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Effect of reduction on Co catalyst active phase highlighted by an original approach coupling ASAXS and electron tomography

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8

9 Abstract

Diversification of liquid fuel sources for transportation plays a key role in the energetic transition. High 10 quality and clean diesels can be produced by Fischer-Tropsch synthesis, which interest is renewed in as 11 12 much it can be performed from syngas (CO+H₂) produced by conversion of natural gas or gasification of lignocellulosic biomass, leading, in this latter case, to fuels from renewable resource. Catalytic 13 14 performance (activity and selectivity) of cobalt-based catalysts are related to the physico-chemical characteristics of the cobalt nanoparticles. The present study propose a new approach to obtain a detailed 15 16 and exhaustive description of the cobalt active phase in FT catalysts. We combined Anomalous Small 17 Xray Scattering (ASAXS) and electron tomography, giving complementary insights on the microstructure 18 and size distribution of both cobalt nanoparticles and aggregates. This approach, carried out on a 19 cobalt/alumina-silica catalysts studied first at the oxide state and then at the reduced state, allowed to 20 highlight the mechanisms involved at the nanoscale during the reduction step. Reduction impact is 21 significant on aggregates morphology, causing their fragmentation and increasing their accessibility. As a 22 perspective, a better knowledge of cobalt aggregates morphology will help to understand their impact on 23 the catalyst performance.

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- 25

26 Keywords

27 Cobalt, catalyst, Fischer-Tropsch, electron tomography, TEM, ASAXS, nanoparticles

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32 **1 Introduction**

33 Diversification of liquid fuel sources for transportation plays a key role in the energetic transition. 34 Fischer-Tropsch Synthesis (FTS), initially devoted to convert coal into fuels, has become attractive as it 35 can be performed from syngas ($CO+H_2$) produced by gasification of lignocellulosic biomass, leading, in 36 this latter case, to fuels from renewable resource [1][2]. Low-temperature FTS, based on cobalt catalysts, 37 produces long-chain hydrocarbons leading, after upgrading, to high quality and clean diesels [1][3] [4]. 38 Catalytic performance (activity and selectivity) of Fischer-Tropsch catalysts is related to the physico-39 chemical characteristics of the Co nanoparticles. In particular, for all studied supports (carbon nanofibers, 40 γ -alumina), authors observe a rapid increase of the catalytic activity when the cobalt particle size 41 increases to a certain diameter and, then, the activity decreases with further increase in particle size showing a volcano shape curve in catalytic activity. Besides, the C_5^+ selectivity is also known to be 42 related to the size of the nanoparticles and increases while increasing the size of the particles, to circa 43 44 fifteen nanometers. Depending of the support, the optimal size for higher activity and selectivity is 45 between 6 and 9 nm [5] [6] [7], smaller particles leading to a significant activity loss. From these results, 46 it is crucial to characterize the nanoparticles, specially their size, which is unlikely to be monodisperse. 47 However, in high-loaded cobalt catalysts (up to 15 wt % Co), the nanoparticles are usually not uniformly 48 dispersed and aggregation of cobalt nanoparticles is often observed, forming aggregates of tens to 49 thousands of nanometers [8] [9]. However, the impact of Co aggregation on activity and selectivity is still 50 unclear / debated. Munnik et al. [8] [9] have observed that the aggregates present in high-loaded catalysts 51 supported on silica were formed during the drying process, after impregnation of aqueous cobalt nitrate 52 precursors. Thus controlling the drying temperature will allow a decrease in aggregate formation rate. By studying catalysts prepared in order to vary the aggregation rate [8], they related an increase of C_5^+ 53 54 selectivity with the increase of the aggregates size.

Basing on this literature, it seems essential to characterize both nanoparticle and aggregate scales to fully
 understand and explain the catalytic performance of the high-loaded cobalt catalysts.

For this purpose, a complete description of the active phase implies measuring i) the size distribution of the isolated nanoparticles, ii) the proportion of Co involved in aggregates or in individual particles, iii) the size of the aggregates and iv) the microstructure of aggregates that controls the accessibility of reactant and product within the aggregates. These steps are essential to calculate the intrinsic activity and selectivity and evaluate the impact of aggregation independently to other parameters such as the particles size and the support effect. 63 Several complementary techniques are commonly used to characterize the Co active phase in FTS 64 catalysts. In particular, Co dispersion is measured by H₂ chemisorption, the mean oxide crystallites size is 65 usually measured by X-ray diffraction (XRD) [10] and the particles and aggregates size distribution can be measured by techniques based on transmission electron microscopy (TEM), such as dark-field TEM or 66 67 HAADF-STEM (high angle annular dark field-scanning transmission electron microscopy) [6,8,9,11]. 68 Concerning the XRD technique, it provides a good evaluation of the average size of the crystallites for the 69 oxide samples, but it is less appropriate for reduced samples as the reduction of the cobalt particles is not 70 total and the peaks corresponding of the CoO and Co° phase overlap. In addition, the stacking faults 71 present in the cobalt metallic phase lead to a broadening of the peaks not related with the finite size of the 72 crystallites and requires advanced approaches to take them into account, such as Rietveld analysis or 73 pattern simulation of mixture of cubic and intergrown Co [12]. The dark-field TEM also allows size 74 distribution determination of the crystallites but cannot be easily carried out on used catalysts due to the presence of waxes. Furthermore, both XRD and dark-field TEM give information on the crystallites size, 75 76 which does not correspond to the nanoparticles size if they are polycrystalline. HAADF-STEM can be 77 used to measure Co nanoparticles, but the contrast between Co and the support is usually too low to be 78 sensitive to the smallest nanoparticles.

Given the sensitivity of the X-ray scattering phenomenon to the solely electronic density variations, 79 80 regardless of the structure or size of the crystal domains considered, small-angle X-ray scattering (SAXS) 81 techniques seem particularly suitable to study the size distributions of metal particles on supports. The 82 first work on the characterization of metallic particles on supported catalysts was carried out in the late 83 1990s by Haubold [13–16] and Benedetti [17,18]. The model systems studied consisted of noble metal particles (Pt or Au) more or less loaded (<1% to 10% by weight of metal) on graphitic matrix. These 84 85 systems have significant differences in electronic densities between the support and the metal phase and 86 therefore a significant contrast is obtained in anomalous Small angle X-ray scattering (ASAXS). Polizzi 87 et al. [19] have also proposed a detailed critical analysis of the methodologies selected for these studies, 88 insisting on the need for protocols based on numerous controls of the measurement parameters in order to 89 ensure a good control of the uncertainties, which is essential in the case of ASAXS measurements. The 90 ASAXS technique is nowadays seldom used for the characterization of the active phases of catalysts, 91 even if it is well appropriate for a three-phases system (pores, alumina and cobalt nanoparticles). 92 Moreover, scattering techniques have already proved their possibilities to characterize the size and 93 morphology of aggregates [20]. Hence, this technique has the advantage of being able to characterize 94 objects with a size between one nanometer and a few hundred nanometers, thus covering the scales of 95 isolated particles and aggregates of particles. In addition, the analysis can be performed on samples
96 embedded in their waxes, which is very relevant for the case of used FT catalysts.

97

98 In order to study the morphology of the aggregates, advanced techniques such as electron tomography can 99 be successfully applied. Based on TEM technique, electron tomography has been developed during the 100 last decades. The principle is to reconstruct a three-dimensional structure from a series of images (or 2D 101 projections) acquired by rotating the sample in the TEM [21] [22]. In the field of heterogeneous catalysis, it was first used to study the shape and the connectivity of mesoporous networks in catalysts supports 102 such as mesostructured silica [23][24] or zeolites [25][26][27][28]. Individual or supported nanoparticles 103 104 can also be characterized by electron tomography: in the first case, the morphology (including faceting) of the particles is sought [23], in the latter case, the location inside the support can be explored. As an 105 106 example, Zecevic et al examined the location of Pt in USY zeolites crystals, showing the presence of Pt particles inside the microporosity [29,29,30]. If the contrast between the nanoparticles and the support is 107 108 too weak in bright field TEM, chemical sensitive techniques have to be used [21]. EFTEM is mainly employed for characterization of biphasic supports [31] or low Z materials [32]. HAADF-STEM is the 109 110 preferred technique for supported catalysts [33]. Compared to bright field TEM mode, HAADF-STEM enhances the contrast between the metallic nanoparticles and the oxide support, in as much as the image 111 intensity is proportional to $Z^{1.7}$ [34]. Arslan and al. [35] studied oxide Co aggregates supported on two 112 different alumina supports. They showed a morphology of fully interlocking aggregates/support in the 113 114 less selective catalyst.

115 Combining SAXS analysis and electron tomography has also been performed by Gommes et al. [36] on 116 Cu catalysts supported on ordered porous silica and allowed illustrating the mesoscale distribution of 117 nanoparticles within supports displaying hierarchical porosity. This study has demonstrated the strong 118 interest of coupling these two kinds of techniques and this approach could be extended to Fischer-Tropsch 119 catalysts provided that SAXS is tuned to ASAXS to specifically focus on cobalt particles and aggregates 120 size.

121

The purpose of the present study is to propose a new approach to characterize the cobalt active phase in FT catalysts. We combined two analytical techniques : ASAXS and electron tomography, giving complementary insights on the microstructure and size distribution of both Co nanoparticles and aggregates. A better knowledge of Co aggregates morphology will help to understand their impact on the catalyst performance. This approach, carried out on a Co/alumina-silica catalysts studied first at the oxide state and then at the reduced state, allowed to highlight the mechanisms involved at the nanoscale duringthe reduction step.

129

130 2 Experimental section

131 **2.1 Catalyst preparation**

For this study, we have chosen to focus on two cobalt-based catalysts. They were prepared on a Siralox 5 silica-alumina support by incipient wetness impregnation of a nitrate cobalt solution, to obtain an amount of 15 wt.% of cobalt. After impregnation, the catalysts were dried and calcined in air at 400°C. The first sample is the oxide catalyst. The second one is obtained after reduction under pure H_2 flow at 400°C during 16h. After reduction, the reduced catalyst is divided into two fractions: the first fraction is passivated under air in order to be analyzed by electron microscopy and the second fraction is embedded in waxes in order to prevent the re-oxidation and to be analyzed in ASAXS.

139

140 2.2 Bulk characterization

141 XRD measurements were performed on the oxide sample on a PANalytical diffractometer. The most 142 intense Co_3O_4 line (311) was used to calculate the average Co_3O_4 crystallite size with the Scherrer 143 equation. The mean oxide crystallite size was found to be 16.6 ± 2.0 nm.

144 XANES analysis was also carried out on the reduced sample embedded in waxes on the ROCK beamline 145 of SOLEIL synchrotron to determine the reduction rate, which is the ratio between the Co^o content (wt 146 .%) and the total Co content. XANES spectrum were decomposed thanks to reference sample spectrum. It 147 has underlined the fact that the reduced sample was partially reduced, with a reduction rate of 47% (see 148 supplementary data, figure 1).

149

150 2.3 TEM and HAADF-STEM observations

By bright-field TEM, the contrast between small cobalt nanoparticles and the silica-alumina support is not sufficient to perform precise size measurements. Thus, cobalt particles size was measured by dark field transmission electron microscopy (DF-TEM). Observations were performed using a JEOL 2100F FEG S/TEM, operated at 200 kV. Samples were crushed in ethanol and sonicated. A drop of the suspension was deposited on a Cu grid covered with a holey carbon membrane. DF-TEM observations are performed using the diffraction spots corresponding respectively to the (311), (660) or (220) planes in Co₃O₄ and (100), (002), (101) or (102) planes in hcp Co°. The particles size was measured on a minimum of 200 Co
crystallites.

Cobalt aggregates were observed and quantified using HAADF-STEM (high angle annular dark field detector in scanning transmission electron microscopy). Before observation, the samples were embedded in an epoxy resin and thin slices (about 70 nm) were cut by ultramicrotomy and transferred onto a Cu grid covered with a holey carbon membrane. Observations were performed using a JEOL 2100F FEG S/TEM, operated at 200 kV. The aggregates size was measured on a minimum of 200 aggregates on several catalysts grains.

165

166 **2.4 Electron tomography**

167 Electron tomography was performed using a JEOL 2100F FEG S/TEM, operated at 200 kV. Samples 168 were prepared as for DF-TEM observations, except that a drop of a gold colloidal suspension was 169 deposited on the grid for further alignment.

Images were taken in HAADF-STEM mode, in order to increase the contrast between aggregates and 170 silica-alumina support. Thus, the morphology of the sole aggregate can be visualized without the 171 contribution of the support. Acquisitions were performed using GATAN Digiscan and Digital 172 Micrograph, by rotating the sample inside the microscope over an angular range of $\pm 60^{\circ}$ using a 1° tilt 173 increment, in equal scheme, which corresponds to 122 pictures for the "oxide sample", and over an 174 angular range of +69° to -67° using a 2° tilt increment, in Saxton scheme, which corresponds to 93 175 pictures for the "reduced sample". Dynamic focus adjustment was used for acquisition of the tilt series. 176 Alignment of the HAADF-STEM tilt series was performed thanks to cross-correlation and gold fiducial 177 markers, 3D reconstruction was realized by filtered backprojection (via etomo in IMOD[37,38,38]). 178 Segmentation based on grey-levels histograms was undertaken using AVIZO[®] software. To facilitate 179 180 segmentation, a pre-processing of the reconstructed volume by a "flowing bilateral filter" [39] was performed. This filter smooths homogeneous regions while preserving the edges of the objects. 181

182 Morphological geodesic operators were applied in order to extract and quantify the porosity inside the 183 aggregates. Co dispersion in the aggregates was calculated by the ratio between the number of voxels at 184 the surface of the aggregate to the total number of voxels in the aggregate.

185

186 **2.5 ASAXS analysis**

- 187 2.5.1 Theoretical aspects of ASAXS
 - 6

For the case of spherical particles with a radius R dispersed in a homogeneous matrix, the scattered intensity is given by:

$$I(q) = \frac{N_p}{V_s} \left(n_p^2 f_p^2 - n_m^2 f_m^2 \right) r_e^2 \int_0^\infty P(R) V(R)^2 F(q,R)^2 \Psi(q) dq$$
(1)

191

with N_p the number of tparticles, V_s the sample volume, n_p and n_m the number density of the atoms in the particles and in the matrix, f_p and f_m the atomic form factors and r_e the Thomson radius ($r_e = 0.282 \ 10^{-12}$ cm).

195 P(R) is the size distribution function, F(q, R) is the form factor and $\Psi(q)$ is the structure factor and 196 equals to 1 when the particles are well-dispersed.

197

For the case of metal particles supported on a porous matrix, for instance on silica-alumina support, the simple SAXS analysis (recording at only one energy) is not sufficient to distinguish and to characterize the nanoparticles as the porous support also contributes significantly to the signal because of the scattering of the pores.

Hence, to isolate the signal specific of each phase, it is therefore interesting to vary the energy of the incident beam and to perform the measurements near and far the metal (cobalt in our case) absorption edge.

205 It is recalled that the atomic form factor of a species *j* can be written:

206

$$f_j(E) = f_0 + f'_j(E) + if''_j(E)$$
⁽²⁾

207 with $f_0 = Z$, the atomic number

The values of $f'_{j}(E)$ and $f''_{j}(E)$ are indicated in the Table 1. We can consider that $f''_{j}(E)$ does vary significantly only after the absorption edge.

210

211Table 1. Energy-dependent parts of the atomic scattering factor f' and f'' [40,41], near the Co K-212edge of absorption (7709 eV).

	Energy (eV)	f'	f"
E ₁	7440	-3.082	0.493
$\mathbf{E_2}$	7648	-4.744	0.470
E_3	7694	-6.356	0.465

213

Considering a system of particles p supported on a porous support s and with air filling the pores, the scattered intensity can be written thanks to the partial structure factors S_{pp} , S_{ss} and S_{sp} described by

218
$$\langle I(q,E)\rangle = \left|f_p(E)\right|^2 n_p^2 r_e^2 S_{pp}(q) + |f_s|^2 n_s^2 r_e^2 S_{ss}(q) + 2\Re (f_p(E)f_s^*) n_p n_{ps} r_e^2 S_{sp}(q)$$
(3)

219 The difference between the intensities measured at two different energies E_1 et E_2 leads to :

220

$$\langle I(q, E_1) \rangle - \langle I(q, E_2) \rangle = n_p^2 r_e^2 \left(\left| f_p(E_1) \right|^2 - \left| f_p(E_2) \right|^2 \right) S_{pp}(q) + 2f_s \left(f_p(E_1) - f_p(E_2) \right) n_p n_s r_e^2 S_{sp}(q)$$

$$= 2n_p r_e^2 \left(f_p(E_1) - f_p(E_2) \right) \cdot \left(n_p \overline{f_p} S_{pp}(q) + n_s f_s S_{sp}(q) \right)$$

$$= 2n_p^2 r_e^2 \overline{f_p} \left(f_p(E_1) - f_p(E_2) \right) \cdot \left(S_{pp} + \alpha S_{sp} \right)$$

$$(4)$$

222

221

223 where $\overline{f_p} = \frac{f_p(E_1) + f_p(E_2)}{2}$ is the mean value of $f_p(E)$ and $\alpha = \frac{n_s f_s}{n_p \overline{f_p}}$

224
$$S_{pp}(q) = \frac{N_p}{V_s} \int_0^{+\infty} P_p(R_p) V(R_p)^2 F(q, R_p)^2 dR_p$$
(5)

225
$$S_{mp} = \frac{N_p}{V_s} \iint_0^\infty P_p(R_p) P_s(R_s) V(R_p) V(R_s) F(q, R_p) F(q, R_s) \frac{\sin(q(R_p + R_s))}{q(R_p + R_s)} dR_p dR_s$$
(6)

As a first approach, we can consider that the term αS_{sp} is negligible before S_{pp} if $\alpha < \frac{\langle V(R_p)^2 \rangle}{\langle V(R_p) \rangle \langle V(R_s) \rangle}$. 227 In our case:

- 228 we have calculated for E_1 and E_2 energies, an α close to 0.55
- 229 the size of the pores and of the particles are in the same order of magnitude, so that $\frac{\langle V(R_p)^2 \rangle}{\langle V(R_p) \rangle \langle V(R_s) \rangle}$ is 230 higher than 1.

It can be concluded that in the present case of the cobalt-based catalysts, the contribution of the supportparticle interaction term is negligible. Indeed, as explained by Morfin et al [10], the particles are big enough and their volume fraction is small compared to the aluminum support consequently the probability of interference between the cobalt particles and the support is low.

235 It follows :

236
$$\langle I(q,E_1)\rangle - \langle I(q,E_2)\rangle \approx 2n_p^2 r_e^2 \overline{f_p} \left(f_p(E_1) - f_p(E_2) \right) \cdot S_{pp}(q) \tag{7}$$

237
$$\langle I(q,E_1)\rangle - \langle I(q,E_2)\rangle \approx \frac{N_p}{V_s} n_p^2 r_e^2 \left(\left| f_p(E_1) \right|^2 - \left| f_p(E_2) \right|^2 \right) \int_0^{+\infty} P_p(R_p) V(R_p)^2 F(q,R_p)^2 dR_p$$
(8)

Finally, when the porosity is filled with waxes, the difference in electron density between particles and waxes should be taken as contrast factor instead of the electron density of the particle itself. Therefore the previous expression becomes :

242
$$\langle I(q,E_1) \rangle - \langle I(q,E_2) \rangle \approx$$

243 $\frac{N_p}{V_s} r_e^2 \left(\left| n_p f_p(E_1) - n_m f_m(E_1) \right|^2 - \left| n_p f_p(E_2) - n_m f_m(E_2) \right|^2 \right) \int_0^{+\infty} P_p(R_p) V(R_p)^2 F(q,R_p)^2 dR_p$ (9)

244

245 2.5.2 ASAXS measurements

246 For ASAXS analysis, the reduced sample was directly embedded in waxes without air exposition in order 247 to prevent reoxidation. The oxide sample was also put into waxes in order to keep the same mode of 248 sample preparation. Then, pellets with a thickness of approximatively 300µm were prepared. Anomalous 249 small-angle X-ray scattering experiments have been performed on the SWING beamline at synchrotron 250 SOLEIL. It involves characterizing cobalt nanoparticles supported on a porous silica-alumina by recording SAXS images around its absorption edge. The advantage of performing ASAXS analyzes is 251 252 that it suppresses the contribution of the porous support. For this purpose, it is necessary to be close to the cobalt absorption threshold in order to vary the diffusion factor of Co and thus of the particles, but also to 253 254 keep constant the diffusion factor of the support. Hence, the scattered intensity was measured at three 255 different energies (see Table 1) 7440, 7647 and 7694eV, slightly below the cobalt K-edge of absorption. 256 The incident beam energy has been calibrated with a metallic cobalt foil. To cover the largest q range, two sample-detector distances (1000 mm and 6000 mm) have been selected. Furthermore, to increase the 257 sensibility of the photons counting on the peripheral part of the detector, two kinds of beamstop (a small 258 259 beamstop with a size of $2x1 \text{ mm}^2$ and larger beamstop with a size of $3x3 \text{ cm}^2$) were used. Using the large beamstop, allowed to recorded 2D images with longer exposure time, by masking the strong diffusion 260 261 around the small beamstop, in order to get better statistics for high q values. In total, four different 262 configurations (detector-sample distance, beamstops) have been used.

The scattering images were recorded using a AVIEX PCCD170170 detector. To increase the statistic of the scattering intensity, thirty images were acquired for each energy and at each detector-sample configuration. The acquisition time was function of the configuration and was chosen in order not to saturate the detector.

267 Through 1D reduction, raw data were corrected with respect to: acquisition time, geometrical effects like 268 the projection of the detector plane on the sphere with radius equal to the sample-detector distance, 269 detector sensitivity (3ADU/ph, Noise≈2ADU),the incoming-photon flux and finally averaged to increase the 270 statistic. The scattering background coming from all components in the beamline and from the air was 271 measured before each series of acquisition, for each energy and at each configuration. The sample 272 transmission was calculating thanks to the measurement of the incoming and transmitted intensities. It 273 allowed correction of the intensity from the real thickness (*e*) of the sample, calculated thanks to the 274 composition of the sample obtained by X-Ray Fluorescence and with the following equation:

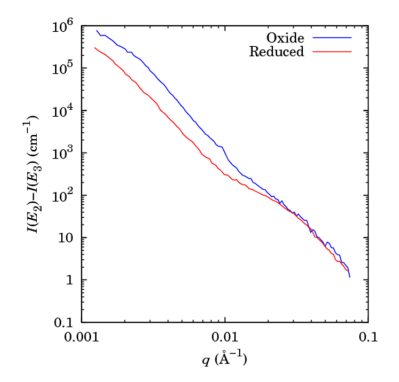
$$e = -\frac{\ln T}{\mu_{sample}} \tag{10}$$

- 275 276
- 277
- 278 T being the transmission and μ_{sample} the linear absorption coefficient (cm⁻¹).
- 279

In order to calibrate the intensity in absolute unities, i.e expressed as differential scattering cross section per unit volume in cm⁻¹, a calibrated NIST glassy carbon [43], was measured at each energy and each configuration. A correction factor has been calculated from the ratio between the NIST data and the experimental data.

284 2.5.3 Modelling of ASAXS data

The ASAXS results for both oxide and reduced catalysts are given in Figure 1. The plotted curves were obtained from the subtraction of the intensity measured at $E_2 = 7647$ eV and the intensity measured at E_3 = 7694 eV near the cobalt K-edge of absorption. Noticeable differences can be observed in the q-range between 2.3 10⁻² Å⁻¹ and 1.2 10⁻³ Å⁻¹.





291 Fig

292

Figure 1. ASAXS curves obtained on oxide and reduced samples

In order to evaluate the size distribution of the cobalt particles, a nonlinear least squares adjustment of the ASAXS signal is performed. The assumptions made for the choice of the regression model are listed below.

Given the shape of the curves (two inflections of the scattered signal), the experimental data could not be fitted with only one function of size distribution. It appears that two kinds of cobalt particles exist in the catalyst, at the oxide and at the reduced state, at two different scales. Based on the electron microscopy observations, these two populations are attributed to isolated, dense and crystalline particles at the small scale, and to larger, polycrystalline and moderately dense aggregates at large scale. This assumption will be corrobated in this paper.

302

For the cobalt phase, we thus consider that the scattering objects can be porous. The porosity is so called ε and is filled with waxes. In the case of the crystalline particles, $\varepsilon = 0$ and in the case of cobalt aggregate, ε 0.5 > 0.5

The number of cobalt atoms in the scattering object and the number of the matrix atoms are respectivelydefined by:

$$N_{Cox} = \frac{N_A \times \rho_{Cox}}{M_{Cox}} \times V \times (1 - \varepsilon)$$
(11)

$$N_m = \frac{N_A \times \rho_m}{M_m} \times V \times \varepsilon \tag{12}$$

With $\text{Co}_{x} = \text{Co}_{3}\text{O}_{4}$ in the case of oxide cobalt and $\text{Co}_{x} = \text{Co}^{0}$ in the case of metallic cobalt, V the volume of the scattering object, N_{4} the Avogadro number (6.022×10²³ mol⁻¹), M_{Cox} and M_{m} the molar masses in g/mol, ρ_{Cox} and ρ_{m} the densities of both phases in g/cm³.

314 Hence, the scattering factor of a cobalt particle is defined by :

315
$$n_p f_p = \left(\frac{N_A \times \rho_{Cox}}{M_{Cox}} \times f_{Cox}\right) \times (1 - \varepsilon) + \left(\frac{N_A \times \rho_m}{M_m} \times f_m\right) \times \varepsilon$$
(13)

316 It follows:

317
$$n_p f_p - n_m f_m = \left(\frac{N_A \times \rho_{Cox}}{M_{Cox}} \times f_{Cox} - \frac{N_A \times \rho_m}{M_m} \times f_m\right) \times (1 - \varepsilon) = (n_{Cox} f_{Cox} - n_m f_m) \times (1 - \varepsilon) \quad (14)$$

319 The term $\frac{N_p}{V_c}$ of the equation (9) can be expressed as:

320
$$\frac{N_p}{V_s} = \frac{w_{Cox} \times \rho_s}{\rho_{Cox}} \times \frac{1}{1 - \varepsilon} \times \frac{1}{\langle V \rangle}$$
(15)

321

With $\langle V \rangle$ the mean volume of the particle aggregate (cm⁻³), w_{Cox} the weight content of the cobalt phase (wt.%), ρ_s the sample density in g/cm³.

The particles are considered to be spherical and isolated: as so, the structure factor is not taken into account while the form factor is. Thus the form factor for a spherical particle is:

326
$$F^{2}(q, R_{p}) = \left[3 \frac{\sin(q, R_{p}) - qR_{p}\cos(q, R_{p})}{(q, R_{p})^{3}}\right]^{2}$$
(16)

327 and
$$V(R_p) = \frac{4}{3}\pi R_p^3$$
 (17)

328

329 The size distributions can be represented by a sum of log-normal distribution:

330
$$P_i(R_p) = \frac{1}{\sqrt{2\pi}R_p\sigma_i} exp\left(-\frac{\left(\ln R_p - \mu_i\right)^2}{2\sigma_i^2}\right)$$
(18)

331

In order to calculate the number and volume average sizes from the adjusted log-normal distribution, the distribution moments M_n must be known [44]:

$$M_n = exp\left(\mu n + \frac{n^2 \sigma^2}{2}\right) \tag{19}$$

335 The volume size distribution P_v is defined by:

336
$$P_{iv}(R_p) = \frac{R_p^{3} P(R_p)}{M_3}$$
(20)

338 The number, the surface and the volume mean sizes can be calculated:

339
$$R = \frac{M_1}{M_0} = \exp\left(\mu + \frac{\sigma^2}{2}\right), R_{surf} = \frac{M_3}{M_2} = \exp\left(\mu + \frac{5\sigma^2}{2}\right) \text{ and } R_{vol} = \frac{M_4}{M_3} = \exp\left(\mu + \frac{7\sigma^2}{2}\right) (21)$$

Given the shape of the curves (two inflections of the scattered signal), we chose to model the distributionof cobalt by two log-normal distributions:

$$\langle I(q, E_{1}) \rangle - \langle I(q, E_{2}) \rangle \\\approx r_{e}^{2} \left(\left| n_{p} f_{p}(E_{1}) - n_{m} f_{m}(E_{1}) \right|^{2} - \left| n_{p} f_{p}(E_{2}) - n_{m} f_{m}(E_{2}) \right|^{2} \right) \\\times \left[\frac{N_{p1}}{V_{s}} \int_{0}^{+\infty} P_{1}(R_{p}) V(R_{p})^{2} F(q, R_{p})^{2} dR_{p} + \frac{N_{p2}}{V_{s}} \int_{0}^{+\infty} P_{2}(R_{p}) V(R_{p})^{2} F(q, R_{p})^{2} dR_{p} \right]$$

$$(22)$$

343 With

344
$$\frac{N_{p_1}}{V_s} = \frac{w_{Cox} \times \rho_s}{\rho_{Cox}} \times w_1 \times \frac{1}{1 - \varepsilon_1} \times \frac{1}{\langle V_1 \rangle} and \frac{N_{p_2}}{V_s} = \frac{w_{Cox} \times \rho_s}{\rho_{Cox}} \times w_2 \times \frac{1}{1 - \varepsilon_2} \times \frac{1}{\langle V_2 \rangle}$$
(23)

345

342

346 $\langle V \rangle$ is the mean volume such as :

347
$$\langle V \rangle = \frac{4}{3} \times \pi \times \int R_p^{3} P(R) dR = \frac{4}{3} \times \pi \times M_3 = \frac{4}{3} \times \pi \times exp\left(3\mu + \frac{9\sigma^2}{2}\right)$$
(24)

348

349 Supposing that the first particle population corresponds to dense, crystalline and isolated particles and 350 that the second one is attributed to aggregates partially dense, the previous equation becomes:

351

$$\langle I(q, E_i) \rangle - \langle I(q, E_j) \rangle = w_{Co_x} \times w_1 \times \frac{\rho_s}{\rho_{Co_x}} \times \Delta \rho_{Co_x}^2(E_i, E_j)$$

$$352 \qquad \times \left[\frac{1}{\langle V_1 \rangle} \int_0^\infty P_1(R_p) V^2(R_p) F^2(q, R_p) dR_p + \frac{w_2(1 - \varepsilon_2)}{w_1} \times \frac{1}{\langle V_2 \rangle} \int_0^\infty P_2(R_p) V^2(R_p) F^2(q, R_p) dR_p \right]$$
(25)

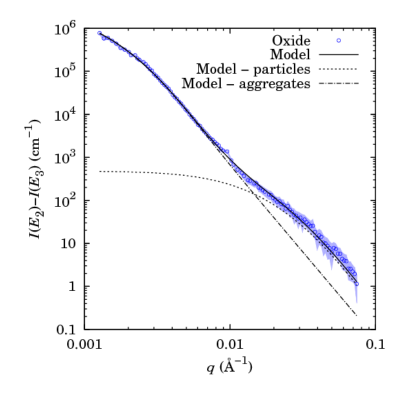
353 With w₁ and w₂ the cobalt weight percent of the first and second population of particles, and

354
$$\Delta \rho_{Cox}^2 (E_i, E_j) = (n_{Cox} f_{Cox} (E_i) - n_m f_m)^2 \times r_e^2 - (n_{Cox} f_{Cox} (E_j) - n_m f_m)^2 \times r_e^2 \quad (26)$$

355 356

This model allows estimating the size distribution of the particles (μ_1 , σ_1), the size distribution of the aggregates (μ_2 , σ_2) and the parameter $\frac{w_2(1-\varepsilon_2)}{w_1}$ which depends on the ratio between the cobalt content included in the first and in the second population, but also on the porosity of the aggregates. An example of fit is shown on Figure 2 for the oxide sample. The first part of the curve (from 10^{-1} Å⁻¹ to 1.4 10^{-2} Å⁻¹) is essentially due to the contribution of the first population of small-size particles whereas the second part of the curve (from 1.4 10^{-2} Å⁻¹ to 1.2 10^{-3} Å⁻¹) is due to the contribution of the second population of larger objects called aggregates.

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Figure 2. Fit of the experimental ASAXS data of the oxide sample.

368 **3 Results**

369 **3.1 Particle scale**

370 Cobalt particle size distribution was measured by dark-field transmission electron microscopy and also

371 obtained by ASAXS.

372 Figure 3a presents a representative image of dark-field TEM of cobalt nanoparticles supported on silica-

alumina. Figure 3b presents the size distribution in both oxide (blue) and reduced (green) catalysts. Both

374 size distributions present the same general shape, and few differences are observed.

375 Besides, the volume size distribution (P_v) of the particles obtained by ASAXS for both catalysts thanks to

the fit of the experimental data are plotted on Figure 4. The two size distributions are slightly different as

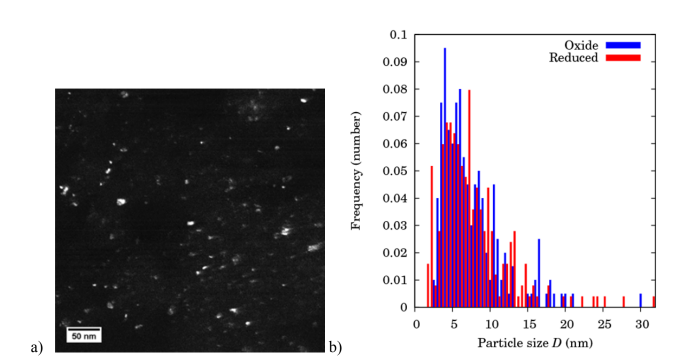
377 revealed by the μ and σ parameters of the lognormal-law (3.29 and 0.56 for the oxide sample versus 3.56 378 and 0.44 for the reduced sample) however the differences are faint.

379

In order to compare TEM, ASAXS measurements and XRD mean size, all results are expressed by the 380 381 volume mean particle radius. ASAXS, TEM and XRD data on oxide samples are totally consistent, 382 considering uncertainties of both techniques. It is noteworthy that a very good agreement is obtained 383 between ASAXS and TEM results for the oxide sample contrary to the case of the reduced sample. This 384 can be accounted for by the 47% reduction rate, measured by XANES. Indeed, less than half of the cobalt is reduced and thus in the metallic Co^o phase. Moreover, it is known that the small particles are less 385 386 reducible than the big ones. Due to its principle, the dark-field TEM allows observing and counting solely Co^o particles, whereas the ASAXS technique characterizes all the cobalt either oxide or metallic. This 387 388 could explain the mean size differences obtained with both techniques, and also the higher mean size 389 particle measured on the reduced sample by TEM compared to the oxide catalyst.

These ASAXS analyses tend to show a decrease in size during reduction, which agrees with the fact that the reduced Co^0 particles are smaller than the oxide Co_3O_4 particles due to the crystallographic structure contraction of around 0.8. However it is difficult to conclude on the effect of the reduction of the particle size as it seems to be very tight and hidden by the large distribution of the particle size and the significant heterogeneity of the samples.





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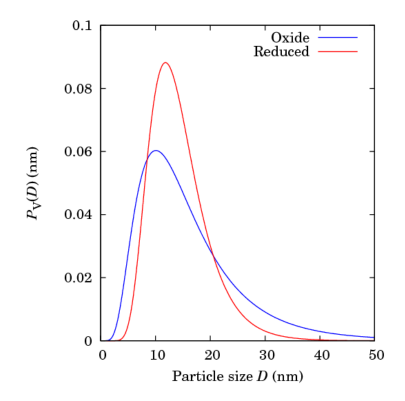
397

Figure 3 : a) Dark-field TEM image of oxide Co/silica-alumina. B) Particles size distribution expressed by number frequency: in oxide catalyst (blue) and reduced catalyst (red)

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 Table 2: Cobalt particle sizes measured by different techniques

	Oxide catalyst	Reduced catalyst
DF TEM – volume mean size (nm)	16.1 ± 2.4	18.0 ± 2.7
ASAXS - volume mean size (nm)	16.2	14.5
XRD – volume mean size (nm)	16.6 ± 2.0	/





406 Figure 4 : Volume size distribution (Pv) obtained for the oxide and the reduced catalysts by
 407 ASAXS.

408

409 **3.2** Cobalt aggregates characterization by HAADF-STEM and electron tomography

410 Cobalt aggregates in the catalysts grains were observed at different scales. First of all, SEM observations 411 on several grains of the oxide catalyst were performed and have showed a strong heterogeneity in the Co 412 aggregates repartition inside a grain and from on grain to another. Furthermore, the scale of 413 characterization was too large to precisely their size. Thus, HAADF-STEM observations were carried out 414 on ultramicrotomy cuts. Representative images for the oxide and reduced catalysts are presented on 415 Figure 5. They have revealed that aggregates repartition and even size can strongly vary from a grain to

416 another. Nevertheless, measurements were performed on more than 200 aggregates in each catalyst and

417 their size distribution is presented on Figure 6. The mean size in oxide catalyst corresponds to 52 nm and

418 to 54 nm in reduced one.

ASAXS analysis has also allowed to obtain the number size distribution (P_N) of the aggregates for both catalysts thanks to the fit of the experimental data (see Figure 7). The maximum of the distribution in the reduced sample is slightly shifted to smaller size than in the oxide one. The mean number size of the oxide catalyst is 85 nm versus 49 nm for the reduced one. However the standard deviation is quite high, meaning that the aggregates size can vary from 20 to almost 200 nm. : the lognormal-law parameters (μ and σ that are, respectively) were found to be 5.91 and 0.55 for the oxide catalyst and 5.16 and 0.83 for the reduced one.

426 ASAXS and STEM measurements are in good agreement concerning the aggregates mean size of the reduced sample (54 and 49 nm) but a small difference exists for the oxide sample, even if the ASAXS and 427 428 STEM values (52 and 85 nm) are in the same order of magnitude. However, we have discussed that the 429 sample heterogeneity and the cobalt distribution from one grain to another was significant, which could 430 explain the differences. Considering heterogeneity of the samples and standard deviations of the 431 distribution, it can be concluded that the aggregate diameter does not evolve significantly during reduction. Hence, reduction under H₂ at 400°C does not lead to aggregate sintering neither to large scale 432 redistribution of cobalt in the support. Cobalt aggregates microstructure was studied by dark-field 433 434 transmission electron microscopy. Both electron diffraction patterns and DF-TEM images reveal an important change of morphology during reduction. Indeed, oxide cobalt aggregates present a dense 435 436 structure with indistinct particles, i.e consisting of coherent domains having rather the same crystalline 437 orientation but with some structural stacking faults (Figure 9). On the other hand, metallic cobalt 438 aggregates present a clearly polycrystalline structure (Figure 10), and crystals forming the aggregates 439 even seem to be disjoint. It is confirmed by high magnification HAADF-STEM images (Figure 8). In 440 oxide catalyst, aggregates look compact, with a uniform contrast, whereas in reduced catalyst, individual 441 particles forming the aggregates on metallic form appear clearly and some porosity is visible in between.

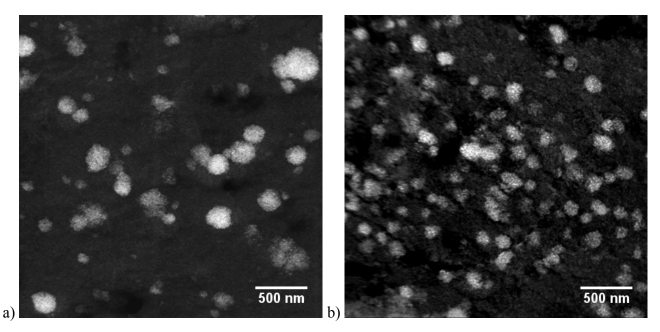


Figure 5 : HAADF-STEM images of a) oxide catalyst ; b) reduced catalyst.

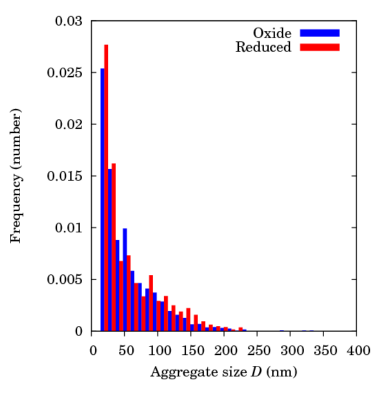


Figure 6 : Cobalt aggregates size distribution, measured by HAADF-STEM, expressed by number
 frequency: in oxide catalyst (blue) and reduced catalyst (red)

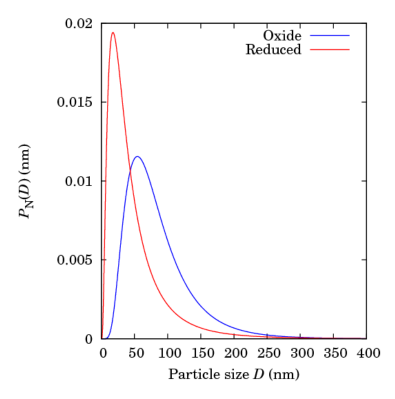


Figure 7 : Number aggregates size distribution (P_N) obtained by ASAXS for the oxide and the reduced catalysts.

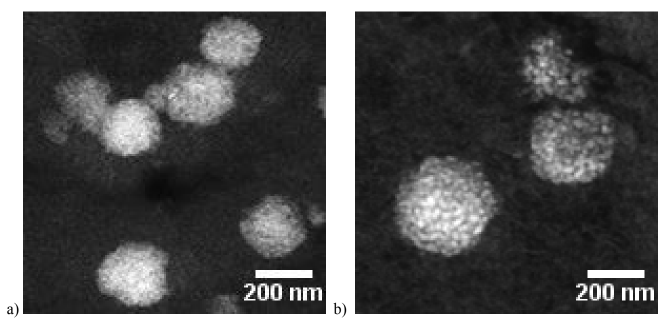


Figure 8 : High magnification HAADF-STEM images of a) oxide catalyst ; b) reduced catalyst.

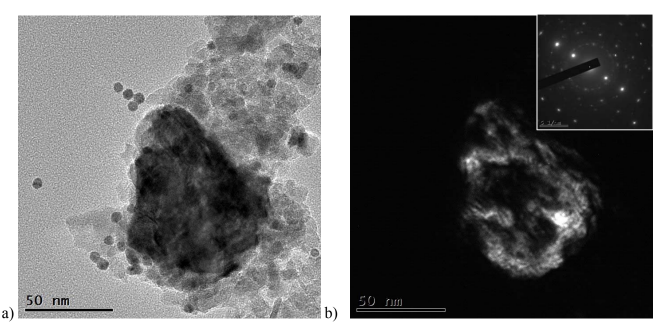
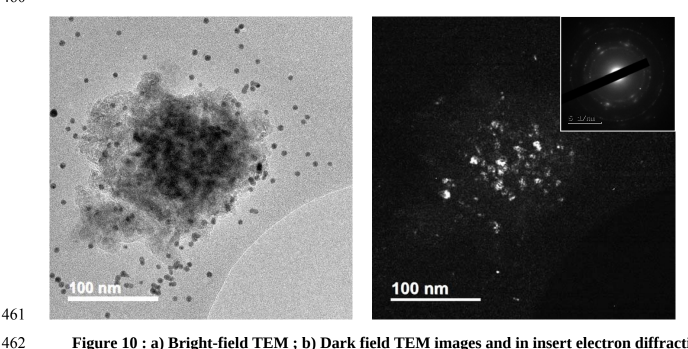


Figure 9 : a) Bright-field TEM ; b) Dark field TEM images and in insert electron diffraction 458 459 pattern of a cobalt aggregate in the oxide catalyst. 460



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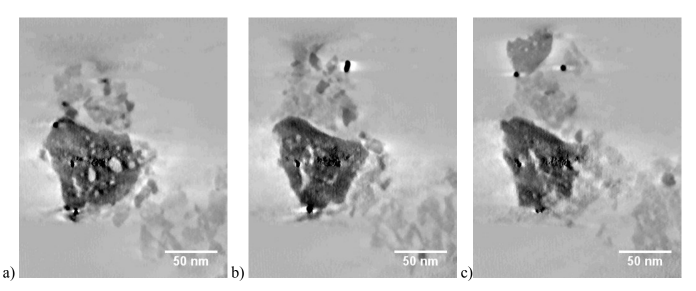
Figure 10 : a) Bright-field TEM ; b) Dark field TEM images and in insert electron diffraction pattern of a cobalt aggregate in the reduced catalyst.

465 Combination of several TEM imaging modes gives an insight of a strong modification of the aggregates 466 morphology during reduction. Nevertheless, these are 2D techniques and they are not able to fully 467 describe their morphology. Three-dimensional morphology can be described by electron tomography.

468 Two different aggregates for each sample were studied by electron tomography. They were chosen 469 according to their size. Too small aggregates (circa 20 nm) may be mistaken to large particles, too large 470 aggregates (> 100 nm) are too thick for electron tomography acquisitions. 50 nm-diameter aggregates
471 were selected, corresponding to the mean size obtained from HAADF-STEM images.

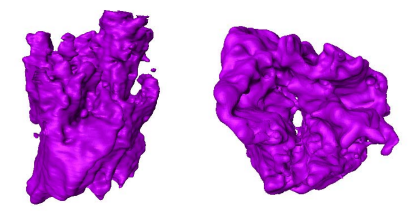
Figure 11 depicts cross sections of the reconstructed volume of a cobalt aggregate in the oxide catalyst. The aggregate appears in dark in the image (the contrast was inverted after the acquisition and before reconstruction) and the support is visible in light grey. The aggregate is a compact and dense phase of cobalt, with some spherical or cylindrical closed pores in its centre.

476



- Figure 11 : a-c) Slices from electron tomography reconstructions for a cobalt aggregate in the oxide
 catalyst.
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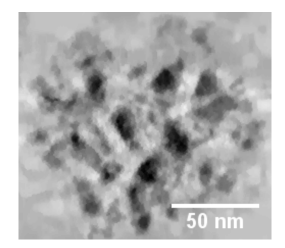
Figure 12: Model of a cobalt aggregate in the oxide catalyst, obtained by segmentation of the
 electron tomography reconstruction. Two different orientations.

485 Segmentation of the reconstructed volume allowed to model the aggregate. A 3D representation of the 486 model aggregate is presented on Figure 12. Only the cobalt phase was segmented independently to the 487 silica-alumina support. Indeed, the contrast of the support compared to porosity is too weak to be 488 segmented without important error. It confirms the compact structure of the aggregate. In particular, the 489 external surface of the aggregate is very smooth. Accessibility of cobalt is a key factor for catalysis. We defined two parameters to estimate the accessibility of cobalt in aggregates. The first one, called 490 491 "aggregate dispersion", is calculated as the ratio between surface cobalt to total cobalt and the second one 492 is the porosity inside the aggregate. From the aggregate model obtained by electron tomography, the 493 "aggregate dispersion" is estimated to 18% and the porosity to 35%. "Aggregate dispersion" takes into account the surface roughness of the aggregate and is different from dispersion classically obtained by 494 495 chemisorption.

Figure 13 depicts a cross section of the reconstructed volume of a cobalt aggregate in the reduced catalyst. The aggregate appears in dark in the image. Here, the support is slightly visible compared to porosity. The aggregate is a porous and airy phase of cobalt, composed of almost independent particles with large inter-particle porosity.

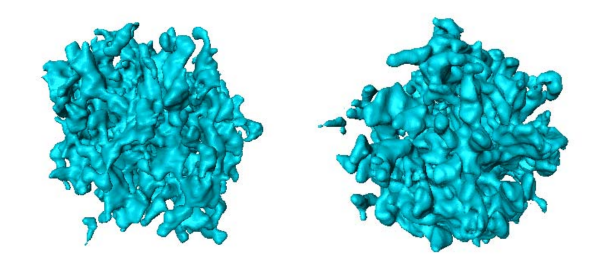
A 3D representation of the model aggregate, obtained by segmentation of the reconstructed volume is presented on Figure 14. Here again, only the cobalt phase was selected for modeling independently from the support. The sponge-like structure of the cobalt aggregate in the reduced sample is clearly highlighted. Contrary to the compact morphology on the oxide state, the aggregate presents a large external surface, leading to high accessibility. From the aggregate model, the "aggregate dispersion" is estimated to 27% and the porosity to 65%.

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507

508Figure 13 : Slice from electron tomography reconstructions for a cobalt aggregate in the reduced509catalyst.



512Figure 14: Model of a cobalt aggregate in the reduced catalyst, obtained by segmentation of the513electron tomography reconstruction. Two different orientations.

Besides these deep electron tomography characterizations, ASAXS also provides information on the 515 516 aggregate structure. Indeed, the model used to fit the ASAXS data makes it possible to calculate the $\frac{w_2(1-\varepsilon_2)}{w_1}$ parameter. It was found to be 3.05 for the oxide sample and 1.11 for the reduced sample. 517 Supposing that the ratio between the cobalt content included in the first and in the second populations 518 519 (particles and aggregates) does not evolve during the reduction step, we can suggest that the ratio $\frac{(1-\epsilon_2)_{reduced}}{(1-\epsilon_2)_{oxide}}$ equals to 0.364, which implies that the compacity of the reduced aggregates ϵ_2 is almost 2.8 520 521 times less significant than the one of the oxide aggregates. Hence, ASAXS analysis also lead to conclude 522 that during the reduction, the aggregate structure becomes airier and more porous.

523 Furthermore, to check the good agreement between ASAXS and electron tomography, setting the oxide aggregate porosity equal to 35% as calculated by electron tomography, the ratio $\frac{(1-\epsilon_2)_{reduced}}{(1-\epsilon_2)_{oxide}}$ of 0.364 524 525 obtained by ASAXS leads to a calculated aggregate porosity of 76% for the reduced sample. Even if the 526 porosity is not exactly the same when calculated by electron tomography or by ASAXS, the same trend is 527 observed with the same order of magnitude. Both techniques are in good agreement but each provides different benefits. Electron tomography provides local information and could suffer from 528 529 representativeness issues, it allows visualizing the object of the study and justify the kind of model 530 employed to fit the ASAXS data (spherical particles and aggregates, log-normal distribution of size). 531 Besides, ASAXS technique is less familiar but as the advantage to be a bulk analysis which provides 532 representative information of the whole sample. Hence, both technique are complementary.

511

533 They underline the strong effect of the reduction step on the aggregate structure: the oxide aggregate are 534 dense with indistinct particles, i.e consisting of coherent domains having rather the same crystalline 535 orientation, but structural stacking faults and with some spherical or cylindrical closed pores, whereas the reduced aggregates are clearly polycrystalline with visible individual particles with large inter-particle 536 537 porosity, which leads to a porous and airy structure. The oxide aggregate porosity is certainly formed during the calcination step under thermal stresses or due to defects relative to the non-uniform cobalt 538 539 impregnation. The reduced aggregate porosity is formed during the reduction step and is supposed to be 540 caused by the crystallographic phase transformation inducing an increase of the crystalline density of the 541 cobalt, leading to voids formation between particles. However, this porosity could not be explained by the 542 sole crystallographic phase transformation as it theoretically induces an aggregate density decrease of 20%, which is inferior to the decrease of 36% measured by ASAXS, or 54% measured by tomography. 543 Thermal stresses could thus be invoked in order to explain the high magnitude of porosity creation. 544

545 This structure change with the porosity increase during the reduction step appears to be very important as 546 it increases the "aggregate dispersion" (from 18% to 25%) and thus increases the number of accessible 547 cobalt active sites. The activity of the FT catalysts depending essentially on this number of accessible 548 metallic cobalt active sites, the optimization of the reduction step conditions in order to optimise the 549 creation of the porosity seems to be an interesting approach to improve the catalyst activity. Furthermore, 550 the FT catalyst selectivity is suspected to be sensitive to the proportion and size of the aggregates [8] that are known to be controlled by the drying and calcination steps, hence to the active sites proximity. Once 551 552 again the aggregate structure and thus the reduction conditions could impact the selectivity. We can 553 suppose for example that varying the gas hourly space velocity, heating ramp, temperature, or the 554 pressure could lead to different aggregate structures.

555

556 4 Conclusion

557 In this study we have proposed a new approach that combines electron microscopy and ASAXS to 558 characterize Fischer-Tropsch cobalt-based catalysts in order to obtain a detailed and exhaustive 559 description of cobalt active phase. This last one is composed of dispersed nanoparticles and aggregates.

Analytical methodology and data treatment have been developed in ASAXS and electron tomography to suit to these materials. Even if their principles are significantly different, ASAXS and electron microscopy give information in term of particle and aggregate size in good agreement. Likewise, both ASAXS and electron tomography provide some insight on the aggregate structure and morphology. 564 These complementary techniques have allowed in particular investigating the effect of the reduction step 565 on the particles and on the aggregates. We have shown that the reduction had a slight effect on the particles, decreasing their mean size due to the crystallographic phase transformation from Co_3O_4 to Co^0 , 566 but no phenomenon such as sintering has been observed. To the contrary, we have underlined that the 567 568 reduction has a significant effect on the aggregates of which morphology, size and structure are certainly defined during the drying and calcination steps. The dense structure with few closed pores of the oxide 569 570 aggregates evolved into a polycrystalline, porous and airy structure. The porosity of the aggregates is 571 doubled which has the effect to increase drastically the dispersion inside the aggregates and thus increase 572 the number of accessible metallic cobalt active sites. Fischer-Tropsch catalysts activity and selectivity are 573 known to be sensitive to the number and the proximity of the active sites hence the aggregate structure and porosity seem to be important parameters. The operating reduction conditions such as the gas hourly 574 space velocity or the temperature ramp are suspected to affect the internal thermal stresses and thus the 575 final aggregate structure and should be specifically investigated in order to try to improve the catalytic 576 properties. 577

Finally, as the reduction step has obviously a strong effect on the morphology, the structure and the 578 579 dispersion of the active phase, it appears clearly that the catalysts have to be characterize in their reduced 580 state in order to explain the catalytic performances. Moreover, it should be even more appropriate to 581 characterize the used catalyst after the catalytic test as the active phase and specially the aggregate structure is potentially affected under Fischer-Tropsch conditions [45]. In this goal, the ASAXS technique 582 583 should be well suitable as it allows to characterize the particles and the aggregates simultaneously, but 584 especially on catalyst embedded in waxes produced by the synthesis and thus without carrying out a 585 specific washing apt to modify the sample.

This study opens prospects for understanding the role of the carrier chemical and textural properties on the cobalt phase dispersion, but also the role of each drying, calcination and reduction unitary step on particle and aggregates size distributions and cobalt phase dispersion, both parameters that control the active sites number and proximity and thus the activity and the selectivity. Hence, confronting particles and aggregates structure properties and the catalytic performances is the next step.

591

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- available: Fig. S1: schematic view of the MCM-41 formation mechanism. Movie S2: Aligned TEM
- tilt series of the SBA-15 particle from Fig. 1 (sample with the lower TEOS to template ratio). Movie

- 660 S3: Aligned TEM tilt series of the SBA-15 particle from Fig. 3 (sample with the higher TEOS to
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