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# Synthesis of material libraries using gas diffusion electrodes†

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The future of energy relies on the advent of electrochemical energy production and storage. A key enabling factor is the effective synthesis of active materials, which, due to the global environmental circumstances and the growing demand, we must ensure are made sustainably. Thus, we unveil a rapid, sustainable, and scalable electrosynthesis route for a whole range of nanocrystalline materials with bright prospects for batteries, solar fuels and fuel cells, among others. For the proof of concept of the synthesis method, gas-diffusion electrocrystallization (GDEx), we synthesize manganese and cobalt oxides and hydroxides: Co-doped Na-birnessite, cubic/tetragonal spinels and layered double hydroxides (CoMn-LDH), owing to their current relevance. An oxygen depolarizing gas-diffusion electrode is used to fuel the oxidative synthesis at the electrochemical interface. Aside from the necessary metal precursors, all reagents are produced in situ with high efficiency. To elucidate the synthesis mechanism, a broad range of materials were produced under the same conditions. By changing the Co and Mn concentrations in the feed solution, the composition (Co/Mn stoichiometry), morphology (spinels vs. nanosheets), structure (tetragonal/cubic-spinel, birnessite, LDH), particle size (15-35 nm), crystallinity (polycrystalline particles vs. single-crystals), and phase purity were precisely tailored. A comprehensive library of nanostructures was built, wherein some materials exhibited outstanding catalytic properties for the oxygen evolution reaction, illustrating the significance of our strategy. To showcase the versatility of the method, we also prove the feasibility for sodium intercalation capacity in the materials, applicable to batteries. This work opens the door to a new systematic way of producing optimized and affordable materials for electrochemical energy applications and beyond.

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#### Introduction

The development of next-generation electrochemical energy technologies plays a key role in the future of sustainable energy. To meet the increasing demand for high-performance energy devices, how we fabricate electrochemically active materials must be transformed. First, we need methods that can develop into potent tools for high-throughput synthesis and screening,

especially of nanostructures with outstanding performance.¹ Today, this mostly implies trial and error strategies, making the development rather sluggish. Second, we are accountable for developing and using sustainable, environmentally innocuous, production methods. Ideally, these should be chemically and energetically thrifty, operate under mild conditions, avoid the use of hazardous chemicals, and not generate waste materials.

Metal-air batteries, solar fuels, and fuel cells are among the technologies driving the study and development of better electrocatalysts. The oxygen reduction (ORR) and evolution (OER) reactions<sup>2-4</sup> are by far the most important transformations in this regard. The optimization of electrochemically active materials in terms of structure and composition is difficult, given the broad range of possibilities. Varying synthesis conditions yield different particle sizes, morphologies, crystallinity, and ultimately, affordability.<sup>2</sup> Due to much lower costs and ready availability, manganese-containing mixed oxides and hydroxides have emerged as alternatives to noble metal catalysts, with certain materials matching or surpassing the performance of IrO<sub>2</sub> (OER) and Pt/C (ORR).<sup>5,6</sup> Thus, we selected such materials to characterize our proof of concept, although our approach is

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far-reaching and can be contrasted with landmark strategies to synthesize nanomaterial libraries.¹

One type of manganese dioxide,  $\delta$ -MnO<sub>2</sub>, forms birnessite and birnessite-type compounds. These are layered oxides, composed of Mn–O octahedral plates that intercalate water. The redox activity of Mn allows some of the metal in the oxide layers to deviate from a 4+ to a 3+ valence, as electroneutrality is conserved by intercalating small cations, usually Mg<sup>2+</sup>, Li<sup>+</sup>, Na<sup>+</sup>, K<sup>+</sup>, *etc.*<sup>7</sup> This makes birnessite-type MnO<sub>2</sub> a natural candidate cathode material for alkali and alkali-earth batteries, supercapacitors, and electrocatalysis (ORR/OER).<sup>8</sup>

Another family of materials that has jumped to the forefront of research is that of layered double hydroxides (LDHs). Their potential as electrode materials arises from their high redox activity, ion intercalation capacity, and high specific capacitance. The design of LDHs, in terms of both the intercalated ion and the metal hydroxide composition, results in the targeting of specific properties, which, if thoroughly controlled, confer them unique synthetic identities. This is an area to be exploited to fully realize the potential of these materials.

Both birnessites and LDHs are excellent precursors for the synthesis of spinels, since they allow for the even distribution of the cations. This a useful feature for synthesizing complex stoichiometries. An uneven distribution of metals is a routine problem in solid-state synthesis, which is the usual method employed for the production of complex spinels. Regardless of the precursors, spinels of the general formula  $\mathrm{Mn}_{3-x}\mathrm{Co}_x\mathrm{O}_4$  are found in numerous applications, including batteries, fuel cells, and (electro)catalysis. <sup>10,11</sup>

Various state of the art Mn/Co-based materials have been synthesized by chemical precipitation,11 and hydrothermal12 and sol-gel methods,13 as efforts are made to steer away from the highly unsustainable solid-state techniques. Particularly in the case of spinels, which are traditionally synthesized at very high temperatures, these approaches yield particles which are crystalline, but often large and irregular.14 Efforts have been recently made to develop synthesis routes at room temperature for the production of spinels, 10 birnessites, 15 and layered double hydroxides (LDHs),16 due to the more sustainable implications. A common thread between these synthesis techniques is the chemical addition of different Co/Mn precursors, oxidants, or reductants, to achieve the desired structure. However, the large spread on the electrochemical performance of these materials reported in the literature emphasizes the lack of control and reproducibility when synthesizing them, especially in terms of their structure.17

Composition, crystallinity, morphology, and more features that define a synthetic identity, are key factors in establishing structure–property relationships for any given structural group of Mn/Co oxides<sup>18</sup>,—as well as for any other nanomaterial. These are closely related to the synthesis process; hence a rational comparison of the materials reported across the literature, resulting from different techniques, is difficult.

Ideally, large ranges of comparable materials with variable structures and compositions, should be synthesized rapidly and under the same operational conditions (*e.g.*, synthesis route, flow, temperature, potential, chemical environment, *etc.*). Only

in this way, reliable benchmarking of catalytic performances, conceptualization of mechanisms, and improvements towards the production of affordable and optimized materials, can be achieved. This systematic approach to the synthesis of a variety of materials with only slight changes to the synthesis conditions represents a valuable platform for future screening of materials. A demonstration of such an approach is provided here, on top of being a sustainable, rapid, and up-scalable alternative.

# **Experimental section**

#### Synthesis by gas-diffusion electrocrystallization (GDEx)

The synthesis was made in an EC Micro Flow Cell (ElectroCell, MFC30009), using a three-electrode configuration, wherein a dimensionally stable anode (DSA) functioned as the counter electrode.

It is to be noted that our choice for the counter electrode material or reaction had no relevance on the process or findings here reported, as only the cathodic half-cell is examined. The electrochemical cell schematic is shown in Fig. 1. A PTFEbonded cold-rolled gas diffusion electrode (VITO CORE®) was used as the cathode; a metal frame in the cathode compartment acted as the current collector. A Zirfon® separator was used between the two electrolyte compartments. An Ag/AgCl reference electrode was placed in the catholyte chamber in close proximity to the gas-diffusion electrode (GDE) surface. The supporting electrolyte solution in all cases was composed of 0.5 M NaCl and sufficient HCl to set a pH of 3.5. The latter was added to avoid hydrolysis of the metal ions. The different solutions were prepared with anhydrous MnCl<sub>2</sub> and CoCl<sub>2</sub> (Sigma Aldrich) with total metal concentrations ranging from 1 mM to 5.5 mM, as measured by Inductively Coupled Plasma Optical Emission Spectrometry (ICP-OES). Each concentration was prepared for the pure metals, and for mixtures of Mn and Co with varying ratios, considering the referred total metal concentrations. The mixtures prepared contained Mn/Co mole fractions from 0 to 1. Anolyte solutions were made in the same

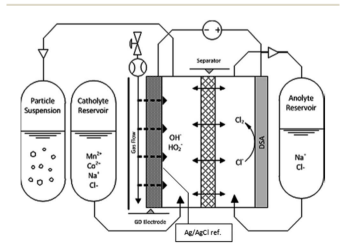


Fig. 1 Electrochemical reactor sketch, showing the primary components used for the synthesis of nanomaterial libraries by gas-diffusion electrocrystallization (GDEx).

way, except that no Mn or Co was added; only the supporting electrolyte and HCl for pH control were supplemented. Air was fed into the cell, through the hydrophobic backing of the GDE, at a flow rate of 1 L h<sup>-1</sup> at an overpressure of 18 mbar (g), set by a water column at the gas outlet of the cell. The electrolyte solutions consisted of 1 L for both anolyte and catholyte, respectively. The catholyte solution was fed from the catholyte external reservoir (Fig. 1) and was collected in a separate bottle at the outlet (Fig. 1, *i.e.*, particle suspension). The anolyte solution was recirculated to and from the same feed bottle (Fig. 1, *i.e.*, anolyte reservoir). Peristaltic pumps (Watson-Marlow) were used to provide a liquid flow rate of 20 mL min<sup>-1</sup>. A constant current of -300 mA was applied for each synthesis.

The electrodes had a projected surface area of 10 cm<sup>2</sup>, facing towards the corresponding flow channel with a thickness of 0.4 cm, and a total of 4 ml of volume in each chamber. Under these conditions the residence time for incoming Mn<sup>2+</sup>/Co<sup>2+</sup> ions in the cell was approximately 12 seconds.

Prior to each synthesis, the electrolyte with no transition metal was fed through the reactor in order to ensure sufficient wetting of the electrode and separator. The inlet of the reactor was then switched from the supporting electrolyte solution, to the catholyte reservoir to begin the synthesis. The chronopotentiometry (CP) was started and the outlet pH was monitored until it reached a steady value. To obtain continuous pH data from the very beginning of the experiments, 50 mL of electrolyte solution at pH 3.5 was added to the particle collection bottle. This ensured the availability of sufficient liquid to wet the pH probe. Once this was achieved, the catholyte outlet was switched to a new collection bottle.

After this startup process, the synthesis began. Once the experiment was finished (when the entire catholyte solution was consumed) the resulting colloidal suspension was centrifuged (Thermo Scientific Sorvall RC 6+), at 11 000 rpm for 15 minutes. The supernatant solution was removed, and the precipitate was washed with a NaOH solution with a pH set to that of the synthesis solution at the outlet, and mixed with a vortex. The new mixture was centrifuged again. This process was repeated 3 times to ensure that leftover NaCl from the electrolyte was removed from the particles. Finally, the supernatant was removed once again and the collected pellets were dried under a nitrogen atmosphere at room temperature, achieving dry powders.

#### **GDE** electrochemical studies

The oxygen reduction reaction (ORR) at the GDE was characterized with Cyclic Voltammetry (CV) and Electrochemical Impedance Spectroscopy (EIS). The supporting electrolyte solution (no transition metals) was used for characterization in the same electrochemical cell, with varying NaCl concentrations (0.25 M and 0.5 M) and varying gases (air  $\nu s.~O_2$ ). EIS measurements were taken at potentials between  $-0.3~V_{Ag/AgCl}$  and  $-0.5~V_{Ag/AgCl}$  on the ORR region of the previously taken CV, between 100 kHz and 10 mHz, at an amplitude of  $\sim 7~mV$  (rms). Prior to each EIS measurement, the system was verified for stability, causality, linearity and finiteness.

#### Structural characterization

The dry samples were analyzed by powder X-ray diffraction (XRD) in a PanAlytical X'Pert Pro diffractometer (Cu Kα radiation,  $\lambda = 1.5406$  Å). Samples were crushed in a mortar and placed in standard silicon monocrystal sample holders. Measurements were performed with a spinner at 40 mA to 40 kV spending 4 s per step, with a step size of  $0.04^{\circ}$   $2\theta$  in the  $10-110^{\circ}$  $2\theta$  range. Rietveld refinements were performed in all samples to fit the profiles and extract the lattice parameters from the data using HighScore Plus software. Crystallite sizes were calculated using the Scherrer equation,  $\tau = \kappa \lambda/(\beta \cos \theta)$ , with crystallite size  $(\tau)$ , shape factor (K), line broadening factor  $(\beta)$ , X-ray wavelength  $(\lambda)$  and Bragg angle  $(\theta)$ . Micrographs of the dry samples were taken with no further treatment, with an FEI Nova NanoSEM 450. Images presented were taken with secondary electrons and an accelerating voltage of 5.00 kV. EDS spectrometry was used to confirm chemical compositions.

#### Chemical characterization

Upon finishing a synthesis the chemical composition of the outlet solution and the solid particles was investigated. 2 mL samples of the inlet and outlet solution were analyzed by Inductively Coupled Plasma Optical Emission Spectrometry (ICP-OES). The chemical composition of the inlet solution was confirmed to be the intended one (from 1 mM to 5.5 mM) and the outlet solution was checked to ensure complete removal of the metals. The dry particles were digested in aqua regia, diluted and then measured in the ICP-OES as well to observe the metal composition of the material and compare it with the inlet solution metal ratio.

#### Testing of electroactive materials

**OER electrocatalysis.** Electrocatalytic inks for coating the glassy carbon electrode were prepared by dispersing the powders in a mixture of 8 mL of ethanol, 12 mL of DI water, and 0.8 mL of a 3 wt% Nafion solution to a final powder concentration of 0.5 mg mL $^{-1}$ . The inks were placed in an ultrasonic bath for 30 minutes, prior to use. Glassy carbon (GC) disk electrodes were used, with 3 mm diameter, encased in polyether ketone (PEEK) rods with gold contacts. 20  $\mu$ L of the inks were drop-cast on the tip of the rods (6 mm total diameter) and dried in an oven at 40 °C, for 1 hour. The coated electrodes were dried at room temperature (18  $\pm$  2 °C). The electrolyte solution consisted of 250 mL of 1 M KOH. The electrochemical setup consisted of the coated GC as the working electrode, carbon paper as the counter electrode, and Saturated Calomel (SCE) as the reference electrode.

IR drop was corrected with high-frequency impedance prior to all measurements. The electrodes were cycled 100 times at 10 mV s<sup>-1</sup>, between 0.3  $V_{\rm SCE}$  and 0.75  $V_{\rm SCE}$  for electrochemical conditioning. After cycling, linear sweep voltammetries (LSV) were performed at 5 mV s<sup>-1</sup> to collect the final data. Measurements were reproduced with samples synthesized separately, and each sample tested underwent the 100 cycle conditioning to achieve a stable value, followed by 3 LSVs at a lower scan rate (5 mV s<sup>-1</sup>) to confirm the values.

Active material for Na-ion battery cathodes. 355 mg of birnessite Na<sub>x</sub>Co<sub>v</sub>Mn<sub>1-v</sub>O<sub>2</sub> issued from the GDEx process was crushed and mixed with 101 mg of conductive carbon and 40 mg of binder. The slurry was doctor blade coated on an aluminum current collector to obtain a mass loading of 1.125 mg cm<sup>-2</sup>. 18 mm diameter, single-side coated electrodes were then tested in a three-electrode, sodium-ion battery half-cell using an EL-Cell (PAT-Cell) chamber. The aprotic mixed-electrolyte, 1 M NaPF<sub>6</sub> in EC: PC (1:1, w/w), and a separator, Freudenberg Viledon (FS-5P, 220 µm), were used. The threeelectrode set-up consisted of a sodium metal disk (18 mm diameter) and a sodium metal ring electrode embedded in the separator as the reference electrode.

# Results and discussion

#### Gas-diffusion electrocrystallization

GDEx is a rapid one-pot reaction crystallization process, electrochemically steered at porous gas-diffusion electrodes (Fig. 2a and b), facing liquid electrolyte and a gas, wherein a triplephase boundary is established (Fig. 2c). The multi-layered gasdiffusion electrodes include a hydrophobic outer layer through which the gas percolates, to ultimately be electrochemically reduced at a hydrophilic active layer (activated-carbon based), acting as a cathode. The active carbon powder constituting the gas-diffusion cathode is porous, including meso- and micropores, with an average pore diameter of 48 Å and a specific surface area of 745 m<sup>2</sup> g<sup>-1</sup>, resulting in a highly reactive electrode. The total porosity of the gas-diffusion cathode manufactured by cold-rolling is 67%, out of which 29% corresponds to hydrophilic porosity and 38% to hydrophobic porosity, facilitating the electrolyte and gas transport, respectively. The specific surface area of the manufactured electrodes is 454 m<sup>2</sup> g<sup>-1</sup>. A comprehensive description of these electrodes and the methods to characterize them can be found elsewhere.19

The synthesis process proceeded in one step: a feed solution of MnCl<sub>2</sub> and CoCl<sub>2</sub> as the metal precursors, and 0.5 M NaCl as the supporting electrolyte, was fed through a flow cell equipped with a gas diffusion electrode (GDE). The formation of nanostructures by GDEx happens through the following steps. First, an oxidant gas  $(e.g., O_2 \text{ in air})$  is forced through the hydrophobic backing on the electrically polarized porous cathode (Fig. 2a). After it diffuses to the electrocatalytic surface (e.g., hydrophilic activated carbon), the gas is reduced. For instance, by imposing specific cathodic polarization conditions (e.g., -0.145 V<sub>SHE</sub>), O<sub>2</sub> is electro-reduced (Fig. 2c), to H<sub>2</sub>O<sub>2</sub> in a 2 electron (2 e<sup>-</sup>) transfer process and H<sub>2</sub>O in a 4 electron (4 e<sup>-</sup>) transfer process, as per the following reactions:

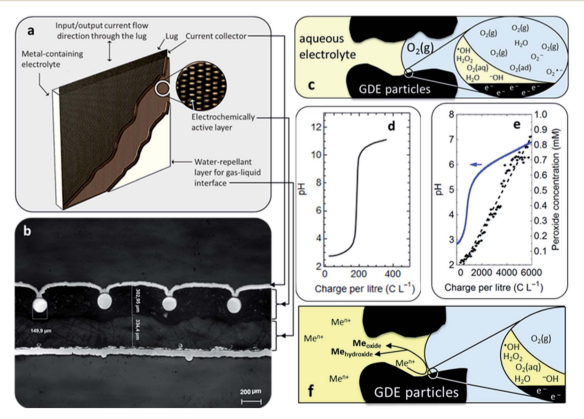


Fig. 2 Gas-diffusion electrode (GDE). (a) Schematic representation showing the different components of a GDE. (b) Micrograph of a VITO CORE® GDE taken by Scanning Electron Microscopy (SEM). (c) Representation of the triple-phase boundary at a GDE, illustrating the oxygen reduction reaction (ORR), an aqueous electrolyte and activated carbon as the electrocatalytic material. High mass fluxes are achieved at the highlighted meniscus region. (d and e) pH and  $H_2O_2$  evolution, respectively, as a function of the charge consumed in a typical gas-diffusion electrocrystallization process starting at pH 2.7, wherein metal ion precursors are absent in the electrolyte bulk (i.e., oxygen reduction reaction at the gas-diffusion electrode). (f) The addition of metal ions to the electrolyte results in oxide and hydroxide formation.

$$O_{2(g)} + 2e^- + 2H_{(aq)}^+ \rightarrow H_2O_{2(aq)}$$
 (1)

$$O_{2(g)} + 2e^{-} + 2H_{2}O_{(l)} \rightarrow HO_{2(aq)}^{-} + OH_{(aq)}^{-}$$
 (2)

Some polyatomic ions, peroxides, and free radicals are transient intermediaries formed together with OH ions (Fig. 2c), as per the mechanism established for O<sub>2</sub> reduction at non-catalyzed carbon electrodes.20 As this happens, abrupt local-pH and local-electrolyte potential changes arise rapidly within the cathode porosity. For instance, based on our experiments it is estimated (from the electric charge consumed), that the profuse amount of OH<sup>-</sup> rapidly produced (e.g., >1300 mol m<sup>-3</sup>), immediately results in a pH as high as 13-14 within the porosity of the cathode. 21 As the hydroxyl ions spread to the bulk electrolyte, systematic pH increases become consistently manifest in the electrolyte bulk (Fig. 2d). In due course, lower amounts of H<sub>2</sub>O<sub>2</sub> are generated (Fig. 1e), e.g. <1-5 mM, depending on the operational conditions employed and the amount of charge transferred per unit volume of catholyte. At the steady state, it is expected that a reaction front is fully developed throughout the hydrodynamic diffusion layer. This sets precedent for local saturation conditions at the electrochemical interface.

Throughout this work, both  $H_2O_2$  and  $HO_2^-$ , will be referred to as  $HO_2^-$ . First, the analytical method employed serves for the indistinguishable quantification of both chemical species. Second, we grant that under the pH conditions prevalent at the steady-state in the bulk of most of the catholyte solutions investigated,  $H_2O_2$  will be the more abundant chemical species (p $K_a = 11.65$ ).<sup>22</sup> However, at the electrochemical interface where the oxygen reduction reaction occurs, a profuse amount of OH-prevails, resulting in a local pH as high as  $14.^{21}$  Thus, the conditions for  $HO_2^-$  to be stable are met in the vicinity of the gas-diffusion cathode and throughout the diffusion layer. It is, in fact, this phenomenon that facilitates the onset of hydroxide supersaturation<sup>23</sup> and hence of the reactive precipitation (*i.e.*, crystallization) of the different nanostructures, at the electrochemical interface (Fig. 2f).

The synthesis of nanostructures by GDEx provides very specific supersaturation conditions: (1) short nucleation and growth periods, (2) at high rates, and (3) avoiding the immediate agglomeration of the nanostructures formed.21 The flowing conditions impose only a temporary contact of metal ion precursors with the reactive species at the saturated electrochemical interface—rendering transient nucleation conditions. The possibilities of growth are feeble, as encountering other metal ions is restricted by their high dilution. The rate of interfacial processes typically scales with the available surface area (i.e., surface active sites), which is provided by the highly porous electrode, facilitating a high rate of production of HO<sub>2</sub>, OH, and their radicals, which quickly react with the metal ion precursors available at the interface. The reaction performance of a gas-diffusion electrode depends on the local environment within the porous active layer and the balance between transport phenomena and reaction kinetics.<sup>22</sup> Specifically, the rate of the gas-reduction reaction depends on the concentration of dissolved gaseous species. In gas-diffusion electrodes, the pores in the active layer are wetted by a thin film of electrolyte, which minimizes the transport resistance of the gas into the electrocatalytic sites while maintaining a good ionic conductivity. Due to these phenomena, the current density in gas-diffusion electrodes can be two orders of magnitude of that of planar electrodes<sup>24</sup> enabling fast reactivity at the electrochemical interface. Lastly, in analogy to growth, aggregation is restricted, as the nanoparticles dispersed in the aqueous solution are few, and then collisions between them are not likely. Otherwise, the particles can be easily transferred to pure water (i.e., using dialysis, or by centrifugation and resuspension), where they can set apart even better, as the repulsive forces become stronger (i.e., linked to double layer expansion as a result of the lower ionic strength—and sometimes pH—vs. those of the primary synthesis medium). The GDEx process is highly reproducible, and it involves mild synthetic conditions (e.g., 291 K and atmospheric pressure).

#### Particle synthesis

Currents were constant at  $-30~\text{mA}~\text{cm}^{-2}$  for all experiments. The materials synthesized were collected at the outlet of the catholyte chamber, as particle suspensions. Fig. 1 shows a schematic of the synthesis setup, highlighting the GDE and the reacting species from the gas phase, as well as the formation of the crystals in the bulk electrolyte. A representative chronopotentiometry is shown in Fig. 3 (bottom). Hydroxide production resulting from the ORR caused an increase in pH. Fig. 3 (top) shows the pH profiles with time (charge) for solutions containing 0 mM, 1 mM, 2 mM, 3 mM, 4 mM and 5 mM of MnCl<sub>2</sub>. All samples and measurements were taken at steady-state, after stable conditions were achieved (<15 minutes).

The current (-30 mA cm<sup>-2</sup>) reduced the incoming molecular oxygen. The first reduction step of the two electron reduction pathway is shown in reactions (1) and (2) for acid and alkaline media respectively.<sup>25</sup> The peroxide ion can be further reduced at the electrode to hydroxide, or, in the presence of Mn<sup>2+</sup> can react to oxidize the metal (reaction (3)).

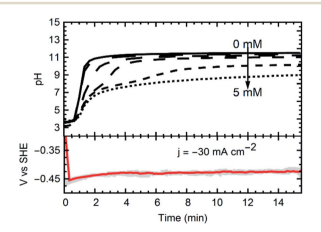


Fig. 3 Top: pH profiles from 5 experiments with increasing initial metal concentration. From top to bottom: 1 mM, 2 mM, 3 mM, 4 mM, and 5 mM. Bottom: Potential of the gas diffusion electrode upon starting the synthesis, with an applied current density of -30 A m $^{-2}$ .

$$Mn_{(aq)}^{2+} + OH_{(aq)}^{-} + HO_{2(aq)}^{-} \rightarrow MnO_{2(s)} + H_2O_{(l)} \tag{3} \label{eq:3}$$

The oxygen reduction reaction fueling the formation of the products was studied on the GDE. Fig. 4 shows the cyclic voltammetry (CV) and impedance (EIS) results registered under the steady-state polarization conditions (time > 30 min). The ORR reaction was studied from a potential of  $-0.3 \text{ V}_{A\sigma/A\sigma\text{Cl}}$  to -0.5

The impedance spectra show two Gerischer-like semicircles<sup>26</sup> with a high-frequency linear segment. The high-frequency semicircle is independent of the applied potential (Fig. 3a), oxygen supply (Fig. 3b), or electrolyte resistance (Fig. 3c). At frequencies between 10 kHz and 100 Hz, this is attributed to the electrode geometry, a behavior characteristic of semi-infinite pores. 26,27 High-frequency signals may fail to penetrate the entire depth of the pores, leading to the straight line in the first part of the spectra. The low frequency semicircle exhibits similar characteristics, but with impedance magnitudes depending on all the mentioned parameters. Increasingly negative potentials, higher oxygen supply, and higher electrolyte

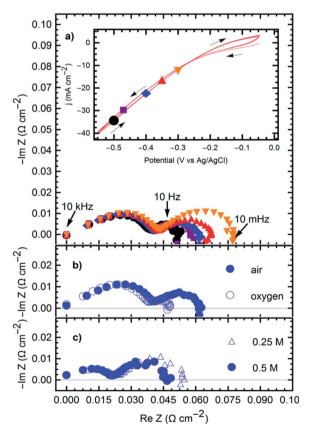


Fig. 4 Complex plane plots for the gas diffusion electrode in NaCl solutions at an influx pH of 3.5 with varying potential, gas choice and electrolyte concentration. (a) Effect of applied potential: 0.5 M NaCl, air, applied potentials as shown by the corresponding markers in the voltammetry. Inset: cyclic voltammetry taken at 10 mV s $^{-1}$ . (b) Effect of oxygen concentration: 0.5 M NaCl, E = -0.47 V, air (filled circles) and oxygen (empty circles) used in the gas chamber. (c) Effect of concentration of conducting electrolyte (NaCl): air, E = -0.47 V, 0.5 M NaCl (filled circles) and 0.25 M NaCl (empty triangles).

concentration, all reduce the overall magnitude of the impedance. Similar behavior is seen in other triple-phase boundary reactions in gas diffusion electrodes,28 suggesting a mechanism that depends on the diffusion through deep electrode pores and adsorption of the reactive species (oxygen) before charge transfer occurs and the reaction products ((1) and (2)) diffuse out of the electrode to start the precipitation process (reaction 3). Indeed, Gerischer-like elements relate to chemical reactions in the bulk electrolyte preceding or following a finite diffusion phenomenon associated with an electrochemical reaction.

Thus, GDEx is defined as the reactive precipitation of intermediaries issued from the ORR at the GDE, with the metal precursors provided. Notably, the synthesis of nanostructures (as explained in the subsequent sections) is fast under the operational conditions here employed, with rates approaching  $\sim$ 2.6 to 7 mg min<sup>-1</sup>. This implies  $\sim$ 3.7 to 10 g per day using a single, inexpensive, electrochemical milireactor under flow regime, which can be scaled-up by replication. The process is highly reproducible, and it involves mild synthetic conditions (e.g., room temperature and atmospheric pressure). Also, the process entails low energy consumption, e.g.,  $\sim$ 2 to 6 mW per h per mg of nanomaterials formed. Finally, it is efficient, i.e., 50% current efficiency for the formation of the reactive intermediaries that fully react with the metal precursors to achieve the targeted nanostructures. As it operates in a flow-cell configuration, it is easily up-scalable by stacking multiple individual cells, without losing in performance. GDEx is a sustainable synthesis method for nanostructures because it uses the cleanest possible reagent, the electron, to fabricate in situ the chemicals needed to react with the metal precursors. It could be optimized for zero-waste generation. It is highly versatile, as a variety of nanostructures are feasible,29 besides providing an elegant and simple way to control many essential nanostructure attributes in the course of the synthesis. Albeit its early stage of development and broad possibilities for optimization, GDEx emerges as a ground-breaking synthesis method for comprehensive libraries of diverse nanostructures.

#### Charge driven structure control

Control over the synthesized materials is easily exerted via the amount of charge applied and changes in the precursor solution. The ratio of charge applied to total metal concentration,  $R_O$ = Q/[M], as well as the mole fraction of Co,  $X_{Co} = [Co]/([Co] +$ [Mn]), determines unique combinations of structure and composition of the products. Fig. 5 shows the structural transitions upon changing  $R_Q$  and  $X_{Co}$ . As the potential, and thus the charge applied, are constant in all the experiments presented, the rate of production of reactive oxygen species (ROS) is the same for all cases. These species (peroxide, hydroperoxyl, etc.) oxidize the metals from 2+ to the target average oxidation state required for each material.  $R_O$  is then controlled by changing the inlet metal precursor concentration, which leads to the desired oxidation state in the products. A higher ratio results in higher average valence of the synthesized materials, with common average oxidation states of the metals: CoMn-LDHs (LDH) +2.3 (reaction (6)), Bir-MnO<sub>2</sub> (BIR) +3.7 (reaction

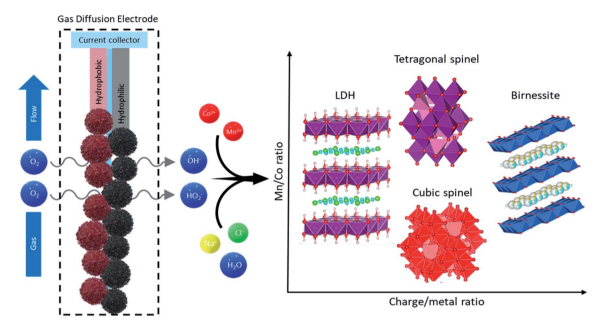


Fig. 5 GDEx system sketch. The main components of the GDE are shown and described, with the gas flow and diffusion path through the electrode leading into the metal containing electrolyte. The result of the synthesis is summarized with the structural control over the resulting structure *via* changes in the charge/metal ratio depicted on the chart.

(4)), and spinels (cubic CSPIN and tetragonal TSPIN) +2.66 (reaction (5)). $^{30-32}$  The specific synthesis rates for these products are  $\sim$ 7.0 mg min $^{-1}$  for the LDHs,  $\sim$ 4.5 mg min $^{-1}$  for the SPINs, and 2.6 mg min $^{-1}$  for the BIRs, respectively. The layered materials result from both extremes of  $R_Q$ , as the layers can have a majority composition of either Mn $^{(IV)}$ O<sub>2</sub> or Mn $^{(II)}$ (OH)<sub>2</sub>, while intermediate oxidation states yield spinels. The proposed formation reactions, starting from the dioxide described by reaction (3), are summarized:

$$MnO_{2(s)} + dX_{(aq)}^{2+} + 2dNa_{(aq)}^{+} + 4dOH_{(aq)}^{-} \rightarrow Na_{2d}X_{d}MnO_{2.2d(s)} + 2dH_{2}O_{(l)}$$
 (4)

$$MnO_{2(s)} + 2X_{(aq)}^{2+} + 4OH_{(aq)}^{-} \rightarrow X_2MnO_{4(s)} + 2H_2O_{(l)}$$
 (5)

$$MnO_{2(s)} + dX_{(aq)}^{2+} + dOH_{(aq)}^{-} + dCl_{(aq)}^{-} + dH_2O_{(l)} \rightarrow X_{cl}Mn(OH)_{3c}Cl_{cl(s)}$$
 (6)

where X represents a divalent, size-compatible, cation such as  $Co^{2+}$  in this case.

A phase diagram (Fig. 6 left) was composed from experiments carried out with -30 mA cm $^{-2}$ . The phase diagram reflects the predominant structure (or mixture) formed by using the annotated concentration of Co and Mn in the inlet solution to the gas diffusion cell. These results are also dependent on the volumetric flow rate (residence time) and the current drawn. As such, in order to reduce the degrees of freedom, the flow rate and current were fixed. Then the two main variables become each metal concentration; choosing a combination of them results in one of the 4 structural groups specified (birnessite, cubic spinel, tetragonal spinel, layered double hydroxide) or a mixture of them. Each structure has a certain tolerance for

defects; once this threshold is overcome, a new phase is formed. This is reflected by the transition zones in the phase diagram with the mixture of colors. Once the metals (and their oxidation states) are in the right ratio for a new single phase product, they enter a new region of the phase diagram. So it becomes an important operational tool in order to predict the materials formed.

Mn-only containing solutions produced birnessites at concentrations lower than 1.5 mM, and single-phase spinels at [Mn] > 2.5 mM. Higher feed concentrations produced unstable Mn(OH)<sub>2</sub>, which quickly oxidized to the spinel, and resulted in leftover Mn<sup>2+</sup> ions in solution, due to insufficient charge for precipitation. Values between the boundaries resulted in mixed compounds with both phases present. The addition of cobalt introduced a new degree of freedom. Stoichiometry changes reflecting the inlet solution occurred until the structural distortions cause too much strain, leading to the formation of a new phase, hence the formation of the cubic spinels (CSPIN) and LDHs. At preponderant cobalt fractions,  $\beta$ -hydroxides were formed; not long after production, the structure partially shifted to the  $\alpha$ -variety, making the synthesis of single-phase  $\beta$ -Co(OH)<sub>2</sub> difficult.

Birnessite-type layered Mn dioxide is rarely formed in the absence of small intercalating cations. As such, the large excess of Na<sup>+</sup> (0.5 M) in the background electrolyte promoted its formation. In addition, the mildly oxidative nature of the synthesis conditions favored the birnessite structure as well, the MnO<sub>2</sub> layers are relatively under-oxidized and electrostatically compensated by the intercalated cations, in contrast to other Mn dioxides ( $\alpha$ -,  $\beta$ -, etc.) which include a higher Mn<sup>4+</sup> content. This resulted in Na-birnessite, Na<sub>x</sub>MnO<sub>2</sub>. On the opposite end of the spectrum, the excess of Cl<sup>-</sup> in solution allowed the

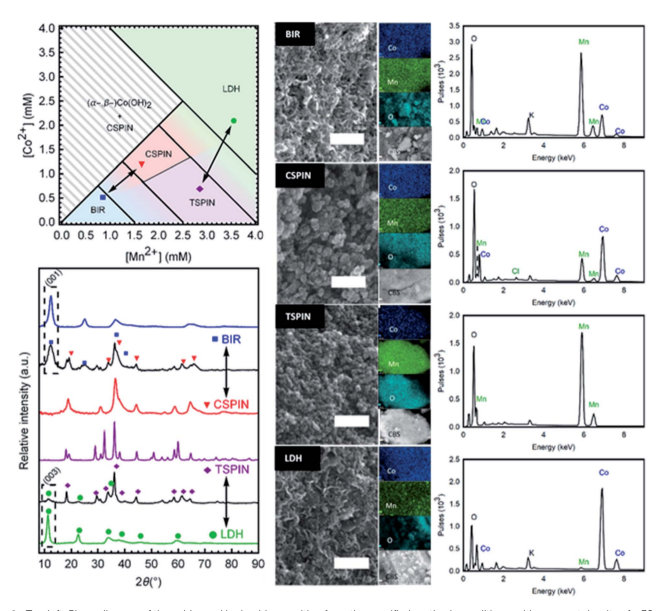


Fig. 6 Top left: Phase diagram of the oxides and hydroxides resulting from the specified synthesis conditions, with a current density of -30 mAcm<sup>-2</sup>. Each colored region is representative of the annotated phase, with transitions between resulting in phase mixtures. Bottom left: XRD traces of 6 samples. From top to bottom: single-phase BIR, mix of BIR with CSPIN (CBIR), single-phase CSPIN, single-phase TSPIN, mix of TSPIN with LDH (TLDH), single phase LDH. The markers show the most prominent peak positions of BIR, CSPIN, TSPIN and LDH phases respectively. Center: SEM pictures of the 4 structures, top to bottom: BIR, CSPIN, TSPIN, and LDH, scale bars are 200 nm. To the right of each image are the 3 EDS elemental mappings of Co, Mn, and O, and the backscatter electron image of the regions examined with EDS for each case. Right: The EDS spectra of each sample shown.

formation of the LDHs over single metal hydroxides when the  $R_O$  is low and the metals are mostly unoxidized. The  $\beta$ -Co(OH)<sub>2</sub> structure was stabilized in the presence of Mn<sup>3+</sup>, forming the Cl<sup>-</sup> LDH.

Single- and multi-phase materials were obtained by operating in the specified concentration regions. Diffractograms of each area in the phase diagram are presented in Fig. 6 (bottom left). Characteristic peaks are marked for each structure, to show the phase purity of BIR, CSPIN, TSPIN, and LDH materials, as well as the different phase mixtures of the phase transitions. All peaks can be indexed by the respective phase. The tetragonal spinel shown was refined to the I4<sub>1</sub>/amd space group.

Peak broadening is observed for the cubic spinel  $(Fd\bar{3}m)$ , attributed to the (nano-)crystallite size. The BIR and LDH traces are also shown to be phase pure. The shift to small angles in the reflections of the LDH, relative to the BIR, serves as an easy identifier of the hydrotalcite structure. Peak splitting around  $65^{\circ} 2\theta$ , seen as a broad hump in the BIR spectra, points to an orthorhombic structure, as opposed to the naturally occurring hexagonal birnessite. The peak broadness, present in the LDH pattern to a lesser degree, is a feature of turbostratic structures.34

Micrographs to examine the morphology and dimensions of samples from each structure are shown in Fig. 6 (center). Clear

distinctions can be made between spinel and layered materials. Spinels exhibit broadly jagged spheres of 15 nm to 35 nm in diameter. Meanwhile, the layered materials can be seen as delaminated sheets, characteristic of turbostratic birnessite,35 with face dimensions in the range of 50 nm. Brunauer-Emmett-Teller (BET) analysis for the specific surface area of the asprepared powders resulted in very large values, greater than 70 m<sup>2</sup> g<sup>-1</sup> for all materials synthesized. Particle sizes estimated from the surface area (approx. 30 nm in diameter) correspond with the range of sizes seen in the micrographs. The large surface area is a feature of the low temperature used to synthesize the materials, as traditional high-temperature processes yield larger, and more inactive, particles. 10,14 Next to each micrograph in Fig. 6 are EDS images showing the elemental composition of the samples, as well as the CBS (circular backscatter detector) image of the region analyzed from each material. The spectra with the approximate elemental ratios are shown in Fig. 5 (right). The composition from the EDS is in agreement with the metal composition of the solution fed to the cell. As the removal of the metals from solution nears 100%, as measured by ICP-OES, the chemical composition of the particles mirrors that of the initial solution. The removal data, as well as the EDS results, were corroborated by the ICP-OES analysis of the (acid digested) products. Table 1 shows the composition ranges associated with the structures shown.

### Structural transformations via doping

The Co/Mn ratio fed to the flow cell is reflected in the final product, as >99% of the metal ions are removed from solution when the total metal concentration is kept below 4.5 mM. For this reason, GDEx can also be regarded as an effective method to remove metal ions from solution (*i.e.*, for remediation or metal recovery purposes).

Cobalt doping can substitute Mn sites (such as for birnessites and LDHs, Fig. 7 left),<sup>7</sup> or it can change the structure as is the case with the spinels<sup>14</sup> (Fig. 7 right).

Fig. 7 shows the traces of the layered materials, from pure Na-birnessite to a hydrotalcite phase for the LDHs, and the β-Co(OH)<sub>2</sub>. All mixed-metal compounds are single phase. As previously mentioned, the Mn-only birnessite exhibits orthorhombic character. Co-doping in birnessites is shown here to decrease interlayer ordering (c-axis) as evidenced by the reduced intensity for (001) and (002) peaks (approx. 12° and 23° 2 $\theta$  respectively) but otherwise preserves the sheet structure. The peaks at larger angles (approx. 37° and 68° 2 $\theta$  respectively) with

no *c*-axis contribution show the conservation of the sheet integrity regardless of interlayer c-stacking. The reduced *d*-space ratio (Mn: 1.68 to MnCo: 1.72) between the (11,20) and (31,02) peaks suggests that Co-doping under these conditions shifts the structure from orthorhombic to hexagonal birnessite. Factorized At the same time, even with increased Co-doping the basal spacing was conserved around 7.3 Å. The addition of lower valent cations ( $\text{Co}^{2+}/\text{Co}^{3+}$  vs.  $\text{Mn}^{3+}/\text{Mn}^{4+}$ ) is balanced by the presence of Na<sup>+</sup> in the interlayer to electrostatically stabilize the structure, causing an expansion to accommodate the ions. The addition of more Co (above  $X_{\text{Co}} = 0.5$ ) resulted in mixed-phase compounds. Larger deviations in the average valence of the metals (Co/Mn) in the nanosheets, as well as the size constraints, disrupt the sheet ordering to the point where unstable delaminated sheets undergo transition into spinels.

The mixture of hydroxides presented in the bottom trace of Fig. 7 (left) is the result of a partial transformation occurring during the collection and cleaning process (post-synthesis) due to the instability of the  $\beta$ -hydroxide structure. The addition of Mn proved effective in stabilizing it, resulting in the layered double hydroxides. Relative to the birnessites, the expansion of the basal spacing continues with the formation of the hydroxides (Fig. 7). Layers of positively charged Mn/Co hydroxides,  $[(x^{2+})_{1-d}(Mn^{3+})_d(OH)_2]^{d+}$ , are balanced by  $Cl^-$  ions in the interlayer spacing. The (003) peak at 11.26°  $2\theta$  is used to calculate the basal thickness, here 7.9 Å on average. LDHs were produced and stabilized with  $X_{Co}$  between 0.50 and 0.90. Low amounts of Co produced a phase mixture with spinels. As such, Na-birnessites were produced with  $X_{Co} = 0$  to 0.5, LDHs with  $X_{Co} = 0.5$  to 0.9.

#### Tetragonal-cubic spinel transitions

Heavy cobalt loading (Co/(Co + Mn) > 0.4) results in cubic spinels (Fd3m). Increasing amounts of Mn (Co/(Co + Mn) < 0.4) creates lattice distortions due to the Jahn–Teller effect, resulting in a clear shift lowering the symmetry to the tetragonal structure ( $I4_1/amd$ ). There are two kinds of transitions seen in Fig. 7, characterized by the diffractograms and the lattice parameter. A sharp cubic–tetragonal transition is expected and seen at (Co/(Co + Mn) > 0.4) with the a/b/c lattice parameter of 8.3 Å splitting into a/b = 5.8 Å and c = 8.81 Å. Within each phase, gradual transition towards the inflection point in the lattice fitting is seen with varying Co/Mn ratio. The symmetry lowering is clearly observed from the splitting of the Fd3m peaks as the Co content decreases, accompanied by an expansion of the c-lattice parameter from 8.81 Å for Co<sub>0.8</sub>Mn<sub>2.2</sub>O<sub>4</sub> to 9.43 Å for pure Mn<sub>3</sub>O<sub>4</sub>. This shows remarkable control of the composition and

Table 1 Chemical composition of structural ranges and samples shown in Fig. 5

Structure abbreviation	Boundary compositions		
	Low Co	High Co	Sample shown in Fig. 5
BIR	$Na_xMnO_2$	$Na_xMn_{0.5}Co_{0.5}O_2$	$Na_xMn_{0.8}Co_{0.2}O_2$
TSPIN	$\mathrm{Mn_3O_4}$	$Co_{1.1}Mn_{1.9}O_4$	$\mathrm{Mn_3O_4}$
CSPIN	$\mathrm{Co_{1,2}Mn_{1.8}O_4}$	$\mathrm{Co_3O_4}$	$\mathrm{Co_{1.8}Mn_{1.2}O_4}$
LDH	$Co_{0.6}Mn_{0.4}(OH)_2Cl_{0.4}$	$Co_{0.9}Mn_{0.1}(OH)_2Cl_{0.9}$	$Co_{0.9}Mn_{0.1}(OH)_2Cl_{0.9}$

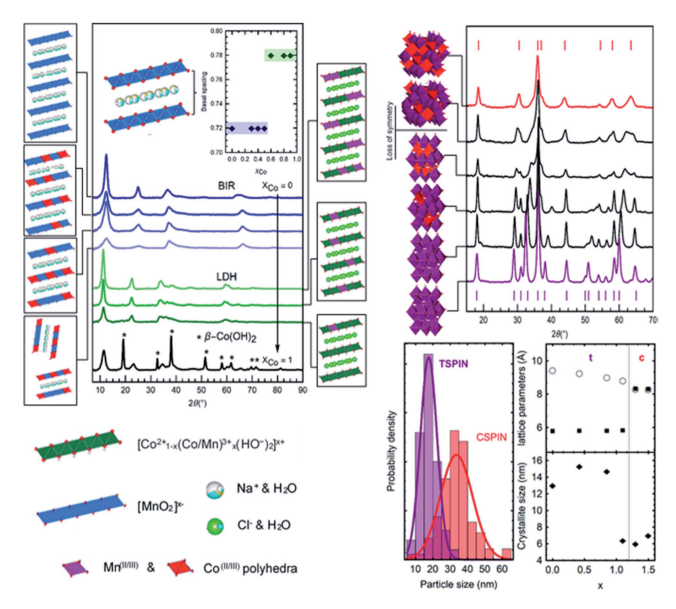


Fig. 7 Left: XRD traces of layered materials. Four birnessites with increasing amounts of cobalt, three layered double hydroxides, and a sample of a two-phase mixture of cobalt hydroxides. The basal spacing of the materials is shown in the inset. Structural representations are shown next to each trace. Right: XRD traces of spinel materials accompanied by their respective structural sketches. Cubic spinel (red) with increasing amounts of cobalt transitions into the tetragonal form (purple) with formula  $Co_xMn_{3-x}O_4$ . (Bottom left) Particle size measurements from SEM images of the boundary compositions are shown. (Bottom right) Lattice parameters and crystallite sizes as a function of Co content, x

structure of the particles produced in the entire range of Co<sub>x</sub>- $Mn_{3-x}O_4$  studied.

All samples are nanocrystalline. The samples heaviest in Co (Co/(Co + Mn) > 0.4) exhibit crystallite sizes in the order of 7 nm, while lower Co content produces more crystalline samples up to the 15 nm range as determined by the Scherrer equation. The latter samples were collected as single crystals of the same 15 nm average size (Fig. 6, bottom left), while the smaller cubic crystallites form polycrystalline particles of 35 nm with a broader dispersion. Using only Co to produce the c-spinel Co<sub>3</sub>O<sub>4</sub> proved unsuccessful as the Co remained under-oxidized by the ROS produced at the electrode, and a phase mixture with hydroxides formed instead of the pure Co<sub>3</sub>O<sub>4</sub>. To summarize,

two sets of nanoparticles are produced, smaller single crystal particles of approximately 15 nm in diameter composed of tetragonal  $Co_xMn_{3-x}O_4$  (x < 1), and larger particles of 35 nm composed of 6 nm crystallites of cubic  $Co_xMn_{3-x}O_4$  (x > 1).

#### **Electrochemical applications**

**OER electrocatalysis.** One composition from each structure synthesized was used to demonstrate the catalytic potential for oxygen evolution of materials synthesized by GDEx. The polarization curves taken after electrochemical conditioning of 100 cycles are shown in Fig. 8a. All samples show a significant current increase from the bare GC electrode, which does not achieve the target  $10 \text{ mA cm}^{-2}$  in the potential window scanned.

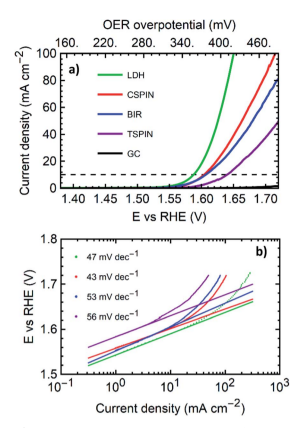


Fig. 8 Oxygen evolution reaction catalysis evaluation for one selected material from each of the 4 structures examined. (a) Linear scan voltammetry sweeps at 5 mV s $^{-1}$  in 1 M KOH solutions for the four structures. (b) Tafel plots of the same LSV data with the slopes in the range of 1 mA cm $^{-2}$  to 10 mA cm $^{-2}$  annotated.

A common metric for OER catalytic performance of flat (GC) electrodes is the overpotential at 10 mA cm $^{-2}$ , shown on the LSVs by the cutoff of the dashed lines. Nonetheless, two of the materials, the cubic spinel and the LDH, exhibit remarkable catalytic properties and are able to produce 100 mA cm $^{-2}$  below 500 mV of overpotential. The polarization curves in Fig. 8 are accompanied by the Tafel plot of the same curves from 1 mA cm $^{-2}$  to 100 mA cm $^{-2}$ . The full collection of parameters is presented in Table 2.

With overpotentials of 355 mV and 420 mV for 10 mA cm<sup>-2</sup> and 100 mA cm<sup>-2</sup> respectively, the LDH sample outperformed all materials tested. The low overpotentials and the low Tafel slope compare favorably with the literature and generally match or sometimes outperform most Co or Mn based metal oxide catalysts and bulk LDHs on GCE or similar substrates.<sup>37</sup> Table

Table 2 OER catalysis characteristic parameters

Structure	Co/(Co + Mn)	η10 (mV)	η100 (mV)	Tafel slope (mV dec <sup>-1</sup> )
TSPIN	0	410	_	46
CSPIN	0.5	375	485	43
BIR	0.25	375	_	53
LDH	0.5	355	420	47

ESI-1† shows a collection of state of the art materials of Co and Mn with similar structures for comparison. Within each structural group, the samples synthesized here exhibit promising performance. Among the spinels, the cubic structure performed far better than the tetragonal sample. The tetragonal structure tested was pure  $\rm Mn_3O_4$ . Here the addition of 50% Co, and as a consequence, the transition to the cubic structure reduces the overpotential at 10 mA cm<sup>-2</sup> by 35 mV with outstanding performance and better metrics than the bulk of mixed metal spinels in the literature.<sup>37</sup>

It is worth noting that the TSPIN sample contained the smaller particles, with approximately 15 nm in diameter (via SEM), as opposed to the 35 nm for the CSPIN sample. Smaller particle sizes improve the catalysis due to the larger surface area. However, the presence of cobalt, together with the structural change it induces, produces better catalytic properties despite the larger particle size of the cubic spinels. The CSPIN and LDH were both made with the same Mn/Co elemental ratio. The sheet-like structure of the LDH played a key role in making a more active material. LDH nanosheets have been shown to possess outstanding OER electrocatalytic properties due to the large active area given by the morphology, and the highly defective mixed-valence structure. At the same time, the BIR nanosheets containing half the amount of cobalt performed similarly to the cobalt-rich CSPIN, with the same  $\eta 10$  but a slightly higher Tafel slope.

Active material for Na-ion battery cathodes. The Co-doped birnessite (Co<sub>0.25</sub>Na<sub>0.75</sub>MnO<sub>2</sub>) sample was tested as a cathode material for Na-ion batteries. The sodium intercalation between the nanosheets offers a natural transport channel for charge-discharge cycles. The pristine samples were used as-synthesized, that is, no post-processing was performed. Similar materials are often calcined<sup>38,39</sup> or synthesized in a hydrothermal process to increase the Na (or Li, K) content to obtain higher capacities.<sup>39-41</sup> The dried sample was mixed with a carbon binder and coated on an aluminium current collector to be tested in a non-aqueous solvent, opposite to a Na anode.

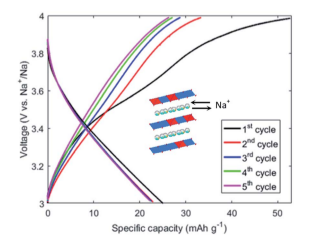


Fig. 9 First 5 cycles of the birnessite  $Co_{0.25}Na_{0.75}MnO_2$  material synthesized by the GDEx process. A current of 0.004 mA cm<sup>-2</sup> (0.1C) was applied with cut-off potentials between 3 V and 4 V.

The open-circuit voltage of the as-prepared electrode was 3.1 V. The electrode was charged and discharged between a lower and upper cutoff voltage of 3 and 4 V, respectively. A current density of 0.004 mA cm<sup>-2</sup> was applied, corresponding to 0.1 C rate. Fig. 9 illustrates the first 5 cycles of the Na//NaMnO cell. A first charge specific capacity of 53 mA h g<sup>-1</sup> and a reversible capacity of 25 mA h g<sup>-1</sup> were recorded. While these values are not among the highest reported in the literature for birnessite-type materials, for room-temperature as-synthesized structures, there is potential for optimization. Post-processing techniques and composition control can be exerted to improve from this

# Conclusions

The level of control achieved over structure and composition with a single system can open the door to the systematic creation of libraries of materials to develop, for instance, better electrocatalysts. The technique presented, GDEx, offers materials with a wide range of structures, compositions, and sizes, with controlled properties, synthesized in the same environment with minimal chemical addition, making it a landmark tool to exploit in material design. The nature of the reactor allows for one set of conditions to be used, and merely changing the inlet solution, to produce the target structure, all with a highly scalable configuration. To summarize, the materials synthesized and discussed include birnessites with Co doping of up to 40% of the Mn sites, tetragonal and cubic mixed Co/Mn spinels with fine control over the structural distortions and transitions, layered Co hydroxides, and CoMn-layered double hydroxides. All these exhibited nanocrystallites, as well as particle sizes down to 15 nm, both for single crystals and polycrystalline particles. In future studies, the elements (i.e., metal ion precursors) can be swapped by other elements of interest (Ni, Fe), and tertiary mixtures could be as easily achieved. Finally, birnessites, cubic spinels, and layered double hydroxides were tested for OER catalysis, and each exhibited at least state-of-the-art catalytic properties.

Partial conclusions can be drawn to emphasize the benefit of the systematic material synthesis employed here. Both layered compounds outperformed the spinel materials made of similar metal compositions. At the same time, within each structural family, the addition of cobalt improved the performance significantly. In this way, structure-property relationships can begin to be built across structures and compositions with a single synthesis process. By fixing the charge applied and residence time in the flow cell, the total metal concentration and Co/Mn ratio can be varied and directly result in a range of materials to optimize. Once the structure and composition result in the desired properties, further degrees of freedom can be added by varying the volumetric flow rate and current, to achieve control over other parameters such as particle or crystallite size. The choice of supporting electrolyte can also be chosen to modulate the basal spacing in the layered materials, as the cations (Na<sup>+</sup>) or anions (Cl<sup>-</sup>) find themselves intercalated between the layers. The choice of larger ions (e.g.: Cs<sup>+</sup>, NO<sub>3</sub><sup>-</sup>) can result in enlarged interlayer distances, which are a factor in

electrocatalytic performance. Above all, GDEx emerges as a sustainable synthesis method for vast libraries of functional nanostructures.

# Conflicts of interest

There are no conflicts to declare.

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