Disk-Shaped Cobalt Nanocrystals as Fischer–Tropsch Synthesis Catalysts Under Industrially Relevant Conditions

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Abstract
Colloidal synthesis of metal nanocrystals (NC) offers control over size, crystal structure and shape of nanoparticles, making it a promising method to synthesize model catalysts to investigate structure-performance relationships. Here, we investigated the synthesis of disk-shaped Co-NC, their deposition on a support and performance in the Fischer–Tropsch (FT) synthesis under industrially relevant conditions. From the NC synthesis, either spheres only or a mixture of disk-shaped and spherical Co-NC was obtained. The disks had an average diameter of 15 nm, a thickness of 4 nm and consisted of hcp Co exposing (0001) on the base planes. The spheres were 11 nm on average and consisted of ε-Co. After mild oxidation, the CoO-NC were deposited on SiO2 with numerically 66% of the NC being disk-shaped. After reduction, the catalyst with spherical plus disk-shaped Co-NC had 50% lower intrinsic activity for FT synthesis (20 bar, 220 °C, H2/CO = 2 v/v) than the catalyst with spherical NC only, while C5+-selectivity was similar. Surprisingly, the Co-NC morphology was unchanged after catalysis. Using XPS it was established that nitrogen-containing ligands were largely removed and in situ XRD revealed that both catalysts consisted of 65% hcp Co and 21 or 32% fcc Co during FT. Furthermore, 3–5 nm polycrystalline domains were observed. Through exclusion of several phenomena, we tentatively conclude that the high fraction of (0001) facets in disk-shaped Co-NC decrease FT activity and, although very challenging to pursue, that metal nanoparticle shape effects can be studied at industrially relevant conditions.

Keywords Cobalt · Nanocrystals · Anisotropy · Disk · Model catalyst · Fischer–Tropsch synthesis

1 Introduction

Controlled synthesis of supported metal nanoparticles is one of the main challenges in designing heterogenous catalysts for major industrial processes as well as fundamental research [1–3]. Traditional catalyst synthesis methods have in common that nanoparticle formation takes place on the support [4]. Recent advances in catalyst synthesis include the use of colloidal techniques to synthesize nanocrystals (NC) in solution followed by deposition on a support [4, 5]. By separating the NC synthesis from deposition onto the support, the properties of the NC can be manipulated with greater precision, leading for example to NC of particular sizes, crystal structures and shapes [6]. Among these properties, nanoparticle shape is arguably the hardest to tune using conventional techniques, so here colloidal methods could offer a clear advantage over existing catalyst synthesis techniques, provided that challenges regarding NC deposition onto the support and ligand removal are solved [7].
Anisotropic NC are often metastable compared to spherical NC, which are usually the equilibrium shape in particular in the case of metals [8, 9]. By using ligands that bind preferentially to specific surface facets, it is possible to inhibit growth in certain directions and to drive the process towards a kinetically controlled product [10, 11]. Distinct crystal facets are exposed at the surface of these particles, which may lead to distinct catalytic performance. Additionally, different shapes also imply different relative amounts of corner, edge and terrace sites, the reactivity of which can vary depending on the type of reaction [12]. The effect of NC shape on catalytic performance has already been investigated mainly for noble metal [13–15] or for metal oxide NC [16], yet few accounts of shape variation for supported base metal NC catalysts exist.

The shape of NC plays an important role if the catalyzed reaction is structure sensitive. One pronounced example of such a reaction is the Fischer–Tropsch (FT) synthesis [12]. In FT, a mixture of H₂ and CO called synthesis gas is converted into hydrocarbon fuels and chemicals over a catalyst, typically based on Fe or Co [17]. The cobalt-catalyzed FT reaction is sensitive to various structural aspects of the Co particles [18]. For example, a size dependency of the catalytic performance of cobalt is generally reported, with the surface-specific activity and C₅⁺-selectivity decreasing sharply below a particle size of 6–8 nm [19–21]. Additionally, several studies indicated that the crystal structure of Co is relevant, with hcp being more active than fcc [22–26], although this topic remains controversial [27]. Given the potential influence of the crystal structure, the shape of Co particles might be important as well, as shape partially governs the surface structure of nanoparticles.

For the potential effect of the NC shape on catalytic FT performance we should resort mainly to theoretical and surface science studies, as little experimental data is available on complex model catalysts under high-pressure conditions. Theoretical studies predict that the B5 surface ensemble is essential for CO-dissociation, which is considered an important step for activity and selectivity. The occurrence of coordinatively unsaturated sites, including B5 sites, varies with particle size [28, 29] and crystal structure [29] and offers one explanation for the observed structure sensitivity. Furthermore, a computational comparison of the FT activity of various Co crystal planes indicated that higher index planes of hcp Co are orders of magnitude more active than the close packed (0001) [25]. Surface science experiments on single crystal planes of hcp Co showed that relatively more CH₄ is produced on hcp (0001) than on higher index planes, leading to a lower selectivity towards C₅⁺-products [30, 31]. However, a recent experimental study indicated that hcp (0001) displayed intermediate activity and C₅⁺-selectivity, with both increasing in the order hcp (1120) < (0001) < {1011} [32]. All of these aspects of surface composition could in principle be affected by the shape of the Co-NC, although it is also possible that surface defects [33], hydrocarbon coverage [34] or CO-induced surface reconstruction [35–41] might in practice annihilate the effects of Co-NC shape.

Surface science cannot always be directly translated to more complex catalysts and high-pressure systems [5]. To bridge the gap between surface science and supported catalysts under industrially relevant reaction conditions, 3D model catalysts containing anisotropic Co particles are required. Examples of anisotropic cobalt nanoparticles in FT include unsupported particles, such as CoMnOₓ nanoprism [42], Co nanocubes [43], bimetallic nanorods [44], and Co₃O₄ nanorods [45], plates or cubes [32]. Furthermore, hcp nanorods supported either on SiO₂–Al₂O₃ [46] or on Ni or Cu foams [47] have also been employed. Clearly, synthetic approaches are still required to produce anisotropic model systems that closely resemble conventional supported cobalt catalyst, with stable Co-NC shapes yet without additives, such as a second metal, that might influence the FT performance.

Here, we report our efforts to advance the synthesis, characterization and testing of shape-controlled Co-NC dispersed on a support as relevant model catalysts for the Fischer–Tropsch synthesis. We consistently obtained a mixture of disk-shaped and spherical Co-NC. The disks were 15 nm in diameter with a thickness of 4 nm and consisted of hcp Co with (0001) terminating the base planes, while the spherical particles consisted of ε-Co. After low-temperature oxidation of the Co-NC, the CoO-NC were deposited on SiO₂ (50 m² g⁻¹). Numerically 66% of the NC were disks, leading to a higher occurrence of hcp (0001) on the cobalt surface compared to purely spherical Co-NC. The FT activity of the disks/spheres mixture was half that of spheres only, while the C₅⁺-selectivity was identical. Remarkably, the shape endured the FT conditions, as 62% of the NC after catalysis were disks of the original dimensions. Nitrogen-containing ligands were largely removed, and both catalysts featured the same crystalline composition, leaving the possibility for enhanced hcp (0001) exposure in disks. Based on these experiments, it was concluded that the disk shape of Co-NC had a strong influence on the FT performance, most probably caused by the introduction of a high number inactive low-index Co planes.

2 Experimental Methods

2.1 Synthesis of Disk-Shaped NC

A mixture of disk-shaped and spherical Co-NC was synthesized based on an adapted procedure originally developed by Puntes et al. [10, 48]. The ligands, 50 mg oleic acid (90% technical grade, Sigma-Aldrich) and 600 mg octadecylamine...
(> 99.0% (GC), Sigma-Aldrich), were transferred to a three-necked 100 mL flask with two septa and a reflux condenser connected to a Schlenk line. The mixture was heated to 80 °C under vacuum using a heating mantle while magnetically stirring at 600 rpm. After 30 min, the setup was flushed three times with N₂ before switching to N₂ for the remainder of the synthesis. Then, 7.5 mL 1,2-dichlorobenzene (99% Anhydrous, Sigma-Aldrich) was added to the mixture without air-exposure, a thermocouple was inserted and the mixture was heated to reflux temperature (188 °C). Meanwhile, 270 mg dicobalt octacarbonyl (95%, stabilized with hexane, Acros Organics) was transferred to a vial in an N₂-glovebox and the vial was left open inside the glovebox for 40 min to age. After ageing, the Co₂(CO)₈ was dissolved in 1.5 mL 1,2-dichlorobenzene and the vial was closed using a cap with a septum and removed from the glovebox. Once the liquid was at reflux temperature, the cobalt precursor was rapidly injected into the center of the flask using a syringe with a Ø 1.2 × 40 mm needle under 600 rpm stirring. The synthesis was quenched after 5 min using a water bath. The water bath was removed after ~3 min when the temperature of the reaction mixture was below 30 °C. Simultaneously, the stirring was stopped and the stirring plate placed 5 cm below the flask to magnetically separate the reaction product. The top 7 mL contained the highest concentration of disk-shaped Co-NC and was transferred to a glass centrifuge tube after 5 min and exposed to air from this point on. Consequently, the NC were oxidized to CoO to facilitate their uniform distribution over a support at a later stage [7]. The centrifuge tube was filled to 20 mL with 2-propanol (≥ 99.5% (GC), Merck) and centrifuged at 2500 G for 10 min. The purple/brown supernatant was decanted and the residue redispersed in 0.5 mL n-hexane (99%, Acros Organics) using sonication. Another precipitation-redispersion cycle was performed by filling the tube to 20 mL with 2-propanol, centrifuging at 2500 G for 40 min, decanting the supernatant and finally redispersing the residue in 10 mL n-hexane using sonication.

### 2.2 Deposition of Disk-Shaped NC

The disk-shaped CoO-NC were deposited on SiO₂ (Aerosil OX 50, Evonik) within 4 h after synthesis. To obtain enough material with sufficient metal loading, two NC synthesis batches were deposited consecutively on the same batch of support. A glass round bottom vessel with an overhead mechanical stirrer was placed in a sonication bath. 300 mg SiO₂ and 20 mL n-hexane were added and the top of the vessel was loosely covered with aluminum foil. Stirring and sonication were applied to suspend the SiO₂ for 20 min and then the first CoO-NC batch in 10 mL n-hexane was added dropwise in 15 min under continued stirring and sonication. The mixture was left for 45 min after the addition and then the n-hexane was slowly evaporated in 15 min by a applying a flow of N₂ through the vessel, all under sonication and stirring. This entire procedure was repeated a second time by suspending the dried uniformly grey powder again in 20 mL n-hexane and adding the second batch of CoO-NC. Afterwards, the dried powder was suspended in 10 mL n-hexane and transferred to a glass centrifuge tube. The solid was precipitated by centrifuging at 2500 G for 1 min and the supernatant was decanted. The residue was 5 times resuspended in 5 mL n-hexane and precipitated by centrifuging at 2500 G for 1 min and finally dried at ambient temperature under vacuum for 16 h. The dry powder was pressed and sieved to a grain size of 75–150 μm (100 mg was obtained).

### 2.3 Synthesis of Spherical NC

Exclusively spherical Co-NC were synthesized according to a comparable hot-injection method, however, only oleic acid was used as ligand according to a procedure described before [49]. The synthesis was performed in N₂ atmosphere using a Schlenk line. 65 mg oleic acid was degassed under vacuum at 100 °C for 30 min under 650 rpm magnetic stirring in a 3-necked 100 mL flask with a condenser and two septa. The flask was flushed with N₂ and the synthesis was continued under N₂. Then, 7.5 mL 1,2-dichlorobenzene was added under N₂ atmosphere and the solution was heated to 165 °C. Meanwhile, 270 mg dicobalt octacarbonyl was dissolved in 1.5 mL 1,2-dichlorobenzene and sealed under N₂ atmosphere inside a glovebox. This cobalt precursor was then rapidly injected (needle: Ø0.9 × 70 mm) into the pre-heated solution at 750 rpm stirring and reacted vigorously. After 20 min, the reaction was quenched and cooled to at ambient temperature using a water bath. One septum was then removed to oxidize the Co-NC to CoO-NC by air-exposure at low temperature for 1 h (650 rpm stirring). The suspension was divided after over two glass centrifuge tubes, which were filled further to 20 mL with 2-propanol and centrifuged at 2200 G for 30 min to precipitate the NC. The supernatant was decanted, the precipitate redispersed in 0.5 mL n-hexane by sonication and the tubes were filled again to 20 mL with 2-propanol and centrifuged at 2200 G for 40 min. This precipitation-redispersion cycle was repeated two more times and afterwards, the residue was redispersed and combined in 2 mL n-hexane in total.

### 2.4 Deposition of Spherical NC

The spherical CoO-NC were deposited on the same low surface area SiO₂ as the disk-shaped CoO-NC, according to a method described before [49]. The SiO₂ had already been pressed and sieved to a fraction of 75–150 μm. The support (500 mg) was transferred to a 100 mL 3-necked flask and a mixture of the Co-NC in n-hexane and 6.5 mL 1-octadecene (90%, technical grade, Sigma-Aldrich) was
added under 400 rpm magnetic stirring. The flask was then equipped with a condenser, glass plug and septum and connected to a Schlenk line. The suspension was degassed under vacuum at 100 °C for 15 min and as a result, the n-hexane in the suspension evaporated. Then, the flask was flushed with N₂ and the deposition was continued under N₂. The suspension was heated to 200 °C for 30 min, after which the heating mantle was removed, causing the solution to cool to room temperature within 30 min. The sample was washed by first centrifuging at 1500 G for 5 min and decanting the supernatant. Subsequently, the precipitate was resuspended in 2 mL n-hexane through sonication, 6 mL acetone was added and the sample was isolated by centrifuging at 1500 G for 5 min and decanting the supernatant. This precipitation-resuspension cycle was repeated five times, followed by a last cycle in which the n-hexane/acetone mixture was replaced by 20 mL acetone. Finally, the sample was dried in three consecutive steps, i.e. 60 °C for 1 h in static air, 120 °C for 3 h in static air and 80 °C for 3 h under vacuum, and pressed and sieved to a grain size of 75–150 μm.

2.5 Characterization

Transmission electron microscopy (TEM) was performed on a Tecnai 20 (FEI) operated at 200 kV. Samples of colloidal suspension were prepared by drop casting a diluted sample directly on a carbon-coated TEM grid. In the case of SiO₂-supported CoO-NC, the powder was first suspended in 2-propanol by sonication and subsequently drop casted on a carbon-coated TEM grid. The TEM images of supported CoO-NC were manually analyzed using ImageJ. In the case of the colloidal suspensions, the images of CoO-NC were automatically classified as either disks or spheres and manually reviewed before analysis. All reported NC sizes are number averaged and recalculated to the equivalent Co⁰ size to compensate for lattice expansion due to a 3 nm thick CoO passivation layer. Because of the limited height of the disks, these NC were assumed to oxidize completely to CoO. The metallic surface area was calculated using the surface area weighted mean diameter, again after correcting for the size to compensate for lattice expansion due to a 3 nm thick CoO passivation layer. A cross-sectional area of 0.0662 nm² per cobalt atom was assumed in order to calculate the metallic surface area [50].

Vitrification of freshly synthesized colloidal suspensions for cryogenic transmission electron microscopy (cryo-TEM) was performed using a Vitrobot Mark IV (FEI) vitrification robot. The Co-NC were vitrified in the original reaction suspension within 30 min from magnetically separating the suspension product. For the vitrification, 3 μL of Co-NC suspension was applied to a 200 mesh, lacey carbon-coated Cu grid at 25 °C in a chamber saturated with toluene vapor. Subsequently, the grid was blotted for 3 s with filter paper and plunge-frozen in liquid N₂ (− 196 °C). The samples were kept in liquid N₂ for several hours and loaded into the Titan Krios (FEI) equipped with a field emission gun operating at 300 kV and energy filter. Images were recorded using a 2 k × 2 k Gatan charge coupled device (CCD) camera. High-resolution transmission electron microscopy (HR-TEM) was performed in areas where no solvent appeared present. Low dose selective area diffraction (SAED) was used for structure analysis and the diffraction patterns were processed according to the method described by Leijten et al. [51].

Electron tomography was carried out using an FEI Talos F200X operated at 200 kV in bright field TEM mode. The CoO-NC/SiO₂ sample was suspended in ethanol by ultrasonication. A few droplets of the suspension were deposited onto a Quantifoil R2/1 carbon film supported parallel-bar Cu grid which already contained Au nanoparticles of 5 nm in diameter. TEM images were recorded with a Ceta camera (4096 × 4096 pixels) over a tilt range of − 6° to + 76° with tilt increments of 2° at a nominal magnification of 74,000 times. The tilt series was aligned using IMOD software [52] using the Au nanoparticles as fiducial markers. The aligned tilt series were binned by 4 and reconstructed using the back-projection algorithm. The resulting reconstructed volumes had a final voxel size of (0.572 nm)³. Segmentation of disk-shaped CoO-NC, spherical CoO-NC and the SiO₂ support in the reconstructed volumes was performed in FIJI [53]. Adequate threshold values were applied to median-filtered reconstructed volumes. Disk-shaped NC, spherical NC and the SiO₂ support were defined in 3D by manually delimiting their boundary. Volume rendering of the segmented volumes was carried out in FIJI.

Inductively coupled plasma-optical emission spectroscopy (ICP-OES) was measured on a SPECTRO ARCOS after digestion of the sample in aqua regia.

X-ray photoelectron spectroscopy (XPS) was measured on a Kratos Axis Ultra DLD with an X-ray source operating at 225 W and 15 keV, and equipped with an Al anode (EKα = 1486.6 eV). The samples were fixed using Ag-tape at ambient conditions and loaded in the setup. High-resolution spectra of Co2p, Si2p, O1s and C1s were measured at 40 eV pass energy, while N1s and the survey scan were recorded at 160 eV pass energy. Kratos Axis sensitivity factors were used for quantification. XPS measurements were performed on three distinct locations on all samples and the results were averaged. In case of the N1s scan, multiple measurements of the same location were accumulated to improve the signal-to-noise ratio. The binding energies were calibrated by setting the C–C/C–H bond peak to 284.8 eV.

In situ X-ray diffraction (XRD) was performed on a Bruker D8 Discover with a Mo (Kα₁ 0.709 Å) source in Debye–Scherrer transmission (capillary) geometry. The X-ray beam was focused on a quartz capillary with 1000 μm OD and a wall thickness of 10 μm using a Göbel-mirror. The setup was equipped with an energy dispersive LynxEye XE
Position Sensitive Detector (PSD). Details on the complete setup can be found in recent publications [54, 55]. The patterns were measured between 5 and 39° 20 with an increment of 0.07° and time per step of 20.5 s. Multiple scans were accumulated to improve the signal-to-noise ratio only if the patterns were identical. Rietveld Quantitative Phase Analysis (Rietveld QPA) was performed on the measured diffractograms using Bruker TOPAS v5 software. Details of the Rietveld refinement procedure are given in Supporting Methods.

Co-S/SiO2 was diluted with 2 mass equivalents SiO2 (Aerosil OX 50) to obtain a similar overall cobalt loading as Co-D/SiO2. Typically, 7 mg sample in the sieve fraction 75–150 μm was loaded in the capillary and fixed with quartz wool at either end, resulting in a catalyst bed length of ~20 mm. The capillary was placed in the setup and the catalysts were reduced in situ at 350 °C for 8 h (1 °C min−1) in a 1.2 mL min−1 flow of 25 vol% H2 in He at 1 bar. After reduction, the temperature was lowered to 180 °C with 3 °C min−1 and at this temperature, a 3.0 mL min−1 flow of synthesis gas H2/CO = 2 (v/v) was introduced. After 10 min stabilization, the back-pressure regulator was set to 10.6 bar. Only after the set pressure was reached and stable, the temperature was increased to 220 °C with 1 °C min−1, which was defined as TOS = 0 h. In situ XRD patterns were recorded both during reduction and FT.

2.6 Fischer–Tropsch Synthesis

The Fischer–Tropsch synthesis performance of the catalysts was investigated using a Flowrence (Avantium) 16 parallel reactor setup. 80 mg Co-D/SiO2 and 12 mg Co-S/SiO2 were first diluted with 100 mg SiC (212–425 μm) and then loaded into stainless steel plug-flow reactors of 2.6 and 2.0 mm (ID), respectively. The samples were dried for 2 h in an He flow and reduced in situ in a 25 vol% H2 in He flow at 350 °C for 8 h after a heating ramp of 1 °C min−1. After reduction, the temperature was lowered to 180 °C with 3 °C min−1 and the reactors were pressurized to 20 bar in an H2 flow. Then, the synthesis gas feed was introduced with H2/CO = 2 v/v and 5 vol% He as internal standard and after 1 h stabilization, the reactors were heated to 220 °C with 1 °C min−1, which was defined as the starting point for FT. After the run, the catalysts were treated in H2 at 200 °C for 10 h to remove carbonaceous products from the catalyst surface and cooled down further under He.

The reaction products were analyzed online using an Agilent 7890A GC with two separate channels. In the first channel, the hydrocarbon products were analyzed using an Agilent J&W PoraBOND Q column in combination with an FID. In the second channel, the permanent gasses were separated on a ShinCarbon-ST column and analyzed using a TCD. The catalytic activity was given as CO conversion (XCO) cobalt-time yield (CTY) and turnover frequency (TOF). The TOF was based on the CTY and the average metallic surface area of the spent catalyst, as determined by TEM. The selectivity (in %C) to C1–C4 hydrocarbons was determined as $S_{C1–C4} = 100 \frac{F_{C1–C4}}{F_{CO, in} X_{CO}}$ with n representing the product’s carbon number and F the flow of the hydrocarbon or CO. The selectivity to C5+ hydrocarbons was calculated as $S_{C5+} = 100\% - S_{C1–C4}$.

3 Results and Discussion

Cobalt nanocrystals (NC) were synthesized by a hot-injection synthesis using oleic acid and octadecylamine as ligands with the aim to synthesize disk-shaped NC and the results were analyzed by TEM (Fig. 1). Both disk- and spherical-shaped CoO-NC were obtained from a single synthesis with the ratio between the two varying strongly with the reaction conditions, which made separation necessary, in line with literature [10]. Enrichment of the disk-shaped over spherical NC was realized by a magnetic field, using the difference in the magnetic properties between the different NC. In our case, the large magnetic moment of the spheres resulted in faster precipitation from the solution compared to the disks under the weak magnetic field of the stirring plate (Fig. S1). This led to enhanced concentrations of disks in the top layer of the reaction mixture (Fig. 1a–e). After magnetic separation, the work-up procedure also affected the disk-to-sphere ratio, as the disk yield decreased with more precipitation-redispersion cycles (Fig. 1a–c). This indicates that spheres were preferentially precipitated during centrifugation or alternatively, that the fraction of disks already varied in the original reaction mixture of different batches.

The elongated, rod-like particles (mainly visible in Fig. 1a) were identified as stacks of disk-shaped particles orientated side to side. From these particles, the average dimensions of the disks were determined (Fig. S2). The average diameter of the disks was 15.5 ± 3.5 nm and the thickness 4.2 ± 0.6 nm, leading to an aspect ratio (height/diameter) of 0.27. As a comparison, all-spherical CoO-NC of 8.3 ± 1.1 nm were synthesized using only oleic acid as ligand, as described before [49] (Fig. 1f).

A batch containing disk-shaped NC after three precipitation-redispersion cycles was analyzed in more detail (Fig. 1c). The NC in the TEM image were semi-automatically divided into two groups (Fig. 1e) based on their contrast, which is a measure for their thickness, in relation to their size and morphology. The large, low-contrast NC were (truncated) hexagonally-shaped and the smaller, high-contrast NC were spherical. To examine the 3D shape of these NC, the stage was tilted 50° inside the TEM (Fig. 1d, S3 and supporting movie S1). Analysis of 100 NC from both groups (Fig. S4) revealed that the size of the particles was 15.4 nm
at 0° and 11.9 nm at 50° in the case of the low-contrast particles, proving that these NC were indeed smaller in the z direction as expected for disk-shaped NC. When considering an aspect ratio of 0.27, the 2D projection length at 50° should be 13.1 nm (Fig. S5), which matches the experimental value quite well. On the other hand, the smaller, high-contrast particles were 10.7 nm at 0° and 10.6 nm at 50°, showing that their diameter was the same in all directions and that these NC were indeed spherical. Because the shapes of the CoO-NC could be accurately distinguished, the numerical disk yield could be determined and was 53% for the batch after three precipitation-redispersion cycles.

Disk-shaped Co-NC are a metastable phase [8, 56], making the synthesis results highly susceptible to small variations in the procedure. Therefore, optimizing the reaction conditions was essential to obtain a high fraction of anisotropic particles. Important synthesis parameters included the speed of magnetic stirring and the speed of precursor injection, as faster stirring or slower injection through a thinner needle led to enhanced formation of 10–30 nm spherical NC and a lower yield of disk-shaped NC. Furthermore, higher disk yields were obtained with batches of Co$_2$(CO)$_8$ that had been aged 40 min in the glovebox prior to use in synthesis. These phenomena have not been reported before for this synthesis method and illustrate the delicate nature of this approach.

The crystal structure of the as-synthesized Co-NC before oxidation was analyzed using a combined cryo- and high-resolution-TEM approach (Fig. 2). A Co-NC suspension was vitrified shortly after NC synthesis and transported and loaded in liquid N$_2$ to avoid oxidation of the Co-NC. The vitrification protected the sample from oxidation and HR-TEM was possible because there was little to no solvent present in the TEM sample. Electron
diffraction of the area in Fig. 2a resulted in the diffraction pattern in the inset of Fig. 2a and was radially averaged to give Fig. 2b. The main peak at \( d = 2.06 \, \text{Å} \) could not be unambiguously assigned to one crystal structure, as this lattice spacing is present in all three metallic cobalt phases (Fig. S6). The diffraction peak at \( d = 2.19 \, \text{Å} \), however, only matched hcp cobalt. Furthermore, the peaks at \( d \)-values of 1.26 and 2.19 Å could be ascribed to \( \{11\bar{2}0\} \) and \( \{10\bar{1}0\} \) of hcp cobalt, respectively, which were more intense than expected. Considering that \( \{11\bar{2}0\} \) and \( \{10\bar{1}0\} \) have periodicity in the \( x,y \) direction and not in the \( z \) direction, this proved that the hcp cobalt was present in anisotropic, disk-shaped NC and that the short \( z \) axis is along the [0001] direction, in line with literature [10]. Based on the intensity ratios of the hcp Co phase, it is likely that the peak at \( d = 2.06 \, \text{Å} \) has another dominant contribution either from fcc or \( \varepsilon \)-cobalt.

Further information on the crystal structure of the NC before oxidation was obtained by investigating an individual NC in more detail with HR-TEM on the same cryogenic sample. The Co-NC in the framed area in Fig. 2c was identified as a disk-shaped NC because of its low contrast and diameter of 15 nm. Lattice fringes of this particle were imaged and the Fourier transform revealed hexagonal symmetry of the crystal lattice, indicative of hcp cobalt. The determined lattice spacings were 2.28 and 1.29 Å, which were again matched to \( \{11\bar{2}0\} \) and \( \{10\bar{1}0\} \) of hcp Co. Additionally, the other prominent particle of 11 nm with higher contrast was identified as a spherical NC and did not show hexagonal symmetry. Its lattice spacings matched \( \varepsilon \)-cobalt best (Fig. S7), indicating that the other crystal structure observed by electron diffraction was most likely \( \varepsilon \)-cobalt, which is consistent with literature [10]. It should be emphasized that this concerned the crystal structure of the Co-NC before oxidation and that it is uncertain what crystal structure would evolve after low-temperature oxidation, in situ reduction and catalysis.

The CoO-NC after two precipitation-redispersion cycles were deposited on low surface area SiO\(_2\) (50 m\(^2\) g\(^{-1}\)) using a method that avoids the use of high temperatures to affect the NC shape as little as possible. This particular type of SiO\(_2\) was selected, because it had yielded very stable Co-NC based catalysts under FT conditions before [49]. The sample containing mainly disk-shaped NC was designated as Co-D/SiO\(_2\). Two washing steps after NC synthesis were the optimal compromise between uniform attachment of the CoO-NC with lower ligand concentration on one hand (Fig. S8) and progressively lower NC yields on the other. Based on ICP-OES, a cobalt loading of 1.4 wt% was obtained using this deposition method. The Co-D/SiO\(_2\) sample was analyzed by TEM (Fig. 3a–c). The NC were largely uniformly distributed over the support, although some areas with clustered disk-shaped and spherical NC were encountered as well (Fig. 3c). Despite this occasional clustering of NC, NC were largely well dispersed on the support, making the catalyst useful for further investigations.

It was challenging to distinguish the disk-shaped NC from the SiO\(_2\) support because of the low contrast of the CoO-NC associated with their limited thickness when orientated perpendicular to the electron beam. However, based on a combination of contrast, size and morphology, it was possible to identify the CoO-NC and discriminate between disks and spheres (Fig. 3a, b). After analysis of 203 NC, 66% of the NC were classified as disks and 34% as spheres. In addition, a reference catalyst containing exclusively spherical CoO-NC was prepared by depositing the 8.3 nm CoO-NC on the same low surface area SiO\(_2\) (Fig. 3d). This catalyst was designated as Co-S/SiO\(_2\). The obtained cobalt loading was 9.6 wt% based on ICP-OES. Furthermore, the CoO-NC were...

**Fig. 2** High-resolution TEM under cryogenic conditions of as-synthesized disk-shaped and spherical Co-NC with little to no solvent present. **a** Overview of several clustered Co-NC with the selected area electron diffraction pattern in the inset. **b** Intensity profile obtained by radially averaging and baseline correcting the SAED pattern. **c** Several Co-NC at higher magnification. The framed area contained a 15 nm, low contrast NC that was classified as a disk; the inset gives the Fourier transform of the framed area.
uniformly distributed over the support and no areas containing clustered NC were observed. This catalyst was described in more detail in an earlier publication [49].

Electron tomography of Co-D/SiO₂ was performed to investigate the morphology of the NC on the support in more detail (Fig. 4, complete tilt series from −66 to +76° included as supporting movie S2). The disks could be clearly distinguished from the spheres using the tilt series, because only the projected aspect ratio of the disks varied with the tilting angle. The accuracy of the method applied before to classify the NC as disks or spheres based on size, morphology and contrast was confirmed by tomography and the obtained results for this limited sample size (only 26 NC in field of view) matched the earlier observation of ~65% of the NC present as disks. Furthermore, a 3D representation of a selected part of the sample was obtained through reconstruction and segmentation of the tilt series (Fig. 4c, reconstruction and 3D-rendered volume as supporting movies S3, S4). From this representation, the morphology of NC can be clearly inferred as well as their interaction with the support. It showed that the disk-shaped NC typically attached to SiO₂ with one base plane. Future work on metal-support interactions for the differently shaped NC could involve, e.g., XPS experiments to explore electronic interactions of the Co-NC with the support.

The FT performance of the catalysts was evaluated over 100 h on stream at similar CO conversion levels (Fig. 5, Table 1). The cobalt-weight-based activity (cobalt-time yield, CTY) against TOS is shown in Fig. 5b. Co-S/SiO₂ was twice as active per unit weight of cobalt as Co-D/SiO₂. Because of the different particle sizes and shapes, the activity was compared as cobalt-surface-based activity (turnover frequency, TOF). The metallic surface area necessary for the TOF was calculated from the NC sizes in the spent catalyst, which will be discussed later (Table 1). The TOF of Co-D/SiO₂ was 49 × 10⁻³ s⁻¹ and the TOF of Co-S/SiO₂ was 86 × 10⁻³ s⁻¹, the former value being in line with other work on Co/SiO₂ using conventional catalyst preparation techniques such as impregnation [57]. However, the TOF of Co-S/SiO₂ was remarkably high, possibly pointing to a higher fraction of more active hcp Co [22–26] resulting from the NC-based synthesis compared to conventional synthesis methods.

The selectivities towards C₅⁺, as well as C₁⁻ and C₂–C₄, products were similar (Table 1) and overall ~5% lower than on conventionally prepared Co/SiO₂ catalysts [57]. Consequently, the ASF distributions and derived chain growth probabilities (α) were comparable as well. Plots of the C₅⁺-selectivity as a function of TOS and the ASF distributions are included as Fig. S9. The effect of crystal structure on selectivity is still under debate, as some reports indicated no difference between hcp and fcc Co [22, 27], while others observed higher C₅⁺-selectivity on hcp Co [24, 58]. However, in our case, similar C₅⁺-selectivity of both catalysts is in line with expectations when considering that hcp (0001) surfaces are virtually inactive compared to higher index surfaces. Therefore, if (0001) concentrations for Co-D/
SiO$_2$ were to be enhanced, these additional surfaces would contribute only marginally to the overall product spectrum and as a result, the selectivity would hardly change. Thus, changing the NC shape from spheres to disks did not influence the selectivity of the catalyst.

Overall, the main difference in performance was the factor 2 lower intrinsic activity of the disk-containing Co-D/SiO$_2$ sample, which could have several reasons. First, the activity could be lower as a result of the shape and thus the exposed facets, which would match the theoretical and experimental finding that hcp (0001) is less active than higher-index hcp or fcc facets [25, 32]. Second, residual ligands could still be present in the case of Co-D/SiO$_2$ and block part of the cobalt surface. Third, the surface area of Co-D/SiO$_2$ could be lower than expected if the degree of reduction is low or if there is substantial NC growth, for example in the areas where clustering of NC was observed (Fig. 3c). These potential explanations were investigated in more detail and are described below.

The structure of the spent catalysts was analyzed after FT and subsequent passivation using TEM (Fig. 6). The sizes and shapes of 160 cobalt nanocrystals in Co-D/SiO$_2$ were quantified (Fig. 6a, b), showing numerically 62% disk-shaped NC with $d = 14.5$ nm and 38% spherical NC with $d = 10.2$ nm. The aspect ratio of the disks could not be reliably determined for this sample and was assumed to have remained unchanged. These results were very similar to those of the pristine catalyst and the disk shape was thus remarkably stable under FT conditions. Furthermore, a more clustered part of the catalyst was also observed where some particle growth had occurred (Fig. 6c). This area probably originated from an area of clustered CoO-NC in the pristine catalyst. The size of the NC in Co-S/SiO$_2$ was unchanged as well with spherical particles of 8.7 nm on average (Fig. 6d). The stability of the Co-NC in terms of both size and shape under reaction conditions meant that catalytic performance might be related to the disk shape.

Two types of ligands were used to synthesize disk-shaped Co-NC (oleic acid and octadecylamine) while spherical Co-NC were synthesized using only oleic acid. For oleic acid, efficient ligand removal has been shown [7], but could be different for linear amines. We assessed the
influence of the octadecylamine ligand in FT by investigating the amount of surface nitrogen species using XPS on the catalysts as-prepared and after 100 h FT followed by passivation. Low concentrations of N were detected on both as-prepared catalysts and on bare SiO2 (Fig. 7a, complete survey scans in Fig. S10). The N-to-Co atom ratio was 4 times higher on Co-D/SiO2 (0.07) than on Co-S/SiO2 (0.02). After FT and passivation, Co-D/SiO2 was partially covered in carbon and only an indication of N was observed. Spent Co-S/SiO2 was too covered in carbon to yield useful data on Co and N contents. The N/Co atom ratio of spent Co-D/SiO2 seemed similar to that of as-prepared Co-S/SiO2, so excess N-species in Co-D/SiO2 from the NC synthesis procedure were removed during reduction/FT. Harmel et al. [46] quantified that the content of the ligand hexadecylamine decreased from 8 wt% to 3 wt% during reduction under H2 at 270 °C, which is substantially lower than the reduction temperature of 350°C applied here. Therefore, it seemed that our reduction procedure was sufficient to activate the catalyst and minimize the effect of remaining N-species.

To get more insight into the structure of the catalysts under FT conditions, in situ XRD was performed on both catalysts at 10 bar, 220 °C and an H2/CO ratio of 2 v/v (Fig. 7b, Fig. S11). To obtain satisfying Rietveld refinement results, it was necessary to allow preferred orientations of hcp Co crystallites, similar to what has been reported before [55]. We ascribe this to polymorphism of Co in the Co-NC/SiO2 samples, since XRD analysis of metallic cobalt is complicated by the propensity of cobalt to include stacking faults [22, 59]. Our analysis method did not quantify this intergrowth structure, but accommodated much of it in the hcp contribution by increasing the intensity of the (0002) peak. The fraction of hcp Co could therefore be overestimated. However, both samples were refined using the same method and given the similarity of the results for both samples, it appears that polymorphism, if present, took place to the same extent in both samples.

The crystalline cobalt phase in both catalysts consisted of approximately 65% hcp Co. The fcc Co and CoO compositions were 21% fcc Co and 12% CoO for Co-D/SiO2 and 32% fcc Co and 3% CoO for Co-S/SiO2. These values are in line with results obtained by Andreev et al. [60] using NMR on Co/SiC at elevated temperatures under static Ar atmosphere and in reasonable agreement with Cats et al. [54] who used operando XRD on Co/TiO2 under FT conditions. In general, however, direct comparison of these results to literature is cumbersome, since the obtained Co phase strongly depends on particle size, metal precursor, support and activation procedure [61, 62]. Furthermore, the average crystallite sizes of metallic cobalt

Table 1 Fischer–Tropsch synthesis results for Co-D/SiO2 and Co-S/SiO2 reported at similar CO conversions and averaged between 90 and 100 h on stream at 20 bar, 220 °C and H2/CO = 2 v/v

| Sample          | GHSV (h⁻¹) | XCO (%) | CTYₐ | TOF (10⁻⁵ molCO·g⁻¹·s⁻¹)ᵇ | C₅⁺-sel. (%)ᶜ | C₂–C₄-sel. (%)ᶜ | C₁-sel. (%)ᶜ | α | Co SA spent catalysts (m²·g⁻¹ Co⁻¹) |
|-----------------|------------|---------|------|---------------------------|---------------|----------------|--------------|--|----------------------------------|
| Co-D/SiO₂       | 1560       | 8.9     | 6.9  | 49                        | 75            | 8.6            | 16           | 0.78 | 56                              |
| Co-S/SiO₂       | 39,900     | 8.1     | 15   | 86                        | 78            | 9              | 13           | 0.76 | 71                              |

ₐ In 10⁻⁵ molCO·g⁻¹·s⁻¹
ᵇ Based on the CTY and the metallic Co surface area of the spent catalysts as derived by TEM

![Fig. 5 Fischer–Tropsch synthesis results of Co-D/SiO₂ and Co-S/SiO₂ at 20 bar, 220 °C and H2/CO = 2 v/v. The GHSV was 1560 h⁻¹ for Co-D/SiO₂ and 39,900 h⁻¹ for Co-S/SiO₂. a CO conversion and b cobalt-time yield as a function of time-on-stream.](image-url)
was 3–5 nm, significantly smaller than the particle dimensions obtained by TEM. This probably indicated that the Co-NC were polycrystalline, which could be related to the occurrence of stacking faults.

The degree of reduction of the crystalline phase was high with 88% for Co-D/SiO₂ and 97% for Co-S/SiO₂ and therefore excluded as a reason for substantial activity differences. Once in the reduced state, the XRD profiles remained unchanged for the duration of the experiment. The crystallites thus remained stable without sintering or restructuring during reduction and the first 18 h FT, in line with the TEM analyses. This also indicated that a significant loss of surface area in Co-D/SiO₂ due to sintering of NC was unlikely. Furthermore, the hcp Co content in Co-D/SiO₂ was similar to that in Co-S/SiO₂, but nevertheless substantial in both samples. Finally, the average crystallite dimensions of 3–5 nm prevented any conclusions on preferred orientations and surface termination in disk-shaped NC with a diameter of 16 nm and 4 nm thickness. Overall, activity differences due to enhanced hcp Co content was excluded, but preferred crystallite orientations within the polycrystalline disk-shaped NC and with that enhanced hcp (0001) terminations were still possible.

Combining the obtained data on the ligands and NC size, shape and crystal structure of the catalysts under reaction conditions, the cobalt specific surface area and the fraction of the surface terminated by hcp (0001) were calculated (Fig. 8). This served as an estimation of the structural difference between the Co-NC disk-containing catalyst and the catalyst containing exclusively spherical NC. To do so, we assumed that the base planes of the disks were terminated by hcp (0001) and that the disks were attached to SiO₂ with one base plane (Fig. 4, supporting movie S4), thereby rendering that plane inaccessible for reactants. This is a conservative estimate, as the curvature of the SiO₂ might in practice make these base planes partially accessible, which would increase the relative occurrence of hcp (0001). Furthermore, the dimensions of the as-synthesized NC were used in surface area calculations, i.e. disks with d = 15.5 nm and h = 4.2 nm and spheres with d = 10.7 nm.

Besides the surface of the disks, also the surface of the spheres has to be considered. According to Wulff reconstructions of spherical hcp cobalt particles by Liu et al. [25], 18% of the surface of a spherical hcp cobalt particle is terminated by (0001). Therefore, the maximal hcp (0001) occurrence in Co-S/SiO₂ was 18%. Compared to an estimated hcp(0001) occurrence of 39% in Co-D/SiO₂, this means that 21% of the metallic cobalt surface area of Co-S/SiO₂ is in a higher index and probably more active form than for the disk-containing Co-D/SiO₂. This holds when all NC consist of hcp Co, while a difference in hcp (0001) exposure of 33% would be obtained if only disks were hcp Co. Considering the uncertainty and error margin in this analysis, we conclude that the difference in hcp (0001) exposure could be of the right magnitude to account for much of the observed activity difference. Therefore, we identified this as a plausible cause for
the differences in catalytic performance of the disk-shaped and spherical NC, also because of the exclusion of other factors that might lead to such a substantial loss of activity, e.g. surface coverage by ligands, or NC restructuring or growth, as discussed before.

This is, to the best of our knowledge, the first time that a supported model catalyst has been synthesized to investigate the effect of Co-NC shape in FT without complications of a secondary metal or interference of ligands and it illustrates the potential of colloidal techniques for model catalyst synthesis. This work shows that nanoparticle shape can be an important design parameter for supported cobalt FT catalysts when operated under industrially relevant conditions. Furthermore, based on these results, we tentatively propose that the extended low-index surface facets of the disks are the main contributor to the lower TOF.

4 Conclusions

We investigated the synthesis of disk-shaped cobalt nanocrystals (Co-NC), their deposition on a support and the subsequent effect of their anisotropy on the Fischer–Tropsch synthesis. Disk-shaped hcp Co-NC were synthesized mixed with spherical ε Co-NC. After oxidation to CoO, these NC were deposited on low surface area silica. Electron microscopy analysis revealed that numerically 66% of the deposited CoO-NC were disks and 34% were spheres. After in situ reduction of the CoO-NC to Co, the Fischer–Tropsch activity of disk-containing catalyst was only half that of a catalyst containing only spherical NC. Interestingly, the catalysts were stable and the size and shape of the NC were retained, even for the disk-shaped Co-NC. XPS showed that most of the amine ligands were removed during activation/FT and in situ XRD revealed that the crystal structure of both catalysts was similar, with NC with 3–5 nm polycrystalline domains and approximately 65% hcp Co and 21 to 32% fcc Co. Based on these experiments, we tentatively propose that changing the Co-NC shape from spheres to disks reduced FT activity because of hcp (0001) facets prevailing. This work clearly demonstrates the potential of colloidal methods to synthesize new, well-defined model catalysts to investigate structure-performance relationships in catalysis even under high-pressure conditions.
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Author Contributions TWD synthesized and characterized the catalysts, evaluated their catalytic performance and drafted the manuscript. JH assisted with sample preparation and JIN synthesized the reference catalyst. HS performed cryo-TEM and interpretation, HY and JZ were involved with tomography and reconstruction, and RO and BMW performed in situ XRD and assisted with data analysis. KPI contributed to experimental design, data analysis and writing of the manuscript. All authors have given approval to the final version of the manuscript.

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Data Availability Data is available upon reasonable request.

Compliance with Ethical Standards

Conflict of interest There are no conflicts of interest to declare.

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