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Preparation of isolated Co_3O_4 and fcc-Co crystallites in the nanometre range employing exfoliated graphite as novel support material†

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The inert nature of graphitic samples allows for characterisation of rather isolated supported nanoparticles in model catalysts, as long as sufficiently large inter-particle distances are obtained. However, the low surface area of graphite and the little interaction with nanoparticles result in a challenging application of conventional preparation routes in practice. In the present study, a set of graphitic carbon materials was characterised in order to identify potential support materials for the preparation of model catalyst systems. Various sizes of well-defined Co_3O_4 nanoparticles were synthesised separately and supported onto exfoliated graphite powder, that is graphite after solvent-assisted exfoliation *via* ultrasonication resulting in thinner flakes with increased specific surface area. The stability of the supported nanoparticles during reduction to metallic cobalt in H_2 was monitored *in situ* by means of X-ray diffraction and smaller crystallite sizes were found to be harder to reduce than their larger counterparts. A low cobalt loading of 1 wt% was required to avoid aggregates in the parent catalyst, and this allowed for the preparation of supported cobalt nanoparticles which were resistant to sintering at reduction temperatures below 370 °C. The developed model catalysts are ideally suited for sintering studies of isolated nano-sized cobalt particles as the graphitic support material does not provide distinct metal–support interaction. Furthermore, the differently sized cobaltous particles in the various model systems render possible studies on structural dependencies of activity, selectivity, and deactivation in cobalt oxide or cobalt catalysed reactions.

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1 Introduction

Traditional support materials have mostly been regarded as inert, but more recently even common metal oxide carriers have been shown to exhibit a strong effect on the catalytic behaviour^{1–4} and on the stability of the active phase specifically in the case of nano-sized material.^{5–9} These supports typically anchor the nanoparticles *via* metal–support interactions sometimes even forming mixed phases.^{10–14} On the one hand, such an interaction is desired as it increases the stability of the nanoparticles under reaction conditions, *e.g.* against sintering, on the other hand it is typically associated with a loss in active phase. Pristine carbon does not exhibit such stabilising properties, but may be functionalised with oxygen or nitrogen containing groups.^{15–17} However, even functionalised carbon shows a relative high chemical inertness except for the exposure to O_2

at increased temperatures which results in combustion.¹⁷ Carbon supports are seldom applied in industry^{17,18} due to their lower mechanical strength compared to Al_2O_3 or SiO_2 supports.¹⁸ However, their inert character makes carbon supports the carrier of choice for fundamental studies on the behaviour and stability of relatively isolated nanoparticles.^{19–22}

Graphitic carbon materials comprise multiple layers of sp^2 -hybridised carbon atoms in two-dimensional planes. The atoms are arranged in the classic honeycomb like structure of graphene and the single layers are typically separated by 3.41 Å and stacked in an ABAB order.²³ The amount of stacked layers can be decreased *via* exfoliation techniques allowing the preparation of few-layer graphite or even graphene, a single two-dimensional plane of carbon atoms.²⁴ The exfoliation of layered structures requires the overcoming of the weak van der Waals attractions between adjacent layers,²³ which may be achieved for graphitic materials *via* immersion in solvents featuring a surface tension of approximately 40 mJ m^{-2} . Dispersion of graphitic material in such solvents results in a reduced interaction of adjacent layers.^{23,25} Exfoliation of immersed graphite may then be achieved by applying shearing forces *via* ball-milling²⁶ or ultrasonication.^{27–29} The latter technique may yield up to 4 wt% of graphene.²⁸ However, pure few-layer graphite with up to 28% single-layer graphene may be achieved when applying a size-

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selective centrifugation.²⁷ A second, chemical approach decreases the van der Waals forces by expanding the distance between the layers allowing for simplified exfoliation. The chemical conversion of graphite to graphene *via* the preparation of intercalated graphite oxide according to a modification of Hummers' method³⁰ with subsequent exfoliation into graphene oxide³¹ and reduction to graphene (commonly referred to as reduced graphene oxide rGO)³² is one of the most widely applied methods due to its low cost in combination with considerable yields of the desired product.^{24,33,34} However, rGO differs strongly from pristine graphene obtained from aforementioned syntheses. The harsh oxidising and reducing conditions during the synthesis of rGO introduce considerable amounts of defects such as hydroxyl and epoxy groups on sp³-hybridised carbon on the basal plane disrupting its electronic properties.^{24,33} Defective sites originate from the incomplete reduction of oxidised tetrahedral sp³ to planar sp² carbon atoms.³³ However, modification of the synthesis may allow for the preparation of rGO with decreased concentrations of residual functional groups.³⁵ Aside from mechanical exfoliation of graphitic precursors with the aid of suitable solvents or *via* intercalation of small molecules, a variety of chemical vapour deposition (CVD) techniques such as thermal CVD, plasma enhanced CVD, or a thermal decomposition on substrates are applied for the synthesis of graphene.^{17,24}

The performance of carbon as catalyst or support material is determined by the surface chemistry of the carbon material.¹⁵ Functionalisation of carbon describes the introduction of functional groups such as carboxylic acids or carbonates on the carbon surface,¹⁵ which allow for an electronic interaction with nanoparticles, *e.g.* in heterogeneous catalysts. The interaction is more pronounced than for the pristine, clean surface and hence a higher stability against sintering, but also a lower reducibility, can be expected. Functionalisation techniques targeting the introduction of oxygen containing groups include acidic treatments in nitric, sulfuric, and/or phosphoric acid, as well as oxidative treatments with air, oxygen, nitrous oxide, potassium permanganate, or hydrogen peroxide.^{15,36–39} Functionalisation can also represent an artefact of the preparation of the carbon material, for example, aforementioned rGO contains various functional groups on the edges and the planar surfaces due to the harsh conditions applied during oxidation to graphitic oxide and reduction after exfoliation.^{24,33} In the present study pristine graphite is exfoliated and applied as novel support material in order to prepare rather isolated Co₃O₄ and, upon reduction in H₂, metallic Co crystallites. The developed model system may allow for the characterisation of the physicochemical properties of isolated crystallites. For example, the stability of the crystallites may be analysed under conditions applied in catalytic processes with a minimised effect of the support material.

2 Experimental

2.1 Chemicals

Acetone, aqueous ammonia solution (25 wt%), and ethanol were purchased from Kimix (South Africa). Benzyl alcohol,

cobalt(II) acetate tetrahydrate, synthetic graphite powder, and 1-methyl-2-pyrrolidinone were obtained from Sigma-Aldrich. Kaylaw graphite powder is a chemically pure carbon material and was supplied by Electrochem (South Africa).

2.2 Exfoliation of graphite

Kaylaw graphite powder was exfoliated to increase its surface area.²⁷ 1-Methyl-2-pyrrolidinone (NMP) facilitates the exfoliation process as its surface energy is very close to the one of graphite.²⁸ An amount of 3300 mg graphite powder was sonicated in 1 L NMP for 12 h.²⁸ The obtained suspension was centrifuged at 500 rpm for 45 min to separate non-exfoliated and larger flakes of graphite.²⁷ The exfoliated graphite (EG) in the remaining supernatant was collected *via* centrifugation at 7000 rpm for 2 h, subsequently washed several times with acetone, and dried at 80 °C in an oven.

2.3 Synthesis of Co₃O₄ nanoparticles

Co₃O₄ nanoparticles were synthesised *via* a surfactant-free, non-aqueous heat treatment of dissolved cobalt acetate in benzyl alcohol in the presence of ammonium hydroxide.^{9,14,22,40,41} Cobalt(II) acetate tetrahydrate was dissolved in benzyl alcohol under magnetic stirring at 500 rpm in a round bottom flask (Table 1). After 2 h of stirring, a 25 wt% aqueous ammonium hydroxide solution was added dropwise to the pink to purple solution forming a brown emulsion. Once the addition of ammonium hydroxide was completed, the flask including the stirrer bar was immediately transferred to a preheated oil bath of a rotary evaporation set-up and heated for a total of 3 h at 165 °C. Air was bubbled through the reaction emulsion making use of the reduced pressure in the rotary flask (900 mbar) in order to ensure adequate mixing of the reaction emulsion.^{9,41}

After cooling-down to room temperature, the volume was tripled with diethyl ether and the mixture was centrifuged for 1 hour at 7000 rpm. The centrifugate containing the nanoparticles was re-dispersed in ethanol and washed at least three times with acetone until a clear supernatant was obtained.^{40,41}

2.4 Supporting of nanoparticles

Co₃O₄ crystallites were dispersed in ethanol in an ultrasonic bath until all nanoparticles were in dispersion (>4 h). Exfoliated graphite was sonicated for 4 h in ethanol after which the dispersion of Co₃O₄ nanoparticles in ethanol was added dropwise targeting the desired loading of metallic Co. After sonicating for another 4 h, the dispersion was transferred to a rotary evaporator and further mixed for 1 h at 240 rpm and 80 °C. Subsequently, ethanol was evaporated from the parent sample at 462 mbar and the sample was dried at 80 °C.^{9,14,22}

2.5 Characterisation

Samples were analysed *via* transmission electron microscopy (TEM) in a Tecnai F20 microscope (Philips) equipped with a field emission gun and operated at 200 kV (Gatan). Images were collected with a US4000 4k × 4k CCD camera (Gatan). Nanoparticles were dispersed in ethanol *via* ultrasonication for



Table 1 Applied parameters in sol–gel syntheses of cobalt(II,III) oxide nanoparticles *via* the benzyl alcohol route

Sample	Benzyl alcohol/mL	Co(C ₂ H ₃ O ₂) ₂ ·4H ₂ O/mg	NH ₄ OH _{aq} (25 wt%)/mL
Co_2.8	140	800	35
Co_3.5	140	400	35
Co_4.9	70	1600	70
Co_7.6	140	3200	140

30 min prior to deposition onto carbon-coated copper grids. In contrast, supported nanoparticles were mixed with acetone and dispersed in the ultrasonication bath for 1 min (parent samples) or 3 min (reduced samples) in order to prevent separation of the nanoparticles and the support. Number-based size distributions were determined by measuring the size of over 500 nanoparticles using the open-source ImageJ 1.51a software package.⁴² The distribution of the nanoparticles was converted to a volume-based size distribution to allow for comparison with volume-based techniques such as X-ray diffraction (XRD). The number mean sizes ($d_{n,TEM}$), the volume-mean sizes ($d_{V,TEM}$), and the relative volume-based standard deviations ($\sigma_{V,TEM}$) were calculated based on the measured sizes d_i (eqn (1)–(3)).

$$d_{n,TEM} = \frac{\sum_{i=1}^N d_i}{N} \quad (1)$$

$$d_{V,TEM} = \frac{\sum_{i=1}^N n_i d_i^4}{\sum_{i=1}^N n_i d_i^3} \quad (2)$$

$$\sigma_{V,TEM} = \sqrt{\frac{\sum_{i=1}^N (n_i d_i^3 (d_i - d_{V,TEM})^2)}{\frac{N-1}{N} \sum_{i=1}^N n_i d_i^3}} / d_{V,TEM} \times 100\% \quad (3)$$

where N is the number of measured particles, and n_i are the particular fractions.

Analysis of samples *via* Raman spectroscopy was conducted in an inVia Raman microscope (Renishaw) with a wavelength of the laser of 532 nm. The Raman laser was fitted with a compact fibre optics probe (Renishaw). Five repetitions were taken at 40 mW laser power and an exposure time of 30 s. The intensity ratio of the D and G band of graphitic samples allows for an estimation of the average in-plane crystallite size of the graphene layers (eqn (4)).⁴³

$$L_a = 2.4 \times 10^{-10} \lambda^4 \left(\frac{I_D}{I_G} \right)^{-1} \quad (4)$$

where λ is the wavelength of the Raman laser and I_D and I_G are the intensities of the D band and G band, respectively.

The surface areas of graphitic samples were analysed *via* physisorption according to the Brunauer–Emmett–Teller (BET) method. Physisorption was conducted using a TriStar II 3020 (Micromeritics) with N₂ as analysis adsorptive and a degassing temperature of 200 °C.

Thermogravimetric analysis (TGA) of graphitic support material was conducted in a SDT 650 DSC/TGA simultaneous thermal analyser (TA Instruments). An amount of 8 to 12 mg of material was loaded into a 90 μ L alumina crucible and heated from 50 to 900 °C at 5 °C min⁻¹ in N₂ to decompose functional groups associated with the surface of carbon. The measured weight loss then corresponds to such functional groups.^{15,44} In another analysis to determine the Co loading, the temperature was ramped from 120 to 900 °C at 10 °C min⁻¹ with a final holding time at 900 °C of 60 min in air. The residual mass corresponds to CoO being the stable cobalt oxide at 900 °C,⁴⁵ which allows for the calculation of the Co loading.

Conventional XRD was conducted at 35 kV and 40 mA in a D8 Advance X-ray diffractometer (Bruker AXS), equipped with a cobalt source ($\lambda_{K\alpha 1} = 0.178897$ nm; slit width = 1.0 mm) and a LYNXEYE XE position sensitive detector (Bruker AXS) from 20–120° at a step size of 0.025° with an exposure time of 1 s per step. XRD analysis was also conducted at the crystallography beamline I711 of the MAX II synchrotron⁴⁶ (MAX IV Laboratory, Lund, Sweden) with a wavelength of 0.9941 Å and a slit width of 0.8 mm (see ESI and Fig. S.1 for experimental set-up†). Obtained XRD patterns were compared to reference patterns of the Powder Diffraction File of the International Centre for Diffraction Data (ICDD; PDF-2 Release 2008;⁴⁷ Co₃O₄: 00-043-1003, cubic CoO: 01-071-1178, fcc-Co: 00-015-0806, hcp-Co: 01-071-4239, graphite: 03-065-6212). Volume-mean crystallite sizes and weight fractions of the particular phases were determined by Rietveld refinement of the XRD patterns (TOPAS 5, Bruker AXS).⁴⁸ The instrumental line broadening was modelled *via* fitting the pattern of corundum. Diffraction line broadening analysis *via* the Scherrer equation⁴⁹ (Co₃O₄: 311 diffraction) with correction for the instrumental line broadening and a shape factor of 0.9 was applied as a secondary technique to estimate the crystallite size of the as prepared nanoparticles *via* XRD.⁵⁰

2.6 In situ X-ray diffraction (XRD)

The used sample presentation device for *in situ* XRD was developed at the University of Cape Town^{51,52} and comprises an infra-red heater and a quartz capillary (outer diameter 1.0 mm, wall thickness 0.01 mm, length 75 mm; Hilgenberg GmbH), which is mounted on the sample stage *via* rubber ferrules and screwing caps. A K-type thermocouple detects the temperature directly in the isothermal zone of the 1.5 cm long catalyst bed in the capillary resembling a fixed-bed reactor.⁵² A Kapton polyimide film (DuPont) is assembled as a cover, which improves temperature control and greatly extends the axial isothermal zone. The catalyst bed is held in place by glass wool plugs.



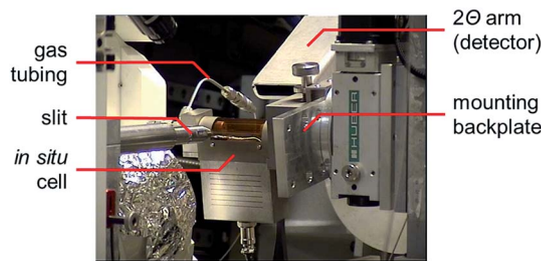


Fig. 1 Photograph of the *in situ* sample presentation device mounted on the experimental table of beamline I711 of the MAX II synchrotron.

The cell was applied in *in situ* XRD reduction studies on a Bruker D8 Advance equipped with a Co source ($\lambda_{\text{CoK}\alpha 1} = 1.78897 \text{ \AA}$). The device can be easily attached to the goniometer of the commercial X-ray diffractometer. The selected step size within the 2θ -window of choice was 0.062° with an exposure time of 0.8 s. In addition, the cell was mounted to the experimental table of beamline I711 (Fig. 1) of the MAX II synchrotron⁴⁶ (MAX IV Laboratory, Lund, Sweden). A wavelength of 0.9941 \AA and a slit width of 0.8 mm were applied for *in situ* reduction studies with a high energy beam.

Exfoliated graphite as the support material of choice was modelled using Rietveld refinement with three separate graphite structures in a modified partial or not known crystal structure (PONKCS) approach (see ESI and Fig. S.2–S.6 for detailed description[†]).⁵³ The thermal expansion of graphitic samples perpendicular to the basal plane^{54,55} was included applying a fitted linear dependency of the c lattice parameter with the temperature (eqn (S.1)–(S.6)[†]).

In a typical *in situ* XRD study, 2–5 mg of the supported parent catalyst were loaded into the quartz capillary. The temperature was ramped in pure H_2 from 50°C to a selected holding temperature at a heating rate of 1°C min^{-1} . A space velocity of $10 \text{ mL min}^{-1} \text{ mg}_{\text{catalyst}}^{-1}$ was applied in all experiments. XRD patterns were obtained every 5 min during the ramp and every 5–20 min during a holding time. After the reduction, the screwing caps are slowly loosened to allow for diffusion of air into the capillary, which can be expected to passivate the catalysts within minutes.⁵⁶

The obtained *in situ* XRD patterns were analysed with the help of a MATLAB (The Mathworks Inc.)⁵⁷ algorithm.⁵⁸ It allows for the automated analysis of all XRD patterns of one experiment by means of Rietveld refinement using the TOPAS kernel. Aside from classical XRD parameters, the output further provides the refined patterns and the single patterns of the particular structures. The algorithm was modified to a small extent in order to accommodate the developed PONKCS approach for the EG support.

3 Results and discussion

3.1 Characterisation of exfoliated graphite

X-ray diffraction (XRD) patterns of graphite samples are dominated by the diffraction of the (002) plane at 30.9° ($\lambda_{\text{Co,K}\alpha 1} = 1.78897 \text{ \AA}$; Fig. 2a). This becomes even more pronounced for

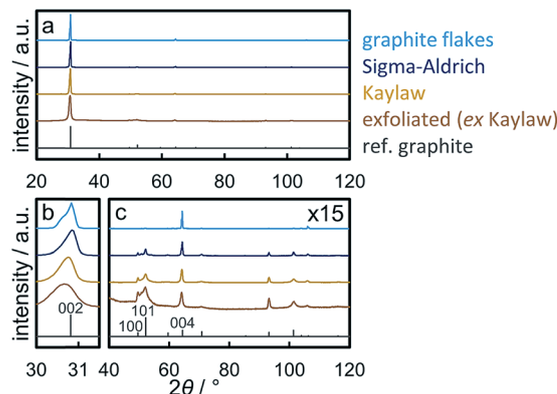


Fig. 2 (a) Normalised X-ray diffraction patterns, (b) enlarged (002) reflections, and (c) the patterns above 40° with 15 times increased intensities of graphitic materials with a reference pattern for graphite.

thick samples with an extended length perpendicular to the basal plane,⁵⁹ *i.e.* with high stacking numbers such as natural graphite flakes. The shape of the (002) reflection depends on the thickness of the graphite sample. Thinner graphite samples result in a broader peak with a maximum at lower diffraction angles due to the increased contribution of intercalated planes on top/bottom of graphitic flakes.^{59,60} XRD analysis of natural graphite flakes exhibits this dependency as the (002) reflection is bimodal indicating a wide range of stacking numbers (Fig. 2b). Application of the Scherrer equation⁴⁹ on the (002) diffraction line allows for an estimation of the average number of layers.^{33,60} In the present study, this method was extended to Rietveld refinement of the patterns from $28\text{--}33^\circ$ applying two separate graphite structures in order to obtain a range of the number of layers based on the calculated crystallite sizes (Table 2). As expected, size analysis of synthetic graphite powder (Sigma-Aldrich) and natural graphite flakes results in the former having less layers. Kaylaw graphite powder, a chemically pure carbon additive for lubricants, shows the broadest (002) reflection. Another indicator for the thickness of a graphite sample is the ratio of the reflection intensity of the interplanar distances (002) or (004) and the intraplanar (100) plane, which increases with the stacking number. This can be demonstrated by a comparison of the ratios for natural graphite flakes and commercially synthesised graphite powder (Table 2; Fig. 2c). The trend in the ratios is inversely proportional to the BET surface area (Fig. S.7[†]), which can be expected to increase with lower stacking numbers.

Exfoliation of Kaylaw graphite powder *via* ultrasonication in NMP^{27,28} and size selection *via* centrifugation²⁹ almost doubled the surface area of the graphite phase (Fig. S.7[†]) while decreasing the calculated average range in the number of layers accordingly by approximately 50% (Table 2). However, the yield upon ultrasonication in fresh, anhydrous NMP and size-selective separation of the fraction of thicker graphite *via* centrifugation is only 5 wt% and decreases with the number of recycling steps of the valuable solvent NMP, presumably due to contamination with water from the ambient atmosphere.²⁸ Analysis of exfoliated graphite (EG) by means of XRD showed



Table 2 Characteristics of reflections in X-ray diffractograms with an estimated number of layers, as well as the surface area of graphitic carbon samples

Sample	Intensity ratio		FWHM (002) ^o	Approx. layers ^a	BET surface area/m ² g ⁻¹
	(002)/(100)	(004)/(100)			
Natural graphite flakes	3225.2	144.7	0.254	270–664	1.0 ± 0.1
Graphite powder (Sigma-Aldrich)	136.3	4.8	0.331	136–400	8.1 ± 0.1
Kaylaw graphite powder	76.0	2.7	0.414	107–207	25.8 ± 0.1
Exfoliated graphite (<i>ex</i> Kaylaw)	22.1	1.0	0.658	58–100	47.8 ± 0.2

^a Estimation of the average number of layers from the crystallite size obtained *via* Rietveld refinement of the (002) reflection with two graphite structures.

decreased ratios of the interplanar reflections relative to the intraplanar reflections confirming a decreased thickness upon exfoliation. Less than 100 layers were estimated from the analysis of the (002) diffraction (Table 2). The actual stacking number may be even lower due to partial re-stacking of thinner few-layer graphite (Fig. S.8†) during the drying procedure,²⁴ which is also indicated by the shift to lower diffraction angles due to an increased *d*-spacing of imperfect stackings.

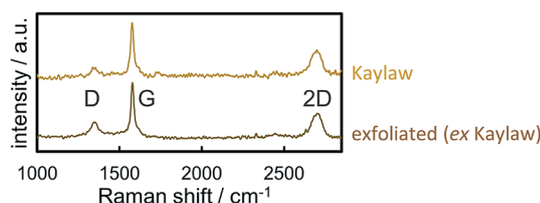
Raman spectra of carbon materials show only few prominent features. However, Raman spectroscopy has become the standard for characterisation and analysis in carbon research.⁶¹ The G band (G for graphite) at approximately 1560 cm⁻¹ for visual excitation is typically assigned to the bond stretching of all pairs of sp²-hybridised carbon atoms in rings or chains and is therefore associated to the graphitic structure of pristine graphite powder.^{61–63} The breathing modes of sp² atoms in carbon hexagons result in the D band (D for disorder or defect) at 1360 cm⁻¹,^{61–64} which is not observed in the centre of a graphite sheets, yet on the edge.⁶⁵ In regard to carbon as catalyst support, the D band generally represents the functionalisation of the carbon carrier due to induced defects. The Raman spectra of Kaylaw graphite powder before and after exfoliation display similar features (Fig. 3). The ratio of the intensities of the D band over the G band (*I_D/I_G*) is inversely proportional to the in-plane crystallite size (*L_a*)^{43,64,66} and increases from 0.18 to 0.29, *i.e.* exfoliation induces a higher defect concentration as more carbon atoms are associated to the edge for a smaller in-plane dimension.⁶⁴ An estimation of the in-plane crystallite size with the intensity ratio of both prominent bands according to Cançado *et al.*⁴³ (eqn (4)) results in 106.8 nm for pristine Kaylaw graphite powder and 66.3 nm after exfoliation. No significant differences can be identified in the 2D band at approximately 2710 cm⁻¹,⁶⁵ formerly referred to

as G' band.⁶⁷ The broad peak of said band indicates a 1 : 1 intensity ratio of the 2D₁ and 2D₂ band suggesting relatively thin graphite flakes,⁶¹ which is supported by XRD analysis (Table 2).

Characterisation of EG and the parent Kaylaw graphite powder under an inert N₂ atmosphere by means of TGA (Fig. S.9a†) strongly supports the conclusions drawn from the analysis of Raman spectra. The weight loss during exposure to increased temperatures can be assigned to the decomposition of particular functional groups on the surface of carbon materials.^{15,44,68} When normalised to the available surface area as determined *via* BET analysis, EG and Kaylaw graphite powder show comparably low concentrations of functional groups (Fig. S.9a†). No more functional groups decompose on the graphitic samples at temperatures exceeding 500 °C. The low degree of functionalisation (weight loss: 0.06% (m² g⁻¹)⁻¹) can be demonstrated when compared to the normalised weight loss of solid carbon spheres,⁶⁹ which is more than 5 times the loss of Kaylaw graphite powder at 500 °C. In addition, significant amounts of functional groups and/or amorphous carbon are converted at increased temperatures. The weight loss due to decomposition of functional groups, when normalised to the BET surface area, is also lower than reported for as-grown multiwalled carbon nanotubes (CNTs; 0.11% (m² g⁻¹)⁻¹).⁶⁸ The weight fraction in said material has been demonstrated to increase to approximately 0.22% (m² g⁻¹)⁻¹ by applying an acidic treatment in HNO₃.⁶⁸ In the present study, TGA was also applied to verify the chemical purity of Kaylaw graphite powder and resulted in a full conversion upon oxidation in air at 900 °C (Fig. S.9b†).

3.2 Characterisation of Co₃O₄ nanoparticles

Four distinct sizes of monodisperse Co₃O₄ crystallites were synthesised *via* the benzyl alcohol-route.^{9,40,41} TEM analysis reveals the high dispersibility of the nanoparticles which are nicely spread out over the grid (Fig. 4a–d). All four sizes show narrow size distributions (Fig. 4e) with relative standard deviations ranging from 16 to 29% (Table S.1†). Analysis by means of XRD confirmed the successful preparation of single-phase Co₃O₄ nanoparticles of high crystallinity (Fig. 4f). All diffractions can be assigned to a Co₃O₄ reference pattern, while no other phases may be detected. Comparison of the width of the diffractions (*e.g.* the most prominent (311) diffraction at 43°)

**Fig. 3** Raman spectra of pristine and exfoliated Kaylaw graphite powder displaying carbon characteristic bands.

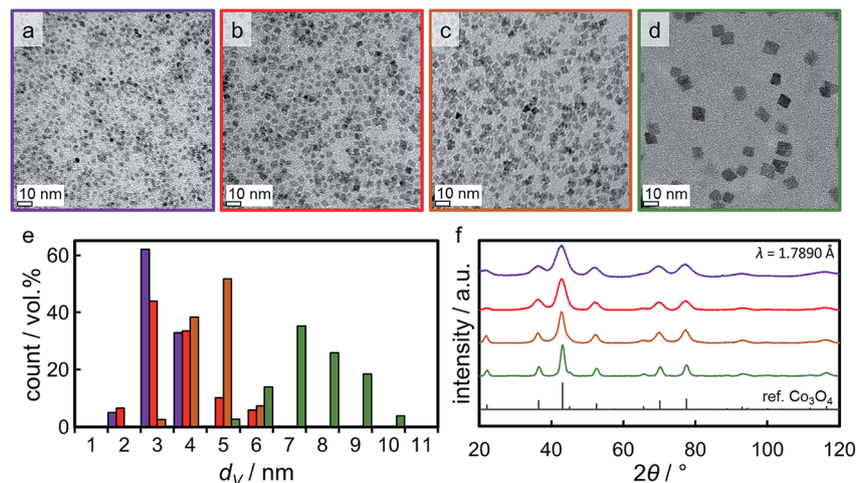


Fig. 4 (a–d) Transmission electron micrographs of the Co₃O₄ nanoparticles obtained in synthesis Co_{2.8}, Co_{3.5}, Co_{4.9}, and Co_{7.6}, respectively, together with (e) the particular volume-based size distributions and (f) X-ray diffractograms with a reference pattern for Co₃O₄. In part reproduced from ref. 9 with permission from The Royal Society of Chemistry.

indicates significant differences in crystallite size. Indeed, Rietveld refinement results in volume-mean crystallite sizes of 2.8, 3.5, 4.9, and 7.6 nm (denoted Co_{2.8}, Co_{3.5}, Co_{4.9}, and Co_{7.6}, respectively), which were confirmed *via* application of the Scherrer equation⁴⁹ and TEM analysis (Table S.1†).

3.3 Size-dependent reduction of 5 wt% Co/EG model catalysts

A couple of experimental procedures can be employed to limit the extent of sintering during the activation of metal oxide-based catalysts in H₂. Minimisation of the reduction temperature and hence the thermal strain is the most obvious one as the sintering tendency strongly correlates with temperature.⁷⁰ Additional parameters of sintering are the composition of reactive atmosphere,^{71–74} the particle size distribution,^{75–77} and the nature/reactivity of the support material.^{76,78,79} The latter is arguably the most important factor in heterogeneous catalysts as unsupported metal oxide nanoparticles experience a severe increase in size without the stabilising effect of a carrier (Fig. S.10 and S.11†). In comparison to oxidic supports, EG and graphite in general exhibit only little functionalisation of the surface.^{24,80} The majority of the functional groups present can be expected to be localised at the edges of the single planes within the graphite structure, which only make up a small fraction of the overall surface area.

The separately synthesised nanoparticles were supported on EG in order to assess the suitability of such novel model systems for the isolated characterisation of metallic Co crystallites. The physicochemical surface properties of the support can be expected to induce differently pronounced interactions with the nanoparticles, which affect their dispersion over the support during the supporting procedure. Three model catalysts with a targeted 5 wt% Co loading were prepared and analysed by means of TGA and TEM. TGA resulted in actual Co loadings in the range of 4.4 to 5.0 wt% (Fig. S.12†), *i.e.* the three samples showed a similar concentration of Co. The thin flakes of

exfoliated graphite and the large density difference between the support material and the Co₃O₄ nanoparticles allow for a clear distinction between the nanoparticles and the carrier (Fig. 5). Comparison of the parent catalysts consisting of the Co₃O₄ nanoparticles Co_{2.8}, Co_{4.9}, and Co_{7.6} supported on EG demonstrates the different dispersion levels of the oxidic cobalt phase over the available surface area of the carrier. The larger nanoparticles in the 5 wt% Co_{7.6}/EG and Co_{4.9}/EG samples are well distributed and only small aggregates are identified. A high dispersion over the surface area of the support was achieved for the smaller nanoparticles, but the surface of the EG appears to be crowded due to the large number of small crystallites required for the 5 wt% loading.

The reduction behaviour of the three model catalysts in H₂ was studied from 50 °C up to the desired reduction temperature *via in situ* XRD. The relatively low energy level of conventional X-ray radiation sources in combination with the overlap of the Co-related reflections with the support pattern result in a lower detection limit of the cobaltous phase of approximately 5 wt%. The largest Co₃O₄ nanoparticles in sample Co_{7.6}/EG feature, as expected, narrow diffractions (most intense reflection at 43.0°) of relatively high intensity and hence a high signal-to-noise ratio, which in turn results in distinct *in situ* XRD patterns (Fig. 6). A successful application of the modified PONKCS analysis⁵³ of EG is indicated by the rather constant pattern of the summed-up graphite structures during the reduction. A first reduction of the Co₃O₄ phase to CoO (reflections at 42.7 and 49.7°) was observed at 235 °C during the 1 °C min⁻¹ ramp to 350 °C. The subsequent formation of fcc-Co occurred rapidly from 310 °C onwards and was delayed when compared to unsupported nanoparticles (245 °C; Fig. S.10 and S.11†). Metallic fcc-Co (reflection at 51.8°) is the only cobaltous phase identified at temperatures exceeding 340 °C.

Even though the low temperature (<400 °C) allotrope of metallic Co is reported to be hcp-Co for bulk phases,⁸¹ the formation of single-phase fcc-Co without a (detectable) fraction



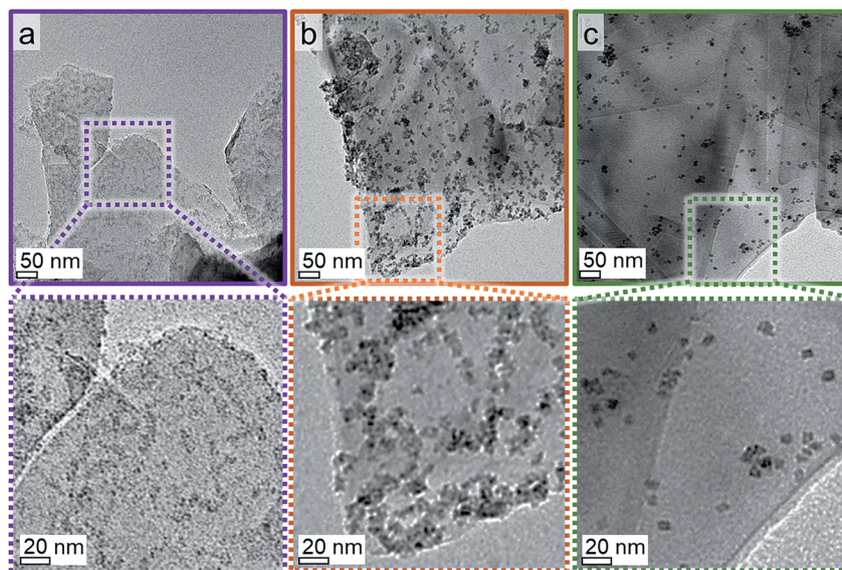


Fig. 5 Transmission electron micrographs of 5 wt% (a) Co_{2.8}, (b) Co_{4.9}, and (c) Co_{7.6} on exfoliated graphite with magnified insets.

of hcp-Co is in good agreement with literature.^{82,83} Metallic Co nanoparticles, prepared *via* neutron sputtering at ambient temperature, are reported to consist of pure fcc-Co for crystallite sizes below 20 nm, while the concentration of the hexagonal allotrope increases significantly with size to approximately 85% in nanoparticles above 40 nm.⁸² However, annealing of 30 nm Co nanoparticles with an initial concentration of hcp-Co of 78% at different temperatures (in the range of 200–500 °C) resulted in an increase of the concentration of hcp-Co at 300 °C, while said phase was transformed to fcc-Co at higher temperatures. A

pure fcc-Co phase was detected at 500 °C.⁸² A preferential formation of fcc-Co in nanoparticles below 20 nm was confirmed for Al₂O₃-supported Co nanoparticles.⁸³ Reduction of several sizes (3–10 nm) of Co₃O₄ to metallic Co in H₂ at 450 °C, similar conditions as applied in this study, resulted in single-phase fcc-Co as well. More recently, a highly intergrown fcc-hcp phase has been reported in cobalt nanoparticles supported on TiO₂ as well.⁸⁴

Sintering of the nanoparticles during reduction of sample Co_{7.6}/EG in H₂ was identified (Fig. 7a) even though the

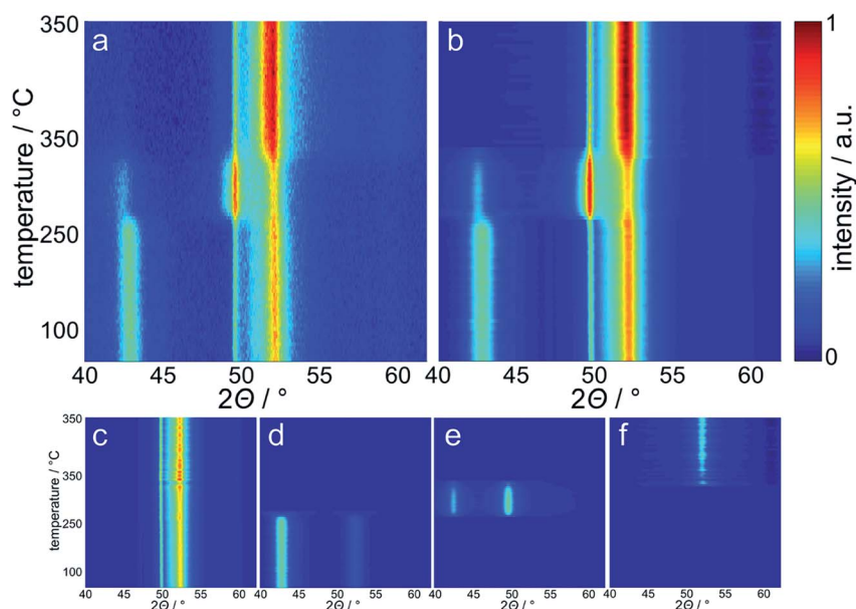


Fig. 6 Top-view of (a) the experimentally obtained *in situ* X-ray diffraction patterns during reduction of 5 wt% Co_{7.6} on exfoliated graphite in hydrogen at 350 °C for 5 h, (b) calculated patterns obtained *via* Rietveld refinement with the background being deducted and the corresponding fitted contributions of (c) the summed-up exfoliated graphite phases from PONKCS analysis, (d) Co₃O₄, (e) cubic CoO, and (f) fcc-Co.



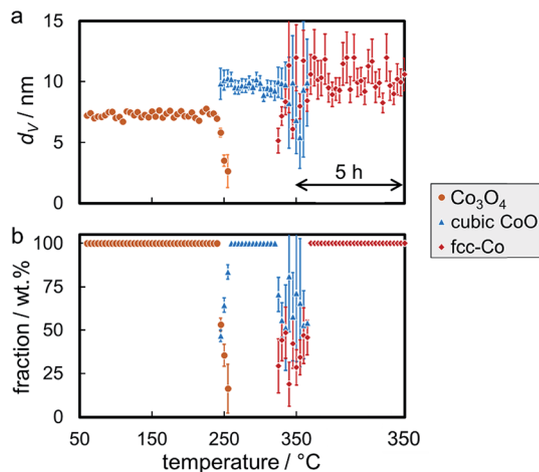


Fig. 7 (a) Volume-mean crystallite sizes and (b) weight fractions with the particular errors of the phases as obtained *via* Rietveld refinement of *in situ* X-ray diffraction patterns during reduction of 5 wt% Co_{7.6} on exfoliated graphite in hydrogen at 350 °C for 5 h.

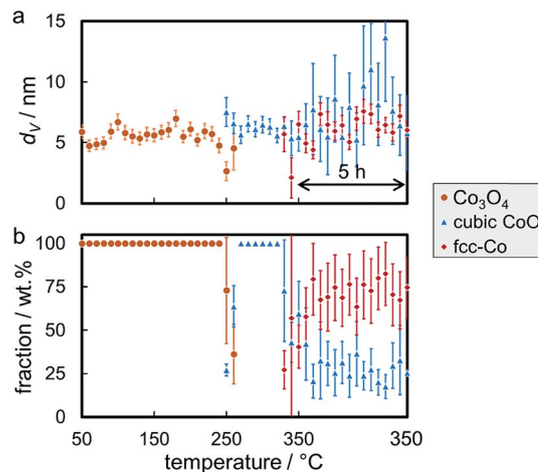


Fig. 8 (a) Volume-mean crystallite sizes and (b) weight fractions with the particular errors of the phases as obtained *via* Rietveld refinement of *in situ* X-ray diffraction patterns during reduction of 5 wt% Co_{4.9} on exfoliated graphite in hydrogen at 350 °C for 5 h.

nanoparticles were expected to be rather isolated with relatively large inter-particle distances. Seemingly, sintering first occurred during reduction to CoO and, to a smaller extent, during the formation of fcc-Co. The presence of aggregated crystallites in the parent sample due to the relatively high Co loading of 5 wt% and consequential crowding of the support surface may explain the observed crystallite growth. Further, carbon supports are reported to provide only little stabilisation of nanoparticles when compared to metal oxide carriers,^{19–21,71} which may allow for pronounced sintering *via* crystallite migration and collision. It has to be noted that the accuracy of the obtained crystallite sizes for fcc-Co is limited due to the overlap of the main diffraction at 51.8° with reflections of the graphitic support. Analysis of the composition of the cobaltous phases during the reduction clearly exhibits a separated two-step reduction of Co₃O₄ to the metallic Co phase *via* CoO (Fig. 7b), which is in line with literature.^{83,85–88}

Reduction of the 4.9 nm Co₃O₄ nanoparticles in sample Co_{4.9}/EG was conducted at 350 °C with a 5 h holding time. The first reduction step to CoO was observed at 245 °C (Fig. S.13†), which is only 5 °C higher when compared to the larger nanoparticles in sample Co_{7.6}/EG. A subsequent slow reduction to fcc-Co was observed at temperatures exceeding 320 °C, while the volume-mean crystallite size only increased by 0.7 nm (Fig. 8a) lying within the error range of Rietveld refinement. Even though the abstraction of oxygen during reduction of Co₃O₄ goes along with a theoretical decrease of the crystallite size during reduction (Co₃O₄ to CoO: 12%, CoO to metallic Co: 43%), the change in crystallite shape from cubes to most likely spheres is expected to approximately balance the shrinkage in size. Rietveld refinement suggests no further reduction of CoO after 1 h at 350 °C (Fig. 8b) as the CoO reflection at 49.6° can be continuously identified throughout the *in situ* XRD pattern.

The required temperature for the first formation/detection of CoO during reduction of the smallest Co₃O₄ nanoparticles in H₂ (Co_{2.8}/EG) was 250 °C (Fig. S.14†) and hence slightly increased

when compared to the larger sizes (Co_{2.8}/EG: 245 °C; Co_{7.6}/EG: 240 °C). This observation may be explained by the small crystallite size resulting in broad and hence less intense reflections or potentially exhibits a weak size-dependency of the reduction behaviour of Co₃O₄ crystallites. Metallic fcc-Co was only formed at approximately 325 °C. Hence, the reduction of CoO to fcc-Co is seemingly hindered for this smallest size as well, which is supported by the continuous identification of the CoO phase *via* the reflection at 49.6°. Sintering is identified during both reduction steps, from Co₃O₄ to CoO and from CoO to fcc-Co (Fig. 9a). However, the sizes of both phases remain smaller than in the samples containing the initially larger crystallites. Rietveld refinement results in a final concentration of fcc-Co of almost 80 wt%, which may be increased further at extended holding times (Fig. 9b).

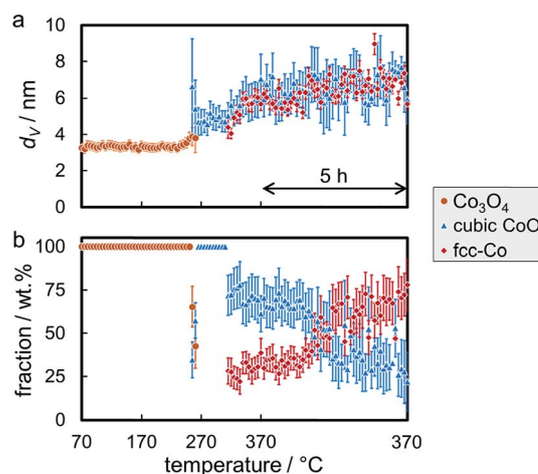


Fig. 9 (a) Volume-mean crystallite sizes and (b) weight fractions with the particular errors of the phases during reduction of 5 wt% Co_{2.8} on exfoliated graphite in hydrogen at 370 °C for 5 h, as obtained *via* Rietveld refinement of the *in situ* X-ray diffraction patterns.



The onset temperatures of the reduction of the supported Co_3O_4 crystallites (240–250 °C) in the three model catalysts are significantly higher than observed for the initial reduction of unsupported crystallites of comparable size (160 °C; Fig. 10, S.10 and S.11†). The identification of sintering, as calculated *via* Rietveld refinement, has been considered in the comparison of the volume-mean crystallite sizes. The second reduction step to metallic Co also requires significantly increased temperatures (305–320 °C *vs.* 245 °C). In addition, the unsupported CoO crystallites reduce rapidly without significant further activation requirement. This may be explained by a H^* spillover process from formed metallic sites to the adjacent oxidic counterpart^{89–91} resulting in an accelerated autocatalytic reduction.^{92–94} Such a mechanism may also be at play during reduction of supported aggregates to metallic Co. However, the differences in the absolute onset temperatures have to be induced by the stabilising effect of the support, which becomes more effective for smaller crystallite sizes. A size-dependent reduction behaviour of cobalt oxide in H_2 has been reported in literature as well.^{95–97} The support limits the tendency of sintering due to the increased inter-particle distance and potentially hinders or limits H^* spillover *via* the surface of the support.⁹¹ Hence, the facile reduction of unsupported crystallites may be driven by the potential decrease in the surface energy^{98,99} *via* a facilitated sintering process during reduction to metallic Co. The hindered reducibility of supported crystallites demonstrates the

stabilising effect of the support on the physicochemical properties of nanoparticles even though carbon supports are expected to provide a low degree of interaction, often quoted as ‘inert’ in literature.^{19–21} Especially the second reduction step forming metallic Co has been reported to be strongly affected by the nature of the support material⁹⁵ and can be expected to shift to increased temperatures for metal oxide carriers.

Post-run analysis of the reduced samples after (passivating) exposure to air by means of TEM confirmed the previously described increases of the volume-mean crystallite size in all three samples with 5 wt% Co loading (Fig. 11). Size analysis of over 500 nanoparticles in the reduced 5 wt% $\text{Co}_{2.8}/\text{EG}$ model catalyst resulted in a volume-mean particle size of 8.5 nm with a standard deviation of 3.9 nm, which is in line with the crystallite sizes obtained *via* Rietveld refinement of the *in situ* XRD patterns (7.5–8 nm). The number-based size distribution shows a broadening of the initial narrow distribution to larger sizes up to 14 nm and resembles to a log-normal size distribution as typically observed after sintering.^{73,100} The size of the sintered crystallites in samples $\text{Co}_{4.9}/\text{EG}$ and $\text{Co}_{7.6}/\text{EG}$ with the initially larger Co_3O_4 crystallites were seemingly underestimated by analysis of the XRD patterns. TEM analysis resulted in volume-mean particle sizes of 10.3 ± 3.2 nm for $\text{Co}_{4.9}/\text{EG}$ and 15.0 ± 4.9 nm for $\text{Co}_{7.6}/\text{EG}$, while Rietveld refinement suggested volume-mean crystallite sizes in the range of 6–7 nm and 9–12 nm, respectively. However, the trend was confirmed and the log-normal-type size distribution indicates the expected occurrence of sintering during reduction of the model catalysts. The low Co loading and the relatively small change of the peak width above 7 nm may be accounted for the underestimation of the sizes by means of XRD.

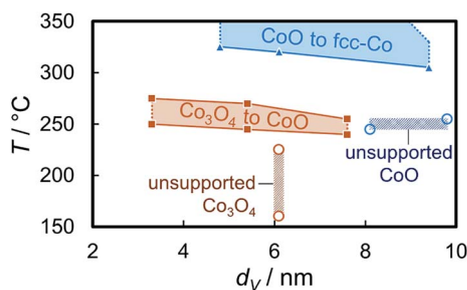


Fig. 10 Temperature ranges for a 1 °C min^{-1} ramp rate of the reduction of exfoliated graphite-supported (5 wt% Co) and unsupported Co_3O_4 crystallites to CoO and the subsequent reduction to metallic Co in hydrogen as observed *via in situ* X-ray diffraction as a function of the volume-mean crystallite size as obtained from Rietveld refinement.

3.4 Reduction behaviour of isolated Co_3O_4 nanoparticles

XRD reduction studies have shown that unsupported Co_3O_4 allows for a facile reduction when compared to supported nanoparticles (Fig. 10). Hence, a significant fraction of the observed reduction of $\text{Co}_3\text{O}_4/\text{EG}$ to metallic crystallites may be attributed to aggregates of nanoparticles, which show a H^* spillover-assisted, autocatalytic reduction behaviour similar to unsupported crystallites and tend to sinter as only little or no stabilisation is provided within the agglomerate. Dispersed nanoparticles exhibit a pronounced stabilisation by the support, which in turn increases the required temperature for

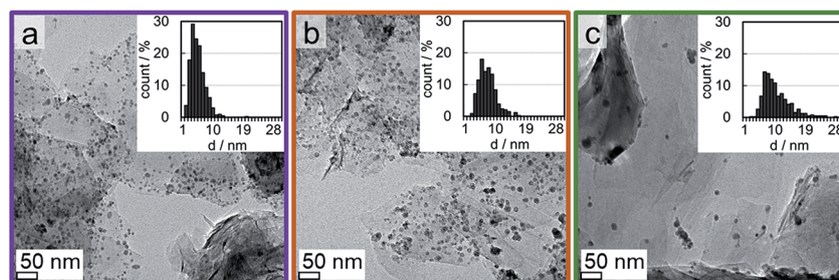


Fig. 11 Transmission electron micrographs and number-based crystallite size distributions of 5 wt% (a) $\text{Co}_{2.8}$, (b) $\text{Co}_{4.9}$, and (c) $\text{Co}_{7.6}$ on exfoliated graphite after reduction in hydrogen at 350 °C during *in situ* X-ray diffraction.



their reduction to metallic Co. The extreme case of an isolated nanoparticle with large inter-particle distances is expected to require the harshest reduction condition, that is the highest temperature. In order to reduce the extent of sintering of the well-defined Co_3O_4 crystallites, the loading of Co has been reduced to 1 wt% and EG has been decorated with several sizes of Co_3O_4 nanoparticles (2.8, 3.5, and 7.6 nm denoted Co_2.8, Co_3.5, and Co_7.5, respectively). TGA resulted in Co loadings of 0.9 to 1.1 wt% (Fig. S.15†). The samples comprising the smaller sizes show a high dispersion of nanoparticles over the support surface (Fig. 12). In contrast, some small aggregates were identified by means of TEM in the samples comprising the larger nanoparticles. However, the fraction of well-dispersed isolated crystallites increased drastically compared to the sample with a 5 wt% Co loading. Synchrotron radiation-based high energy XRD allows for the detection of loadings of 1 wt% of Co on EG as long as the crystallite size is sufficiently large, *i.e.* the XRD patterns feature well-defined reflections. However, the difference between the bare support and the parent Co_2.8/EG sample is marginal prohibiting a viable *in situ* analysis (Fig. S.16†). Especially the main (311) reflection of Co_3O_4 at 23.5° can be identified for the two larger sizes. Hence, *in situ* XRD reduction studies in H_2 were only conducted for the larger supported nanoparticles utilising synchrotron-generated X-rays with the selected wavelength of 0.9941 Å.

A reduction temperature of 330°C was selected for the 1 wt% Co_7.6/EG sample based on the observed rapid reduction of the 5 wt% Co_7.6 nanoparticles on EG at 350°C (Fig. 6). Even though this model catalyst has a relatively large size of Co_3O_4 nanoparticles, most of the features in the top-view of the *in situ* XRD patterns during reduction in H_2 are related to the support (Fig. S.17†). Superimposition of the XRD pattern obtained prior to reduction, the pattern exhibiting the maximum concentration of CoO during the temperature ramp, and the pattern after reduction aids the visual identification of the particular phases

present in the sample (Fig. 13). The (311) plane in Co_3O_4 crystallites results in a diffraction at 23.5° , which shifts to marginally lower diffraction angles upon first reduction due to the (111) plane in CoO. Furthermore, the major (200) diffraction of CoO at 27.0° results in a shoulder at lower angles for the support reflections in the range of $26.8\text{--}30.5^\circ$, which also increases the intensity of the reflection of EG at 26.9° . The (111) diffraction of fcc-Co at 28.0° increases the intensities of the reflections of EG at 28.2° and results in an increased intensity between the major reflections of EG between 27 to 28° .

Rietveld refinement, including the application of a modified POKCS approach⁵³ for EG, results in viable patterns for the particular cobalt phases, as well as for the support (Fig. S.17†). However, no clear transition region could be modelled for the first reduction step, *i.e.* only one of the cobalt oxides may be fitted per pattern even though the experimentally obtained *in situ* XRD patterns suggest a transition period with the simultaneous presence of both oxides. The required temperature of approximately 265°C for the reduction to CoO is significantly higher than for the same nanoparticles in the 5 wt% Co_7.6/EG sample (240°C). This shift to higher temperatures, when compared to a higher loading, demonstrates an improved dispersion and hence a pronounced stabilisation of the nanoparticles by the support. Increased inter-particle distances due to the decreased loading and less aggregates can cause this effect, which was already displayed when comparing the 5 wt% samples to unsupported crystallites (Fig. 10). The intensity of the (111) reflection of CoO peaks at 280°C . Low concentrations of fcc-Co can be fitted from 300°C onwards, but significantly higher concentrations are only obtained at the maximum reduction temperature of 330°C . The onset temperature is comparable to the 5 wt% loaded sample (305°C), however the rate of reduction is slower. As observed for higher loadings, no formation of hcp-Co is detected upon reduction to metallic Co. The obtained crystallite sizes of the cobalt oxides from *in situ*

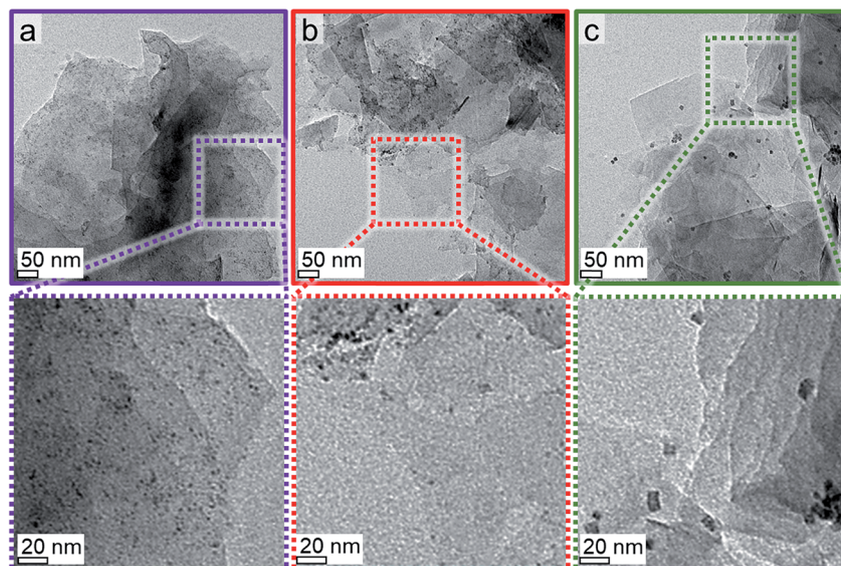


Fig. 12 Transmission electron micrographs of 1 wt% (a) Co_2.8, (b) Co_3.5, and (c) Co_7.6 on exfoliated graphite with magnified insets.



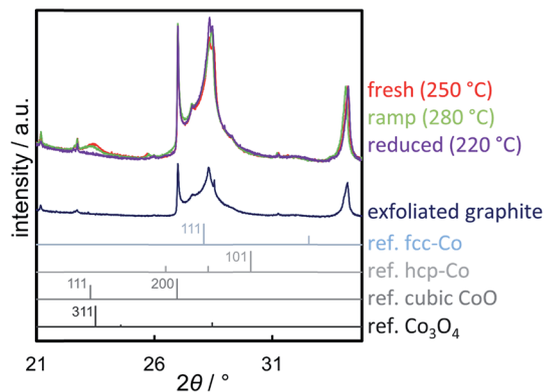


Fig. 13 Synchrotron radiation-based ($\lambda = 0.9941 \text{ \AA}$) X-ray diffraction patterns of 1 wt% Co_{7.6} on exfoliated graphite before reduction in hydrogen (fresh), during the initial ramp at 280 °C displaying the maximum concentration of CoO, and after reduction at 330 °C normalised to the (004) reflection of exfoliated graphite (34.2°) with the pattern of exfoliated graphite and reference patterns for fcc-Co, hcp-Co, cubic CoO, and Co₃O₄.

synchrotron radiation-based XRD of 4.5–5.5 nm represent an underestimation of the actual size seeing the initial size of 7.6 nm (identified *via* TEM and Rietveld refinement of the XRD pattern of the unsupported nanoparticles; Table S.1†). However, no significant increase of the crystallite size is identified during reduction to CoO and fcc-Co indicating the absence of major sintering (Fig. 14a). A continuous reduction of CoO to fcc-Co was modelled at the reduction temperature of 330 °C during the first 2 h of the holding time (Fig. 14b).

The smaller Co₃O₄ nanoparticles in sample Co_{3.5}/EG were reduced at 350 °C, which is sufficiently high to obtain high concentrations of fcc-Co as indicated by the increased intensity of the diffraction of the support at 28° (Fig. S.18 and S.19†). It has to be noted that no measurements were taken during the

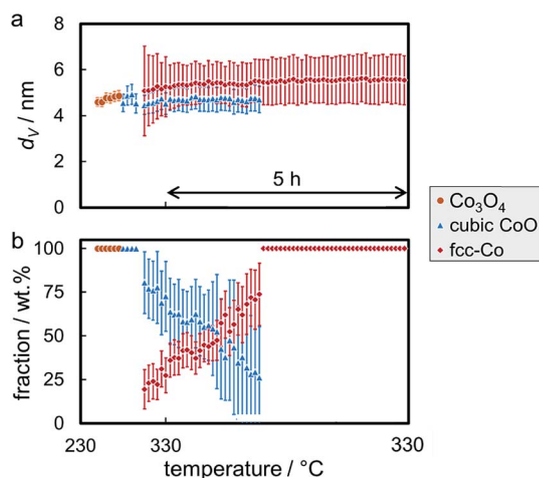


Fig. 14 (a) Volume-mean crystallite sizes and (b) weight fractions with the particular errors of the phases as obtained *via* Rietveld refinement of synchrotron radiation-based *in situ* X-ray diffraction patterns during reduction of 1 wt% Co_{7.6} on exfoliated graphite in hydrogen at 330 °C for 5 h.

first hour at 350 °C due to beam instability. As for the larger size, the transition period from Co₃O₄ to CoO could not be modelled with both oxidic phases in one pattern. The onset temperature of the reduction to CoO was approximately 270 °C and therefore higher than for sample Co_{2.8}/EG with 5 wt% loading (250 °C). The maximum concentration of CoO was observed at an increased temperature when compared to the larger sized crystallites in Co_{7.6}/EG (295 vs. 280 °C). Both observations indicate an improved isolation of the crystallites due to the decreased loading of Co. Metallic fcc-Co was first identified at 310 °C, which is lower than in 5 wt% Co_{2.8}/EG at 325 °C. This observation may be due to the increased sensitivity of the synchrotron radiation-based XRD seeing the relatively low fraction of Co of 10–15 wt% in combination with the slow reduction at said temperatures and the marginally increased crystallite size (Fig. 15). Approximately half of the Co has been reduced to its metallic state after one hour at 350 °C. The initial size of Co₃O₄ crystallites is seemingly underestimated by 1 nm, while the resulting final size of fcc-Co lies within the expected range of 3.5–4 nm. Rietveld refinement further suggests a constant crystallite size of the metallic phase during the last 4 h of the holding time at 350 °C. The absence of reflections of the two oxide phases in the post-reduction XRD pattern indicates a high DOR (Fig. S.18†) even though Rietveld refinement suggests a fraction of 25 wt% CoO (Fig. 15b). Once again, the difficulty in the identification of the Co phase demonstrates the operation close to the detection limits for the studied model system. A decreasing intensity ratio of the support reflections at 27.0° and 28.4° in the superimposed patterns provides further proof for the formation of metallic fcc-Co (Fig. S.19†).

Post-run analysis of the reduced and passivated samples Co_{7.6}/EG and Co_{3.5}/EG with a 1 wt% loading of Co by TEM confirmed the high resistance of both samples against sintering. Volume-mean crystallite sizes of 7.1 ± 1.4 (1 wt% Co_{7.6}/

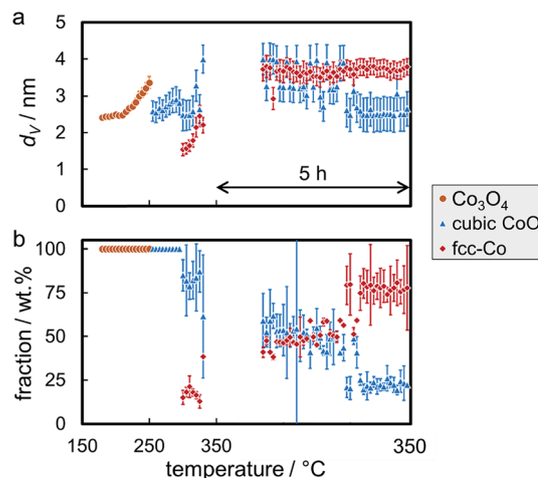


Fig. 15 (a) Volume-mean crystallite sizes and (b) weight fractions with the particular errors of the phases as obtained *via* Rietveld refinement of synchrotron radiation-based *in situ* X-ray diffraction patterns during reduction of 1 wt% Co_{3.5} on exfoliated graphite in hydrogen at 350 °C for 5 h.



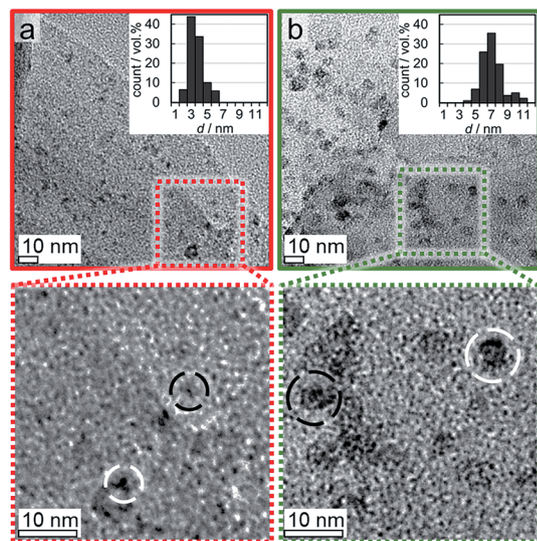


Fig. 16 Transmission electron micrographs of 1 wt% (a) Co_{3.5} and (b) Co_{7.6} on exfoliated graphite after reduction in hydrogen at 350 °C and 330 °C, respectively, during *in situ* X-ray diffraction with volume-based size distribution and enlarged areas displaying reduced and passivated crystallites of unchanged sizes (encircled black) and after sintering (encircled white).

EG) and 3.6 ± 1.0 nm (1 wt%. Co_{3.5}/EG) were measured (Fig. 16). The size distribution of the larger crystallites in Co_{7.6}/EG suggests a slight bimodal distribution, which is an indicator for sintering of a small fraction of the nanoparticles.^{73,100} Less than 10 wt% of the Co fraction participated in sintering, *i.e.* most of the nanoparticles displayed a high stability during reduction in H₂ as supported by *in situ* XRD. In contrast, almost all of the smaller nanoparticles in Co_{3.5}/EG maintained their initial size, potentially due to a superior initial distribution of the nanoparticles over the support. Pronounced metal–support interaction due to the decreased size can also be expected to result in an enhanced stabilisation of the nanoparticles by the support.

4 Conclusions

Graphitic carbon materials were characterised and tested for their suitability as support material for cobaltous nanoparticles in model catalyst systems. When compared to pristine graphite powder, exfoliated graphite was shown to provide a significantly increased surface area allowing for a high dispersion of nanoparticles over the support surface. However, a cobalt loading as low as 1 wt% is required in order to minimise the formation of aggregates which sinter easily during reduction in hydrogen. A mildly size-dependent reduction behaviour of Co₃O₄ crystallites in H₂ was identified utilising this newly developed model catalyst system exhibiting a delayed reduction of smaller nanoparticles to CoO, as well as to fcc-Co in the second reduction step. The model catalysts were highly reducible at moderate temperatures due to the weak interaction of the relatively inert exfoliated graphite support with the nanoparticles. The developed model catalyst provides great

opportunities in order to study isolated Co₃O₄ and fcc-Co nanoparticles in various applications. Exfoliated graphite as support material further allows for fundamental insight into the physicochemical properties of supported nanoparticles by limiting the influence of the carrier material.

Conflicts of interest

There are no conflicts to declare.

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