reduce this barrier[50-54], but this was beyond the scope 165 of this study. The initial reduction of nitrate towards nitrite follows a three-step electrochemical-166 chemical-electrochemical (ECE) mechanism described by Eqs. (6)-(8)[22].

$$NO_{3}(ad) + e^{-} \rightarrow NO_{3}^{2^{-}}(ad)$$
 limiting step (6)

$$NO_3^{2-}_{(ad)} + H_2O \rightarrow NO_2^{\bullet}_{(ad)} + 2OH^-$$

$$\tag{7}$$

$$NO_2^{\bullet}_{(ad)} + e^- \rightarrow NO_2^{-}_{(ad)}$$
(8)

167

168 Nitrate reduction by-products kinetics did not show appreciable accumulation of NO₂intermediates; a maximum concentration of 2.2 mg NO₂-N L⁻¹ was reached at 120 min (Figure 1b). 169 170 The observed trend occurs due to the high electrocatalytic activity of Pt reducing nitrite to nitrogen 171 gas and/or ammonia following reactions (9) and (10), respectively [55,56]. The net reduction of 172 nitrate presented takes into account that nitrite yielded may be re-oxidized at the anode in the 173 undivided cell (see Fig. S1a). According to reactions (9) and (10), hydroxide ions (i.e., consumption 174 of protons during nitrate reduction) are also generated, increasing the pH from 5.86±0.07 to 175 11.3 \pm 0.1 over time as shown in Figure 1a (dashed line). In this pH range, N₂ and NH₃/NH₄⁺ are the 176 thermodynamically most stable forms of nitrogen according to Frost-Ebsworth diagrams[22,57]. 177 Note that ammonia volatilization was not observed in blank experiments. Fundamental 178 electrochemical studies typically do not buffer pH, and effects of constant pH and other water

quality parameters (e.g., alkalinity, ion composition) could be evaluated in a second-tier framework study. Note that during the reduction of nitrate higher amount of OH⁻ is yielded by electron (1.3 OH⁻/1e⁻for nitrogen gas and 1.2 OH⁻/1e⁻ for ammonia) than H⁺ produced in the anodic oxidation of water (1 H⁺/1e⁻). Thus, the pH change observed also allows inferring that nitrate reduction takes place.

$$NO_{2^{-}(ad)} + 2H_2O + 3e^{-} \rightarrow \frac{1}{2}N_{2(gas)} + 4OH^{-}$$
 (9)

$$NO_{2^{-}(ad)} + 5 H_{2}O + 6e^{-} \rightarrow NH_{3} + 7OH^{-}$$

$$\tag{10}$$

184

After 360 min of reaction, a 58% nitrate loss yielded 49.5 mg L⁻¹ NH₃-N and 5.8 mg L⁻¹ N₂-185 N. The calculated selectivity towards nitrogen evolution $(S_{N_{gas}})$ was only 10% with preferential 186 187 production of ammonia (S_{NH_3} = 87%). The Pt selectivity is closely related to crystallographic 188 availability at the interface[58]. Fundamental studies using monocrystalline electrodes have 189 identified Pt(100) domains as essential for selective reduction towards N₂ following the Duca-Feliu-190 Koper mechanism[59–61]. However, synthesis of monocrystalline Pt electrodes limits the electrode 191 size to a few millimeters. Introducing defects in crystalline structure affects selectivity of Pt 192 catalysts. Nitrogen evolution can be drastically reduced by the defects in symmetry as observed at 193 Pt [(1 0 0) x (1 1 0)] and [(1 0 0) x (1 1 1)] surfaces[62]. This is evident in commercial 194 polycrystalline Pt electrodes where all crystallographic facets are present in a random distribution 195 rather than a preferential orientation such as the one reported in single crystal Pt. This is the reason 196 for reduced selectivity towards nitrogen evolution observed in polycrystalline Pt electrodes, with 197 the characteristic behavior depicted in Figure 1.

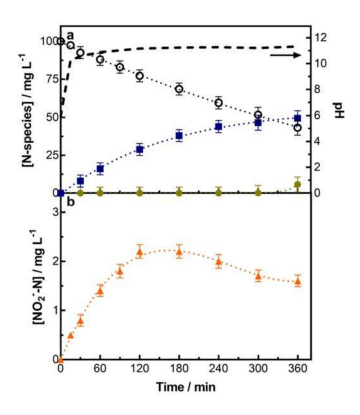
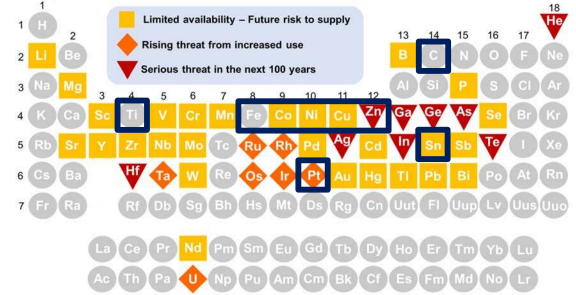


Figure 1. (a) Evolution of the nitrogenated species ((O) NO₃⁻-N, (\blacksquare) NH₃-N, and (\bullet) N₂-N) and pH (long-dashed line) over time for the electroreduction of 100 mg L⁻¹ NO₃⁻-N in 50 mM Na₂SO₄ at 20 mA cm⁻², using Pt as cathode material. (b) Evolution of nitrite ((\blacktriangle) NO₂⁻-N) over time.

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203 **3.2** Nitrate reduction kinetics of electrocatalysis materials upon moving up the periodic table

204 Figure 2 shows the periodic table and identifies "endangered elements", which reflect their 205 relative earth abundance[38]. While platinum is considered a standard in electrocatalysis due to its 206 excellent properties, it is unfortunately an endangered element because of its limited abundance and 207 increasing use in products or processes[38]. The high electrocatalytic activity of Pt is associated 208 with the unclosed d-orbital shells of the metal, which contributes to the charge injection to the 209 lowest unoccupied molecular π^* orbital of nitrate[22,63]. Transition metals resembling the electron 210 configuration of Pt are suitable for nitrate reduction but have not been the focus of comparative 211 studies. Thus, we compared the performance of transition metals for nitrate removal. By "moving 212 up the periodic table", catalysts using transition metals are more earth abundant and lower cost than



213 Pt. The elements from the fourth period of the periodic table accomplish that requirement.

Figure 2. The periodic table's endangered elements (adapted from open source[38]). Elements
availability: (●) high abundance, (■) limited availability, (◆) rising threat, (▼) serious threat in
next 100 years.

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Figure 3 depicts the performance of several transition metals materials to reduce nitrate under the same operating conditions. Fitted k_1 values fell within a narrow range (1.3 to $6.6 \times 10^{-5} \text{ s}^{-1}$) (statistical fit details shown in SI, Figure S2). The low variability suggests a possible mass transfer rate limit that can be overcome through reactor engineering design.

Titanium (Ti) and iron (Fe) are widely available elements without a close depletion risk (Figure 2), but few direct comparisons of nitrate removal exist in the literature. Figures 3a and 3b illustrate that Ti ($k_1 = 2.8 \times 10^{-5} \text{ s}^{-1}$, R²=0.993) and Fe ($k_1 = 1.8 \times 10^{-5} \text{ s}^{-1}$, R²=0.997) electrodes both have ~2.0-fold slower nitrate reduction kinetics when compared with Pt. Table 1 summarizes key fitted and calculated parameters from all the experiments. For the Fe electrode, nitrite concentrations peaked at 2.7 mg L⁻¹ NO₂⁻-N and followed trends similar to Pt. After 360 minutes of reaction, the residual ammonium ion was 26 mg NH₃-N L⁻¹. $S_{N_{gas}}$ and S_{NH_3} using Fe were about 11% and 80%, respectively, also similar to Pt. For Ti electrode, nitrite accumulation peaked at 5.6 mg L⁻¹ NO₂⁻-N at 180 min and decreased slowly to reach 4.5 mg L⁻¹ NO₂⁻-N at 360 min. Consistent with the literature[55,64–66], our Ti and Fe results showed ammonia generation was electrocatalytically preferential over nitrogen gas evolution.

234 Cobalt (Co), nickel (Ni), and copper (Cu) are within the group of elements with limited 235 availability and possible future supply risk (Figure 2). Nitrate kinetics for these metals are shown in 236 Figures 3c, 3d, and 3e; Table 1 summarizes associated key parameters, and Figure S3 shows pH 237 evolution over time. Co had the highest nitrate reduction kinetics (~1.8-fold higher than Pt) and produced the lowest nitrite (<0.08 mg NO₂⁻-N L⁻¹) throughout the experiment. Ni electrodes 238 achieved a high $S_{N_{gas}}$ (36%) but had amongst the slowest kinetics. Cu electrodes performed 239 240 between Ni and Co. Electrocatalytic reduction with Cu exhibited a slightly higher k_1 than Pt, accumulated 3.8 mg NO₂⁻⁻N L⁻¹, and had $S_{N_{gas}}$ and S_{NH_3} of 28% and 64%, respectively. Cu 241 catalyzes the initial charge injection illustrated by Eq. (6), which is a rate limiting reaction for nitrite 242 243 formation. However, prolonged electrolysis preferentially reduces nitrite to ammonia instead of 244 nitrogen gas.

Zn electrodes had slightly higher k_1 than Pt (Table 1) and reached 60% nitrate reduction after 360 min of treatment (Figure 3f). Zn showed the highest nitrite accumulation among the evaluated elements in the fourth period of the periodic table, with maximum value of 17 mg NO₂⁻-N L⁻¹ at 240 min, decreasing during the following 120 min to 13 mg NO₂⁻-N L⁻¹.

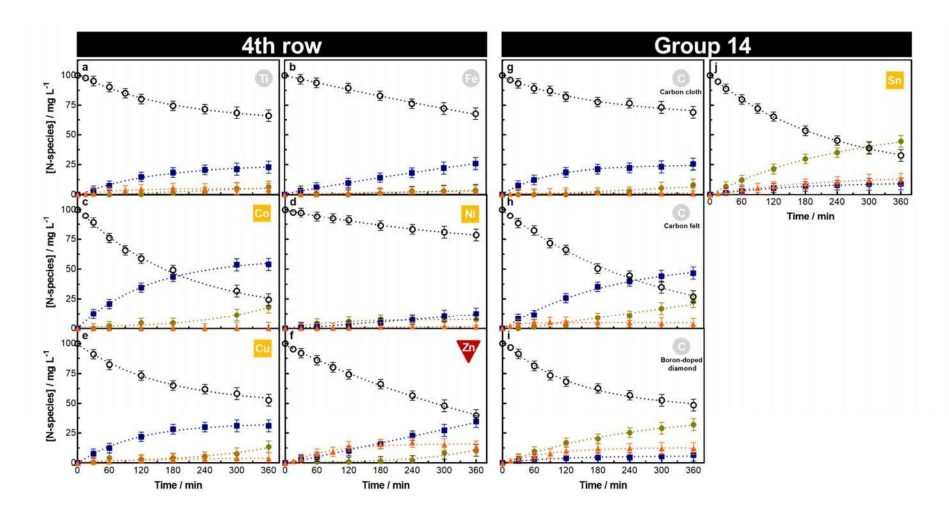




Figure 3. Evolution of the nitrogenated species ((O) NO₃⁻-N, (\blacktriangle) NO₂⁻-N, (\blacksquare) NH₃-N, and (\bigcirc) N₂-N) over time for the electroreduction of 100 mg L⁻¹ NO₃⁻-N in 50 mM Na₂SO₄ at 20 mA cm⁻² using (a) Ti, (b) Fe, (c) Co, (d) Ni, (e) Cu, (f) Zn, (g) carbon cloth, (h) carbon felt, (i) boron-doped diamond, and (j) Sn.

Electrode material	$k_{l}, \times 10^{-5} (\mathrm{s}^{-1})$	NR (%)	S _{Ngas} (%)	S _{NH3} (%)	Maximum (and t=360 min) nitrite (mg NO ₂ ⁻ -N L ⁻¹)
Pt	3.7±0.2	57±2	10±1	87±3	2.2±0.2 (1.6±0.3)
Ti	2.8 ± 0.4	34±1	19±1	68±2	5.6±0.3 (4.5±0.5)
Fe	1.8±0.1	31±1	11±1	80±2	2.7±0.2 (2.7±0.2)
Со	6.6±0.5	76±3	25±2	75±1	0.08±0.01 (0.03±0.01)
Ni	1.3±0.2	21±1	36±1	58±1	1.6±0.3 (1.3±0.2)
Cu	4.2±0.3	47±2	28±2	64±3	3.8±0.4 (3.7±0.3)
Zn	4.1±0.3	60±1	18±1	59±2	17±2 (13±1)
Sn	5.4±0.4	67±3	67±3	14±1	13±1 (13±1)
Carbon					
Cloth	2.8±0.1	31±1	24±1	74±2	1.8±0.3 (0.7±0.1)
Felt	5.9±0.3	73±3	31±2	64±2	5.7±0.4 (3.6±0.3)
BDD	5.5±0.2	51±2	63±2	13±1	12±1 (12±1)

 $100 \text{ mg } \text{L}^{-1} \text{ NO}_3$ -N in 50 mM Na₂SO₄ at 20 mA cm⁻² and 360 min of treatment time.

Tin (Sn) is not a 4th period d-block element in the periodic table, but it is widely reported as an electrocatalyst[22]. Figure 31 shows that the tin electrode achieved 67% nitrate removal and yielded the highest mass production of N-gases (44.5 mg N L^{-1}). Tin enhanced nitrate removal with kinetics 1.5-fold higher than Pt and achieved the highest nitrogen gas concentration among the electrode materials studied.

Each electrode material led to different trends for nitrate reduction and by-product formation. The mechanism by which the material transfers electrons more easily to adsorbed nitrate determines its suitability for pollutant removal. Nitrite is the first stable intermediate produced during nitrate reduction. The further electrochemical reduction of this species can yield ammonia (reaction (10)) and innocuous nitrogen gas (reaction (9)). Nitrite and ammonia are harmful to human health and cause operational problems in drinking water systems, therefore making them undesired by-products from nitrate reduction[22].

269 **3.2.2. Carbon-based materials**

270 Carbon is one of the most abundant elements on Earth[38]. Carbon-based materials have a 271 diversity of structures, good electrical, thermal conductivity, and high mechanical strength, making carbon stand out as an electrode material[67]. Carbonaceous electrodes, such as carbon cloth and 272 273 carbon felt, can be 1000-fold cheaper than PGEs. Carbon cloth and felt (Figures 3g and 3h) yielded a 1.8-fold lower k_1 than Pt (Table 1). Nitrite concentrations with carbon cloth reached a maximum 274 of 1.8 mg L⁻¹ NO₂⁻-N at 120 min and decreased to 0.73 mg L⁻¹ NO₂⁻-N at 360 min. Overall, carbon 275 276 felt performed better (higher k_I values) than Pt, carbon cloth, and most of the transition metals. One 277 factor that may explain the superior performance of carbon felt is that its high porosity and large 278 pores formed by interlace fibers allow the electrolyte to flow through the felts, favoring the 279 reactants transport and reducing the cell resistance[68-70].

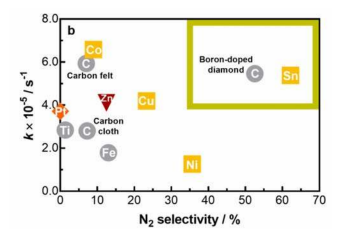
BDD electrodes achieved the most favorable outcomes among the carbon-based electrodes. Most notably, BDD achieved the highest selectivity towards desired gaseous N by-products $(S_{N_{gas}}=63\%)$ while also degrading nitrate 1.5-fold faster than the Pt electrodes (Figure 3i and Table 1). Due to its high stability under aggressive high acidic/alkaline media[71], corrosion resistance, low electrocatalytic inhibition[72], and belonging to the group of elements most available, BDD is a promising electrode for the electrocatalytic reduction of nitrate (ERN).

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287 **3.3.** Benchmarking electrocatalysts towards higher electrokinetics and selectivity

Electrochemical reduction applicability is defined by treatment kinetics and selectivity (i.e., preferential formation of N₂). Technology competitiveness in market settings will be defined by techno-economic aspects related to material cost and availability, which goes beyond treatment performance. While market costs are good indicators of electrode costs for some elements (e.g., Pt) (see Fig. S4), the actual capital costs for low-cost elements may be further influenced by electrode synthesis processes and net manufacturing costs. For example, the three carbon-based electrodes 294 demonstrated that even for a single earth-abundant element (i.e., carbon), the electrode morphology, 295 element oxidation state, and element bonding configuration played a major factor in overall 296 electrode performance. Manipulating the element's morphological and chemical properties 297 influence the synthesis or manufacturing costs of electrodes. This is true for most of the elements 298 studied herein. For example, while elemental Ti plate was used here, other researchers observed that 299 Magneli phase of titanium oxide (e.g., Ti₄O₇) can also degrade nitrate[73]. The numerous oxidation 300 states of many of the earth-abundant elements, compared with the almost exclusive reliance on elemental metals in application of PGE electrodes, offer a remarkable "design space" for use of 301 302 non-endangered elements as electrode materials.

Figure 4 benchmarks each electrocatalyst against Pt (\blacklozenge) in terms of kinetics and selectivity obtained at 20% of nitrate removal, while considering sustainability aspects of different elements. Note that several electrocatalysts (Co, Cu, Zn, carbon felt, BDD, and Sn) outperformed Pt for NO₃⁻ reduction kinetics, which is illustrated by their location in the upper half of Figure 4b.



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Figure 4. Pseudo-first order kinetics obtained vs. selectivity towards N₂ during electrochemical reduction of 100 mg L⁻¹ NO₃⁻-N in 50 mM Na₂SO₄ at 20 mA cm⁻² (values set at 20% of initial NO₃⁻ abatement). The electrocatalysts included in the upper right quadrant present the fastest reduction kinetics and highest selectivity towards nitrogen gas. Elements earth availability color coded based upon Figure 2: (•) high abundance, (•) limited availability, (•) rising threat, (•) serious threat in next 100 years. For water treatment, the viability of electrodes fabricated from different elements depends strongly on by-product selectivity towards innocuous nitrogen gas rather than selectivity toward nitrite or ammonia. In Figure 5a, the average of selectivity towards nitrogen is plotted with error bars showing that this parameter varies minimally across the duration of the electrochemical treatment for most of the elements studied. In fact, this behavior can be verified in Figure 5b, with BDD and Sn attaining the highest S_{Naas} .

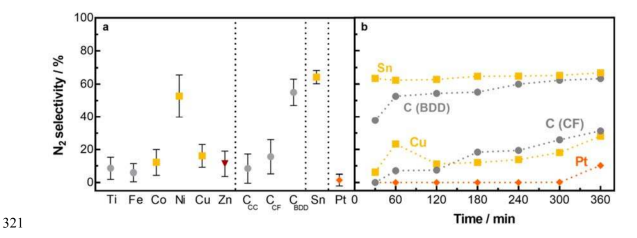


Figure 5. (a) Average and standard deviation of the selectivity towards N₂ obtained with the different cathodic materials studied during the ERN treatment. (b) Selectivity towards N₂ over time for selected electrodes. Note: CC, CF, and BDD stands for carbon cloth, carbon felt, and boron-doped diamond, respectively. Conditions: 100 mg L⁻¹ NO₃⁻-N in 50 mM Na₂SO₄ at 20 mA cm⁻². Elements availability: (•) high abundance, (•) limited availability, (•) rising threat, (\checkmark) serious threat in next 100 years.

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Effective electrocatalysts should have the attributes of rapid nitrate removal (i.e., high k_1) with a corresponding high selectivity towards desired N-gas by-products (i.e., high $S_{N_{gas}}$). For electrode materials illustrated in Figure 4, the more effective materials are located in the upper-right quadrant. Thus, Sn and carbon-based BDD would be classified as the most effective for nitrate

removal and are clearly superior to commercial polycrystalline Pt electrodes, which achieved only
1% selectivity towards N-gases.

335 Other material factors can impact electrocatalyst effectiveness for use in drinking water. 336 Most notably, corrosion of some electrodes may impart regulatory concerns or long-term 337 operability challenges. Among the metals investigated, only Cu is included in primary drinking water regulations, where levels above 0.8 mg L⁻¹ begin to pose health concerns. Such high levels 338 would be detrimental to the electrode itself and would therefore need to be studied in more detail. 339 340 Fe and Zn have similarly high secondary drinking water limits, set by aesthetics rather than health 341 limits, of 0.3 and 5 mg L⁻¹, respectively. Other metals (Ti, Co, Ni, Sn) are not regulated in drinking 342 water. While carbon is not directly regulated, numerous organic carbon compounds (e.g., 343 polyaromatic hydrocarbons) are regulated as carcinogens, but their evolution from graphitic or 344 diamond forms of carbon are unlikely. Thus, considering drinking water contaminant limits 345 wouldn't change the above recommendation to focus on BDD- or Sn-based electrodes.

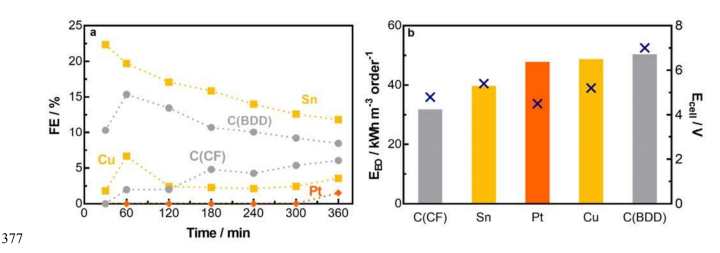
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3.4 Energy-based figures of merit considerations

348 Beyond performance (i.e., nitrate removal rates and by-product formation), energy 349 efficiency will also contribute to electrode element selection. Electrochemical reduction 350 effectiveness is commonly evaluated in terms of Faradaic efficiency (FE, Eq. (3)) for N₂ evolution. 351 Figure 6a shows FE over time for selected electrocatalysts and demonstrates competitive 352 alternatives to conventional Pt electrodes in terms of superior kinetics and selectivity. 353 Electrocatalysts with low selectivity for N_2 (i.e., Pt, Cu, carbon felt) accordingly had low FE, 354 ranging between 0% and 7%. Meanwhile, promising BDD and Sn electrodes had FEs between 10 and 22%. Both BDD and Sn showed an analogous decrease in FE over electrolysis time. This 355 356 behavior can be related to the nitrate concentration decrease over time, whereas the electric current 357 is the same. Consequently, the kinetics of parasitic reactions, such as hydrogen evolution, may be 358 enhanced[72,74], resulting in a lower final FE.

Engineering figures of merit provide a different, practical method to assess process energy 359 360 efficiency across multiple options. Because the WHO recommends $\sim 11 \text{ mg L}^{-1}$ as NO₃-N and our experiments had initial nitrate concentrations of 100 mg NO_3 -N, the E_{EO} is shown for one order of 361 362 magnitude decrease in nitrate (kWh m⁻³ order⁻¹, Figure 6b). Under identical electrochemical cell reactor configurations, E_{EO} ranged between 30 and 50 kWh m⁻³ order⁻¹. For the two most effective 363 364 materials from Figure 4 (i.e., Sn and BDD), E_{EO} was 39 and 49 kWh m⁻³ order⁻¹, respectively. E_{EO} at a given applied current is controlled by two main factors: (a) nitrate reduction rate k_1 and (b) the cell 365 366 potential (E_{cell}). Although Sn and BDD had a comparable kinetic constant for nitrate removal, the 367 cell potential average (E_{cell}) for each is different. The E_{cell} was 5.4 V for Sn and 7.0 V for BDD, which resulted in a higher E_{EO} for the BDD cathode. The interdependence between k_1 and E_{cell} 368 369 explain why the E_{EO} for Pt (47 kWh m⁻³ order⁻¹) is between the ones for Sn and BDD. Other 370 research shows that E_{EO} can be decreased by >10-fold from the observed values through engineering design of electrode morphology, electrode spacing, and hydrodynamics[22,23,73]. For 371 372 example, flow-through electrodes are superior to flow-by electrodes because the former exhibits 373 lower mass transfer limitations for target pollutants/by-products towards/from the electrode surface. 374 Fortunately, many of the elements, including BDD and Sn, can be engineered into almost any 375 integrated electrode and reactor configuration.



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Figure 6. (a) Faradaic efficiency (FE) over time, (b) energy consumption per order (bars) and cell potential average (×) for the electroreduction of 100 mg L⁻¹ NO₃⁻-N in 50 mM Na₂SO₄ at 20 mA cm⁻², using Cu, C-CF, C-BDD, Sn, and Pt as cathode materials. Note: CF and BDD stand for carbon felt and boron-doped diamond, respectively. Elements availability: (\bullet) high abundance, (\blacksquare) limited availability, (\diamond) rising threat, (\bigtriangledown) serious threat in next 100 years.

383

384 4. Conclusions

385 This work applies a framework for screening earth-abundant electrode materials under 386 identical operating conditions to obtain figures of merit suitable to identify more sustainable 387 electrocatalytic systems that can remove nitrate from drinking waters. Platinum (Pt), a well-studied 388 electrocatyst in the literature, was selected as a benchmark electrode representing the PGEs. 389 Electrokinetic pseudo first-order nitrate degradation rates (k_1) were slightly better (~50% larger k_1) 390 for Cu-, C-, and Sn-based electrodes relative to Pt. The difference in selectivity towards the 391 preferential by-products (i.e., N-gases) showed tremendous promise for more earth-abundant 392 elements, being up to 7-fold more selective (S_{Naas} of 55% and 64% for BDD and Sn, respectively) 393 than the PGE-based electrode ($S_{N_{aas}} \sim 1\%$ for Pt). Degradation rates and selectivities impacted 394 Faradaic efficiency and E_{EO}. No adverse effects were shown on these figures of merit when 395 transitioning electrode material from PGE to more sustainable elements (e.g., Sn, BDD).

Each element used in the electrodes has a commodity market cost ($\$ kg^{-1}$). The structure of catalytic materials, with crystallites of different sizes and orientations, are critical to maximize the selectivity towards the preferred by-product species (N₂ gas). Electrode manufacturing to transform the base element into the plethora of electrode architectures (e.g., high porosity electrodes) will add to the cost of the final electrode. For example, while carbon is the cheapest commodity element, the three carbon-based electrodes showed that different morphologies, oxidation states, and crystallites can lead to very different reaction outcomes. Thus, while acknowledging it may be premature to draw a specific selection of electrodes in terms of performance-cost analysis, this work provides an
initial roadmap on expected cost depending on the element source. A complete techno-economic
analysis will be conducted in a well-developed electrocatalytic system considering electrode life
and its manufacturing associated costs.

407 The above framework identified the application of earth-abundant materials, which global 408 availability is not endangered, to electrochemical nitrate reduction. This represents a game changer for developing low-cost electrocatalytic-based treatment water systems, where Pt-based catalysts 409 410 have been shown to be cost prohibitive (e.g., [40,41]). Sn-based and carbonaceous electrodes (i.e., 411 BDD) were recognized as viable earth-abundant electrocatalysts alternatives to endangered 412 elements. Innovations in electrode manufacturing are still needed to reduce capital costs and E_{EO} by 413 optimizing mass transport. This may occur by maximizing surface area of the more reactive and stable surfaces of earth-abundant elements as electrodes. Future research using nanotechnology with 414 415 earth-abundant materials can exploit opportunities [75], enabling catalytic sites that specifically 416 tailor nitrate reduction towards innocuous nitrogen gas, further increasing the competitiveness of 417 POU systems.

418

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