Modifying the Surface Structure of Perovskite-Based Catalysts by Nanoparticle Exsolution

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Abstract: In heterogeneous catalysis, surfaces decorated with uniformly dispersed, catalytically-active (nano)particles are a key requirement for excellent performance. Beside standard catalyst preparation routines—with limitations in controlling catalyst surface structure (i.e., particle size distribution or dispersion)—we present here a novel time efficient route to precisely tailor catalyst surface morphology and composition of perovskites. Perovskite-type oxides of nominal composition ABO$_3$ with transition metal cations on the B-site can exsolve the B-site transition metal upon controlled reduction. In this exsolution process, the transition metal emerges from the oxide lattice and migrates to the surface where it forms catalytically active nanoparticles. Doping the B-site with reducible and catalytically highly active elements, offers the opportunity of tailoring properties of exsolution catalysts. Here, we present the synthesis of two novel perovskite catalysts Nd$_{0.6}$Ca$_{0.4}$FeO$_{3-δ}$ and Nd$_{0.6}$Ca$_{0.4}$Fe$_{0.9}$Co$_{0.1}$O$_{3-δ}$ with characterisation by (in situ) XRD, SEM/TEM and XPS, supported by theory (DFT+U). Fe nanoparticle formation was observed for Nd$_{0.6}$Ca$_{0.4}$FeO$_{3-δ}$. In comparison, B site cobalt doping leads, already at lower reduction temperatures, to formation of finely dispersed Co nanoparticles on the surface. These novel perovskite-type catalysts are highly promising for applications in chemical energy conversion. First measurements revealed that exsolved Co nanoparticles significantly improve the catalytic activity for CO$_2$ activation via reverse water gas shift reaction.

Keywords: exsolution; nanoparticles; perovskites; reverse water-gas shift reaction; tailored surfaces

1. Introduction

In heterogeneous catalysis, controllable surface properties and a maximum amount of stable uniformly-dispersed catalytically highly active sites on the surface of a porous material are of key importance. Typically, catalysts consist of metal, alloy, or oxide nanoparticles embedded in an oxide support material. Depending on the type of catalytic reaction, active sites are either just nanoparticles themselves or the combination of nanoparticles and oxide support. Ideally, catalysts exhibit long-term stability and activity during reaction. Therefore, resistance to catalyst poisons, inhibition of carbon deposition (blocking of active sites), and prevention of particle agglomeration and sintering (loss of active surface area) is essential.

Typically, these structures are prepared by deposition, impregnation or precipitation techniques [1,2] followed by catalyst activation prior to reactions via oxidation and reduction [3]. Although these approaches are applied widely, they offer limited control over size, distribution, and anchorage of deposited species, not only during preparation but also during catalyst activation or operation, and they might be time consuming and costly [4].
Alternatively, an emerging concept is to grow the nanoparticles in situ, by reduction or during catalytic reaction, directly from the (porous) oxide backbone support itself [4–6]. Perovskites (nominal composition \( \text{ABO}_3 \) with A and B being a large and small cation, respectively) can incorporate catalytically highly active elements as cations on the B-site of the perovskite lattice (e.g., Ni [4,7], Fe [4,8], Co [9], Cu [4,9], Pt [10], Pd [10,11]). Upon exposure to reducing conditions, these elements can be partly exsolved as nanoparticles, thus opening the possibility of in situ growth of active catalysts [12], see sketch in Figure 1. In comparison to traditional deposition techniques, this process produces finer and higher dispersed catalyst nanoparticles [4,13,14] and is more time- and cost-effective, as it does not require multiple ‘deposition’ steps or expensive precursors [15]. Furthermore, it has been shown that nanoparticles formed by exsolution exhibit improved sintering stability during catalytic reaction [16]. In addition, due to a possible reversibility of the nanoparticle exsolution process, particle agglomeration can be avoided by reoxidation cycles [17], hence, greatly enhancing the catalyst lifetime [12,18].

![Diagram of Perovskite and Exsolution Process](image)

**Figure 1.** (a) Perovskite lattice structure; (b) simplified exsolution process.

Mechanisms for nanoparticle exsolution from perovskites were investigated already by multiple groups [4,11,19–21]. It is generally accepted that exsolution catalysts need easily reducible dopant cations [22], such as Ru, Co, Pd, or Ni [23], that will form metallic deposits on the perovskite surface. Thus, depending on temperature and oxygen chemical potential in the gas phase (i.e., the strength of reduction), dopants and in some cases also lattice cations are reduced and exsolved as nanoparticles [8,24]. As an explanation for exsolution mechanisms, Neagu et al. [4] proposed that oxygen vacancies created by reduction destabilize the perovskite lattice, especially when their concentration is high and when additionally A-site cation vacancies are present. Such combined oxygen and A-site cation vacancy formations may locally cause spontaneous exsolution of B-site species in an attempt to re-establish stoichiometry across all sites. Oh et al. [19] implied that the morphological evolution of nanoparticles could be explained using a simple energy-based model, which accounts for interplay between surface free energy and strain energy induced by the included metal nucleate. By quantitative strain field modelling, they could demonstrate that this provides the driving force for exsolution processes. Similarly, Han and co-workers applied classical nucleation theory to the exsolution process and found the strain between metal particle and supporting oxide being a potential influencing factor for the particle size.

Usually, formation of nanoparticles on the surface is accomplished via a reductive treatment. For example, Arrivé et al. [25] showed for Ni doped \( \text{LaSrTiO}_3 \) that pre-reduction in \( \text{H}_2 \) at 1573 K leads to exsolution of Ni particles on the surface of solid oxide electrochemical cells leading to improved electrochemical performance. Additionally, Papargyriou and Irvine demonstrated Ni exsolution from Ni-doped \( \text{La}_{0.75}\text{Sr}_{0.25}\text{Cr}_{0.5}\text{Fe}_{0.5}\text{O}_3 \) perovskite by \( \text{H}_2 \) reduction [14]. Nanoparticles with a size of 30–50 nm were formed on the surface, leading to an improvement of catalytic properties of the material.
Similarly, Sun et al. [13] could observe Ni nanoparticle exsolution from Ni-doped La$_{0.7}$Sr$_{0.3}$CrO$_3$ upon pre-reduction in H$_2$. Most of the recent studies on exsolution phenomena are reported and studied by the electrochemical community (mainly with respect to fuel cell technology), but utilization of this effect for preparing tailored heterogeneous catalyst materials is still rare—even though the initial idea came indeed from the catalysis community [12].

Here we present synthesis and characterization of the novel perovskites Nd$_{0.6}$Ca$_{0.4}$FeO$_{3-δ}$ and Nd$_{0.6}$Ca$_{0.4}$Fe$_0.9$Co$_{0.1}$O$_{3-δ}$. Their exsolution properties (by reductive treatment in humidified H$_2$) were studied utilizing (in situ) X-ray diffraction (XRD), X-ray photoelectron spectroscopy (XPS), scanning electron microscopy (SEM), and transmission electron microscopy (TEM). To highlight the effect of nanoparticle exsolution on catalytic activity, the novel perovskites were then tested regarding their suitability for reverse water gas shift reaction (rWGS), for the Co-doped perovskite reductive treatment caused the formation of finely dispersed metallic Co nanoparticles, which greatly enhanced the rWGS reactivity.

2. Results and Discussion

2.1. Materials Design

For synthesis of the novel materials, we made a judicious choice of composition of the perovskite host lattice, enabling us future use of a variety of catalytically-active elements as dopants. An optimized perovskite composition enables exsolution with controlled nanoparticle size. For the A-site neodymium and calcium have been chosen with a 6:4 ratio. Usually La is one of the most common elements for A-site. However, to avoid XPS peak overlap with some future potentially interesting dopant elements (especially Ni) when performing in situ near ambient pressure XPS (NAP-XPS) measurements, we used Nd instead. Acceptor doping of the A-site with Ca introduces electronic defects (holes) and oxygen vacancies, which improves the electron and oxygen anion conductivity of the material. Density Functional Theory (DFT) calculations (utilizing WIEN2k [26]) suggest that this electronic defects are delocalized over both oxygen and iron atoms. Oxygen participation in the delocalization is revealed by empty O-p states above the Fermi level, that in the bulk are occupied (details, Section 2.2), which is also in accordance with results from X-ray absorption spectroscopy (XAS) on a very similar material (La$_{0.6}$Sr$_{0.4}$FeO$_{3-δ}$) [27]. The contribution of iron atoms is indicated by a reduction of their magnetic moment (around 0.5 µ$_B$) corresponding to partial oxidation. Additionally, a Jahn–Teller distortion for the iron atoms in question is found (where four bonds get shortened by about 6.9%, while the other two are elongated by about 6.6%), which corresponds to the changed electronic configuration and the smaller size of Fe$^{4+}$ ions.

Variation of the A-site dopant concentration allows additional fine-tuning of the exsolution process [28]. In a preliminary study, we saw that for perovskites without B-site doping the temperature necessary for exsolution is higher with an increasing amount of Ca doping. The B-site cation is iron, which was replaced by 10% of cobalt for the doped catalyst. As discussed later in detail, the undoped perovskite exhibited iron nanoparticle exsolution at 948 K. When the material is additionally doped on the B-site with Co, exsolution of both Fe and Co nanoparticles, or even formation of FeCo alloy nanoparticles would be possible. Therefore, we settled for the considerably high amount of Ca A-site doping of 40%. Consequently, the host lattice is expected to be quite stable, leading to preferential exsolution of cobalt upon reductive treatment of the doped material.

After synthesis, Nd$_{0.6}$Ca$_{0.4}$FeO$_{3-δ}$ and Nd$_{0.6}$Ca$_{0.4}$Fe$_0.9$Co$_{0.1}$O$_{3-δ}$, were characterized by XRD for identification of the crystal structure. For both freshly synthesized materials (Figure 2a,b bottom diffractograms), only the pure perovskite phase was observed. All the reflexes in the pattern at RT correspond to this phase, they are marked with P in the graph, and their hkl-indices are given in Figure 2a. The patterns of the undoped and doped material were almost identical, indicating equivalent structures. The main reflexes were at 20 of 32.9°, 47.0°, and 58.7°, respectively. Rietveld refinement in combination with database structures for similar materials was utilized to propose the crystal structure.
An orthorhombic lattice (lattice parameters: \( a = 5.533 \, \text{Å}, b = 7.726 \, \text{Å}, c = 5.449 \, \text{Å} \)) with space group \( Pnma \) (Nr. 62) was revealed. For an ideal cubic perovskite structure, ionic radii of all elements have to fulfil a certain size relation, which is reflected by a Goldschmidt tolerance factor of 1. For our materials, this factor is slightly below 1, leading to a distortion of the ideal structure (tilted B-site coordination octahedrons) and a reduced symmetry [29].

![XRD patterns for (a) \( \text{Nd}_{0.6}\text{Ca}_{0.4}\text{FeO}_3-\delta \) and (b) \( \text{Nd}_{0.6}\text{Ca}_{0.4}\text{Fe}_{0.9}\text{Co}_{0.1}\text{O}_3-\delta \). The hkl-indices of the undoped perovskite phase correspond also to the Co-doped material. The bottom diffractograms were collected at RT after oxidation in \( \text{O}_2 \) at 873 K. Patterns at elevated temperatures were collected in situ during reduction in humidified \( \text{H}_2 \) (ratio \( \text{H}_2:\text{H}_2\text{O} \approx 32:1, p = 1 \, \text{bar} \)) at increasing temperature. For the undoped perovskite, Fe exsolution was observable at 923 K. The Co-doped material shows formation of a metallic bcc Fe/Co phase above 823 K.](image)

### 2.2. DFT Calculations

To gain insight on the effect of A-site doping of \( \text{NdFeO}_3 \) with \( \text{Ca} \) on atomic and electronic structure of the perovskite, DFT calculations of \( \text{NdFeO}_3 \) and \( \text{Nd}_{0.5}\text{Ca}_{0.5}\text{FeO}_3 \) were performed. A previous theoretical study of such a system by Wang et al. [30] treated the 4f electrons of Nd in the frozen core approximation.
Here, however, a full potential (FP-(L)APW+lo) calculation was done. The doped material shows more strongly distorted Fe–O-octahedra compared to the octahedra in bulk NdFeO₃: while Fe–O bond lengths in the bulk materials differ only by 1.1%, the longest Fe–O bond in the doped material is 5.4% and 14.5% longer than the shortest one for Fe³⁺ and Fe⁴⁺, respectively. DFT+U finds a bandgap of 2.38 eV for bulk NdFeO₃, while the doped material is metallic. The Fermi level is shifted into bands that are fully occupied in the bulk due to the doping—as shown in the density of states (DOS) in Figure 3 (states, that are occupied in the bulk, are empty in the doped material because replacing Nd³⁺ with Ca²⁺ reduces the number of valence electrons). Both oxygen p states and iron d states contribute to the empty states directly above the Fermi level, indicating a partial delocalization of the electron hole over many oxygen atoms on one hand and a partial oxidation of Fe atoms (Fe⁴⁺) on the other hand.

![Figure 3](image-url)

**Figure 3.** Total (black solid line) and partial (dashed colored lines) DOS for Nd₀.₅Ca₀.₃FeO₃. The dashed vertical line represents the Fermi level. The total DOS of bulk NdFeO₃ is included for reference as grey area. The inset focuses on the DOS around the Fermi level. For Nd, Fe³⁺, and Fe⁴⁺ the partial DOS is shown only for one spin, the antiferromagnetic counterparts are omitted.

The Fe atoms in the bulk unit cell are pair wise equivalent (spin-up and spin-down atoms differ only with respect to the magnetic structure of their next nearest Nd neighbors, which shows that the magnetic ordering on the Nd sublattice has virtually no effect on the Fe atoms in the bulk)—in the doped material, this equivalency is lifted: The iron atoms (both spin-up and spin-down) with slightly larger Fe–Ca distances show a reduction of their respective magnetic moments (going from 4.0 µB to 3.4 µB). This hints toward a partial oxidation of Fe³⁺ atoms. For Fe⁴⁺ the larger moment derives from fully occupied spin-up states; this occupation is reduced for Fe⁴⁺. The partial oxidation is further evidenced by a Jahn–Teller distortion around the oxidized atoms (four bonds are shortened by 6.9%, two bonds are elongated by 6.6%) corresponding to the change in electronic configuration and the smaller ionic radius of Fe⁴⁺.

### 2.3. Exsolution Properties

For the following reductive treatment we used humidified H₂ at 1 bar with a H₂/H₂O ratio of ~32:1, which corresponds to an oxygen partial pressure of only 5.3 × 10⁻²⁶ bar. The role of water in the gas phase is to provide an oxygen source for a well-defined oxygen partial pressure (pO₂, explanation see SI). This is required to be able to exactly compare our results to future planned electrochemical studies and to other work on exsolution in literature [31]. For the undoped perovskite, in situ XRD measurements revealed the formation of a metallic Fe bcc phase above 948 K at 2θ = 44.4° (Figure 2a). Important is, that although we observed nanoparticle exsolution, the perovskite itself stayed intact, as indicated by the XRD pattern. SEM combined with energy dispersive X-ray spectroscopy (EDX)
confirmed the formation of larger Fe particles with a size of ~470 nm on the surface, Figure 4. It is clearly visible that the exsolved particles are well shaped crystals growing from the surface (Figure 4a) and that they primarily consist of iron (Figure 4b). Formation of metallic iron on the surface was also visible in the Fe 2p XPS signal at 707.0 eV (SI, Figure S1) [8].

Figure 4. SEM images of the Nd$_{0.4}$Ca$_{0.6}$FeO$_3$ perovskite after reductive treatment with humidified H$_2$ at 973 K (ratio H$_2$:H$_2$O ≈ 32:1, p = 1 bar). Image (a) displays the exsolved particles, growing from the surface. The magnified particle (inset) has a diameter of ~470 nm. In image (b), an EDX mapping of Nd and Fe (overlay) confirms the iron nature of the nanoparticles (more Fe and less Nd at the positions where bright particles are visible in the secondary electron image).

For comparison, doping with cobalt facilitated exsolution already at lower temperatures. Upon reductive treatment a Fe/Co bcc diffraction line evolved between 823 and 848 K in the XRD pattern at 2θ = 44.8° (Figure 2b). Unfortunately, it was not possible to distinguish clearly between Fe and Co with XRD due to an overlap of the diffraction lines of both metals. SEM images showed the formation of finely dispersed nanoparticles on the surface with an average size of 30 nm to 40 nm, Figure 5a.

Further analysis of the doped perovskite with TEM and scanning TEM (STEM) combined with EDX revealed the presence of Co nanoparticles on the surface. For these studies, the thin film samples were reduced at 923 K in humidified hydrogen (2.5% H$_2$ /2.5% H$_2$O/balance Ar) atmosphere in a tube furnace. A core-shell structure is visible in Figure 5b. In high resolution mode (HRTEM, Figure 5c), lattice planes can be observed, the Fourier transformation of this area (FFT, Figure 5d) reveals two phases, CoO$_3$O$_4$ and Co. These observations can be explained by the air exposure of the sample between exsolution and TEM analysis. The original particle consisted of metallic cobalt. Its surface was oxidized upon air exposure, forming the cobalt oxide shell with a diameter of 30 nm. The core is the remaining unoxidized cobalt and has a diameter of 18 nm. An EDX mapping (Figure 5e) verified that the predominant metal in the particle was cobalt. The cobalt concentration (color intensity) is higher in the core and lower in the shell, in agreement with the proposed Co and CoO$_3$O$_4$ phases. Formation of metallic cobalt upon nanoparticle exsolution could also be confirmed with in situ NAP-XPS. A metallic Co2p signal at ~776 eV evolved when exposing the perovskite to H$_2$/H$_2$O at 873 K (SI, Figure S2).
Our results highlight that, already, the host perovskite lattice is capable of Fe particle exsolution at higher reduction temperatures (>948 K). The addition of an easier reducible dopant element (here, cobalt) leads to its preferential exsolution at significantly lower temperatures (823 K) in the form of finely dispersed nanoparticles. It cannot be ruled out that by further reduction at even higher temperatures also iron may be exsolved to the surface, either in form of additional Fe nanoparticles, or by alloy formation with the already present Co particles (further studies are planned). In a related study, Zhu and co-workers only observed CoFe alloy formation at temperatures above 1023 K from La$_{0.8}$Sr$_{0.2}$Co$_{0.45}$Fe$_{0.45}$Nb$_{0.1}$O$_{3-δ}$ [32].

In comparison to our studied perovskites, other groups reported generally much higher reduction temperatures for nanoparticle exsolution. Gao et al. showed Ni exsolution from La$_{0.52}$Sr$_{0.28}$Ti$_{0.94}$Ni$_{0.06}$O$_{3-δ}$ after reduction at 1173 K [33]. Similarly, Kim et al. highlighted Ni exsolution from La$_{0.25}$Sr$_{0.75}$Ti$_{0.9}$Ni$_{0.1}$O$_{3-δ}$ at the same temperature [34]. Earlier work of Irvine and co-workers demonstrated nanoparticle formation (Fe, Ni, Cu) upon reduction at temperatures between 1173 K and 1273 K [4]. Recently, they proved the possibility to lower the exsolution temperature by inducing lattice strain to perovskite thin films resulting in Ni exsolution at 823 K [25]. A lower temperature was also reported by Steiger et al. for LaFe$_{0.8}$Ni$_{0.2}$O$_{3-δ}$ with Ni nanoparticle exsolution by reduction at 873 K [35]. Joo et al. reduced a cobalt doped Mn-based perovskite at 1023 K to achieve Co nanoparticle exsolution [36].

A reason for the reported high exsolution temperatures is that these studies are mainly conducted in the field of solid oxide fuel cells with their operation temperatures in the 873–1173 K range. For catalytic applications, however, lowering the exsolution temperature is utmost important, especially for future application in industrial processes. Moreover, it has been shown that nanoparticles formed...
by exsolution exhibit improved sintering stability [16], preventing one of the major problems of catalyst deactivation in heterogeneous catalysis.

2.4. Catalytic Test Measurements

As outlined in the introduction, nanoparticle exsolution is an elegant and fast route for catalyst preparation. To demonstrate the capability of our tailored perovskite materials, first catalytic tests for CO\textsubscript{2} activation via rWGS reaction were performed. We have chosen rWGS as it can be considered a promising technology for converting CO\textsubscript{2} to CO, which can then be further converted via CO hydrogenation to liquid fuels (e.g., diesel, gasoline, and alcohols). This approach has the advantage of high throughput rates, selectivity, and technological readiness [37]. Furthermore, we have shown in previous work that perovskites are excellent materials for CO\textsubscript{2} activation, e.g., by electrocatalytic CO\textsubscript{2} reduction [31].

After introducing the starting reactant gas flow with a CO\textsubscript{2}/H\textsubscript{2} ratio of 1:1, the temperature was increased stepwise up to 973 K. Onset of CO formation could be observed above 573 K, Figure 6. Already the undoped perovskite Nd\textsubscript{0.6}Ca\textsubscript{0.4}FeO\textsubscript{3-δ} showed catalytic activity (blue line). This was expected, as iron is known to be catalytically active for rWGS [37]. However, additional doping with Co and the resulting Co nanoparticle exsolution from Nd\textsubscript{0.6}Ca\textsubscript{0.4}Fe\textsubscript{0.9}Co\textsubscript{0.1}O\textsubscript{3-δ} resulted in a strong improvement of the catalytic activity (green line). Additionally, for Co, it has been shown that it is catalytically highly active for rWGS [37]. The combination of an already active perovskite host material as support and the catalytically active dopant material (here Co), that forms finely dispersed nanoparticles on the surface, lead to a strong improvement of catalytic reactivity.

![Figure 6](image_url)  
**Figure 6.** rWGS activity of undoped and Co doped catalyst. The temperature was stepwise increased with reaction onset above 573 K. For better comparability, the CO formation is given as specific activity (normalized by active catalyst surface area). Co nanoparticle exsolution from Nd\textsubscript{0.6}Ca\textsubscript{0.4}Fe\textsubscript{0.9}Co\textsubscript{0.1}O\textsubscript{3-δ} improved the catalytic activity significantly.

By correlating nanoparticle exsolution with catalytic testing, we could demonstrate the impressive capabilities of perovskite based catalysts. In combination with doping of the B-site by catalytically highly active elements these materials enable manifold routes for catalyst engineering, opening the possibility of tailoring catalyst surfaces to actual needs.
3. Materials and Methods

3.1. Synthesis of Novel Perovskites

The perovskite powders were synthesized via the Pechini route [38]. The cations were mixed in the desired stoichiometric ratio. The starting materials were Nd$_2$O$_3$ (99.999%, Alfa Aesar, Haverhill, MA, USA), CaCO$_3$ (99.999%, Alfa Aesar, Haverhill, MA, USA), Fe (99.99%, Sigma-Aldrich, St. Louis, MO, USA), and Co (99.995%, Sigma-Aldrich, St. Louis, MO, USA). The compounds were weighted accordingly and dissolved in HNO$_3$ (double distilled, 65%, Merck, Darmstadt, Germany). Cation complexes were formed by adding citric acid (99.9998% trace metals pure, Fluka, Honeywell International, Charlotte, NC, USA) in a molar ratio of 1.2 with respect to the cations. After evaporation of H$_2$O, the resulting gel was heated until self-ignition took place. The obtained powder was calcined for three hours at 1073 K. After grinding (i.e., with a mortar to obtain homogeneous powders for better characterization), the resulting products were used for the (in situ) XRD, standard XPS, SEM studies, and catalytic testing.

For the in situ NAP-XPS and TEM experiments, thin film samples of Nd$_{0.6}$Ca$_{0.4}$Fe$_{0.9}$Co$_{0.1}$O$_3$ were prepared. Therefore, the powder was pressed into a pellet using a uniaxial press. Sintering of the pellet was carried out at 1523 K for 12 h. Dense perovskite thin films were then produced by pulsed laser deposition (PLD), using the Nd$_{0.6}$Ca$_{0.4}$Fe$_{0.9}$Co$_{0.1}$O$_3$ pellet as a target. Ablation was carried out at 873 K on (100)-oriented yttria stabilized zirconia (YSZ) single crystals (CrysTec GmbH, Berlin, Germany) (10 × 10 × 0.5 mm) as substrates in 0.04 mbar oxygen background pressure using a KrF excimer laser (Lambda COMPexPro 201F, λ = 248 nm, Coherent, Santa Clara, CA, USA) with 400 mJ laser pulses at 5 Hz.

3.2. Characterization Methods

Standard XPS measurements were performed using a photoelectron spectrometer (SPECS Surface Nano Analysis GmbH, Berlin, Germany) equipped with a PHOIBOS-100-MCD-5 hemispherical energy analyser and an X-ray source XR50 with a double Al/Mg anode. For measurements, the samples were supported onto double-sided conducting carbon tape.

In situ XPS measurements were performed on a lab-based in situ NAP-XPS system equipped with a Phoibos 150 NAP hemispherical analyzer and a XR 50 MF X-ray source (microfocus), all from SPECS (Berlin, Germany). Spectra were recorded with monochromatic Al K$_\alpha$ radiation and data were analyzed with CasaXPS software (Casa Software Ltd., Teignmouth, UK). Peaks were fitted after Shirley or linear background subtraction with Gauss−Lorentz sum functions. Peak positions and full width at half-maximum (FWHM) were left unconstrained. Peak positions were referenced either to the valence band signal (Fermi edge) or to the carbon signal.

The powder XRD measurements were carried out on a PANalytical X’Pert Pro diffractometer (Malvern Panalytical, Malvern, UK) in Bragg–Brentano geometry using a mirror for separating the Cu K$_{α1,2}$ radiation and an X’Celerator linear detector. For the in situ experiments, an XRK 900 chamber (Anton Paar, Graz Austria) was used. The pristine samples were pre-treated in O$_2$ at 875 K for 30 min. After cooling to room temperature, the atmosphere was changed to humidified H$_2$ (using a bubbler at room temperature, p = 1 bar, H$_2$:H$_2$O ≈ 32:1) and the temperature was increased stepwise by 25 K. At each step, an in situ XRD measurement was performed after waiting for 30 min. The data were analyzed using the HighScore Plus software (Malvern Panalytical, Malvern, UK) [39] and the PDF-4+2019 database (ICDD - International Centre for Diffraction Data, Newtown Square, PA, USA) [40]. The reflexes were assigned according to database entries.

The SEM experiments were done on a Quanta 250 FEGSEM (FEI Company, Hillsboro, OR, USA) equipped with an Octane Elite X-ray detector (EDAX Inc, Mahwah, NJ, USA) using an acceleration voltage of 5 kV to achieve sufficient surface-sensitivity. The images were recorded using secondary electrons.
For the (scanning) TEM experiments, the sample was first coated with protective C and Pt layers. Then, an ultra-thin sample was cut containing single, well-separated nanoparticles using a focused Ga ion beam in a Quanta 200 3D DBFIB (FEI Company, Hillsboro, OR, USA). The TEM measurements were performed with an acceleration voltage of 200 kV in a Tecnai F20 microscope (FEI Company, Hillsboro, OR, USA) equipped with a RIO camera (Gatan, Pleasanton, CA, USA), an Apollo X-ray detector (EDAX Inc, Mahwah, NJ, USA), and a GIF Tridiem energy filter (Gatan, Pleasanton, CA, USA). First, the high-resolution images were recorded in TEM mode. Then, the X-ray measurements were performed in the same area in STEM mode.

3.3. DFT Calculations

Spin-polarized DFT (DFT+U) calculations were performed using the program package WIEN2k (Institute of Materials Chemistry, TU Wien, Vienna, Austria) [26], which is an implementation of the augmented plane wave + local orbital method [41] treating all electrons in a full potential (FP-(L)APW+lo). The GGA functional PBE [42] was used to treat exchange-correlation with an additional Hubbard-U [43] to account for localized Fe 3d and Nd 4f electrons, respectively. In both cases a value of $U_{\text{eff}} = 4\,\text{eV}$ was used [44,45].

Using the XRD data presented in Section 2.1 (cp. Figure 2a) a unit cell containing four formula units of NdFeO$_3$ was set up. This cell was used to simulate Nd$_{0.5}$Ca$_{0.5}$FeO$_3$ (corresponding to a 50% doping) by substituting two Nd atoms with Ca (see supporting information, Figure S3). The atomic positions were optimized until residual forces were below 1 mRy/bohr.

Both Nd and Fe order magnetically: Spin-up and spin-down Fe atoms make up a type G [46] antiferromagnetic sublattice [47], where the spin of any given iron atom is aligned antiparallel to all its nearest neighbors. Although Nd atoms in NdFeO$_3$ show long range antiferromagnetic order for very low temperatures (experiments find ordering for temperatures below 2 K [47,48]), initial calculations were done with a ferromagnetic using setup to save on compute time, since the effect of ferromagnetic Nd ordering on the moments of the other atoms is small (in the order of 0.01 $\mu_B$). Final calculations were performed with optimized positions from initial runs and antiferromagnetically ordered Nd atoms—since all residual forces remained small (below 1.5 mRy/bohr) no additional structure optimization was carried out.

The choice for atomic sphere radii for all calculations was 2.20, 1.92, and 1.60 bohr for Nd/Ca, Fe, and O respectively. A plane wave cutoff of $R K_{\text{max}} = 8$ was used to determine the basis set size, where $R$ is the radius of the smallest atomic sphere and $K_{\text{max}}$ denotes the largest considered vector in reciprocal space. The Brillouin zone of the doped material was sampled using a $9 \times 6 \times 9$ Monkhorst–Pack k-mesh [49].

The calculations of the doped material were compared with results of undoped bulk NdFeO$_3$ [50], derived using the same computational parameters.

3.4. Catalytic Testing

Catalytic tests for CO$_2$ activation via rWGS reaction (CO$_2$ + H$_2$ ⇌ CO + H$_2$O) were performed in a 6 mm tubular flow reactor (quartz glass, inner diameter 4 mm) at atmospheric pressure. Online gas analysis was conducted by a Micro-GC (Fusion 3000A, Inficon, Bad Ragaz, Switzerland) which was sampling every ~2 min. The total flow was 12 mL/min, with educt flow of 6 mL/min with a ratio of 1/1 for CO$_2$/H$_2$ and argon added as carrier gas (all gases provided by Messer Group GmbH, Bad Soden, Germany). A blind test was performed without catalyst to rule out any catalytic activity of the reactor itself. For the reactions, the pure perovskite powder catalysts (20–60 mg) were supported directly on a glass wool bed. A PID controller (EMSR EUROTHERM GmbH, Vienna, Austria) was used for control of the reactor heater. For a better comparability of the two perovskites, the CO formation was normalized by the active catalyst surface area (BET area) and is, therefore, given as specific activity (mol/m$^2$/s).
4. Conclusions

We have demonstrated that exsolution is an elegant process to produce catalytically active surfaces with well-dispersed metal nanoparticles. This is especially interesting for fast and simple catalyst preparation, as we have shown for Nd$_{0.6}$Ca$_{0.4}$FeO$_{3-δ}$ and Nd$_{0.6}$Ca$_{0.4}$Fe$_{0.9}$Co$_{0.1}$O$_{3-δ}$. Whereas the undoped material exhibits Fe nanoparticle exsolution at 948 K, the doped material facilitates Co nanoparticle exsolution already starting at 823 K. Furthermore, these nanoparticles are smaller (average size 30–40 nm) and well dispersed over the surface. The lower exsolution temperature makes the material especially interesting for catalytic applications as demonstrated for CO$_2$ activation via rWGS. The undoped Nd$_{0.6}$Ca$_{0.4}$FeO$_{3-δ}$ is already active for rWGS and with additional Co doping the perovskite exhibits significantly increased reactivity due to Co nanoparticle exsolution. Our novel materials open unpredicted possibilities for customized engineering of support materials, including doping with catalytically-active elements that are capable of forming nanoparticles on the surface.

Supplementary Materials: The following figure is available online at http://www.mdpi.com/2073-4344/10/3/268/s1, Figure S1. XPS spectra of Nd$_{0.6}$Ca$_{0.4}$FeO$_3$, Figure S2. XPS spectra of Nd$_{0.6}$Ca$_{0.4}$Fe$_{0.9}$Co$_{0.1}$O$_3$, Figure S3. DFT structure of Nd$_{0.5}$Ca$_{0.5}$FeO$_3$.

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