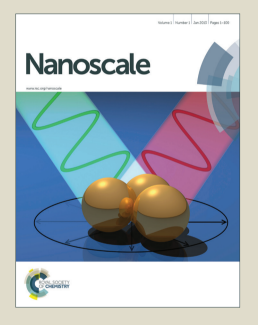
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Uniform 2 nm Gold Nanoparticles Supported on Iron Oxides as Active Catalysts for CO Oxidation Reaction: Structure-Activity Relationship

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Uniform Au nanoparticles (~2 nm) with narrow size-distribution (standard deviation: 0.5–0.6 nm) supported on both hydroxylated (Fe_OH) and dehydrated iron oxide (Fe_O) have been prepared by either deposition-precipitation (DP) or colloidal-deposition (CD). Different structural and textural

- characterizations were applied to the dried, calcined and used gold-iron oxide samples. The transmission electron microscopy (TEM) and high-resolution TEM (HRTEM) described the high homogeneity of supported Au nanoparticles. The ex-situ and in-situ X-ray absorption fine structure (XAFS) characterization monitored the electronic and short-range local structure of active gold species. The synchrotron-based in-situ X-ray diffraction (XRD), together with the corresponding temperature-
- 15 programmed reduction by hydrogen (H₂-TPR), indicated the structural evolution on the iron-oxide supports correlating to their reducibility. An inverse order of catalytic activity between DP (Au/Fe_OH < Au/Fe_O) and CD (Au/Fe_OH > Au/Fe_O) has been observed. Effective gold-support interaction results in high activity for gold nanoparticles locally generated by sintering of dispersed Au atoms on the oxide support in the DP synthesis, while hydroxylated surface favors the reactivity of externally introduced Au
- ²⁰ nanoparticles on Fe_OH support for the CD approach. This work reveals why differences in the synthetic protocol translate to differences in the catalytic performance of Au/FeO_x catalysts with very similar structural characteristics in CO oxidation.

1. Introduction

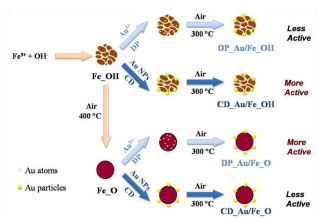
Low-temperature CO oxidation over supported nanosized 25 gold catalysts has been one of the most extensively studied systems in heterogeneous catalysis. 1-3 However, this simple reaction, catalyzed by nanostructured gold, is still very difficult to understand in depth. It is known that various factors, including the size of gold nanoparticles, 1,4,5 the valence state of gold species 30 (metallic Au 0 and ionic Au $^{8+}$), $^{6-8}$ the oxide support, $^{9-14}$ and the metal-support interaction, 15,16 can influence the catalytic performance of supported gold catalysts. However, previous findings are rather controversial with respect to the origin of high activity of gold catalysts, and the debate over the reaction 35 mechanism of low-temperature CO oxidation has continued for more than two decades because of the ultra sensitivity and complexity of the oxide-supported gold system. One of the origins of this debate lies in the interrelationship of the different factors mentioned above, which are very difficult to be 40 distinguished from each other in gold catalysis.

Gold on iron-oxide (Au/FeO $_x$) is one of the highly active catalysts for CO oxidation, and also is a typical system for the

study of the chemistry of gold on oxide supports. 4,5,17 Regarding the widely studied coprecipitated (CP) Au/FeO_r catalysts those 45 have wide size-distribution of gold particles (from sub-nanometer to more than 20 nm), 4 it has been discovered that as-dried samples are much more active for CO oxidation than those after sintering, indicating the negative effect of calcination with respect to activity. 4,17 In fact, the calcination of gold-iron oxide 50 catalysts usually results in simultaneous structural changes of both the gold and the oxide support, i.e., small gold clusters (< 1 nm) with dominant cationic component are transformed into purely metallic nanoparticles (> 1 nm); in addition, the phase of the support changes from hydroxylated ferrihydrite to dehydrated 55 hematite. ¹⁷ For such a catalyst system, the accurate analysis of the origin of CO oxidation activity is a substantial challenge due to the combination between various structural and textural effects. Thus, the separate synthesis of gold nanoparticles with narrow size distribution and iron oxide supports with specific structure is 60 crucial to rationally investigate the structure-activity relationship of Au/FeO_x catalyst.

Therefore, in this paper, we try to identify the support effect and the formation of gold nanostructure of Au/FeO_x catalysts for CO oxidation by using deposition-precipitation (DP)¹⁸ or

colloidal-deposition (CD)13 syntheses for anchoring gold on hydroxylated (Fe_OH) or dehydrated iron oxide (Fe_O) supports (Scheme 1). Highly dispersed gold nanoparticles (~ 2 nm) with narrow size distribution (standard deviation: 0.5-0.6 nm) were 5 obtained. The catalysts were studied by a combination of transmission electron microscopy (TEM) and high-resolution TEM (HRTEM), ex-situ and in-situ X-ray absorption fine structure (XAFS), and in-situ X-ray diffraction (XRD) techniques. These unique samples allow us to distinguish the 10 different factors (particle size, metal-support interaction, surface hydroxyl effect) governing the catalytic activity of gold-iron oxide catalyst. It is also demonstrated how important the formation pathways (locally generated or externally introduced) of gold particles are for the generation of active sites.



Scheme 1 Schematic demonstration of the formation pathways of the different Au/FeOx catalysts.

2. Experimental

15

2.1. Catalyst preparation

20 2.1.1 Preparation of iron oxide supports⁷

In a typical procedure, 0.25 mol L⁻¹ Na₂CO₃ aqueous solution was added drop-wise to 200 mL of 0.1 mol·L⁻¹ Fe(NO₃)₃ aqueous solution under stirring at 80 °C until pH = 8.2, and left stirring for another 1 h. The precipitate was collected by filtration 25 and washed with deionized (DI) water at 80 °C. This precipitate was dried at 120 °C in air for ca. 12 h to generate the hydrated iron oxide support (Fe_OH). The oxide support (Fe_O) was obtained via calcination of Fe_OH in air at 400 °C for 2 h.

2.1.2 Preparation of gold-iron oxide catalysts via colloidal 30 deposition¹³

For colloidal deposition, poly(vinyl alcohol) (PVA, M_w 10,000 from Aldrich, 80% hydrolyzed) was used as the protecting agent. Typically, 0.675 mL of 0.5 wt.% PVA solution (Au:PVA = 1.5:1 in weight) and 2 mL of 0.0125 mol·L⁻¹ HAuCl₄ solution 35 were added into 50 mL of Millipore water (18.25 M Ω) at room temperature under vigorous stirring. After stirring for 10 min, a rapid injection of 1.3 mL of 0.1 mol·L⁻¹ NaBH₄ aqueous solution led to formation of a dark orange-brown solution. 0.5 g Fe_OH or Fe_O was then added to the colloidal gold solution immediately 40 under vigorous stirring, which was continued for 6 h until complete adsorption of the gold (1 wt.%), which was indicated by decoloration of the solution. The solids were collected by

filtration and washing with Millipore water to remove dissolved impurities (Cl-, e. g.). After drying at 60 °C in air overnight, 45 CD_Au/Fe_OH and CD_Au/Fe_O were obtained. All the above steps were carried out in the absence of light by covering all containers with aluminum foil.

2.1.3 Preparation of gold-iron oxide catalysts via deposition precipitation

Typically, 0.5 g support powder (Fe_OH or Fe_O) was suspended in 23 mL Millipore water. 2 mL of 0.0125 mol·L⁻¹ HAuCl₄ aqueous solution was then added to the solution under stirring at ca. 60 °C. After 30 min, 25 mL of aqueous solution containing 0.5 g of urea was quickly added into the stock solution. 55 Thereafter, the solution temperature was increased to 80 °C and kept under vigorous stirring for 3 hours to allow decomposition of urea, which resulted in a gradual increase of the pH value from 4.0 to 8.6. The solution was aged at room temperature for another 20 h. The as-obtained solids were collected by filtration and then 60 washed with Millipore water at 60 °C. After drying at 60 °C in air overnight, DP_Au/Fe_OH and DP_Au/Fe_O were obtained.

2.2 Characterization

The gold loadings of catalysts were determined by inductively coupled plasma atomic emission spectroscopy (ICP-65 AES) on an IRIS Intrepid II XSP instrument (Thermo Electron Corporation).

Transmission electron microscopy (TEM), high-resolution TEM (HRTEM) and the related high angle annular dark fieldscanning TEM (HAADF-STEM) were conducted on a Philips 70 Tecnai F20 instrument at 200 kV and a field emission TEM (JEOL 2100F) machine equipped with a 2k × 2k CCD camera at 200 kV. All the tested samples were ground in a mortar alone rather than suspended in ethanol before deposition on an ultrathin carbon film-coated copper grid.

Temperature-programmed reduction by hydrogen (H₂-TPR) was carried out in a Builder PCSA-1000 instrument (Beijing, China) equipped with a thermal conductivity detector (TCD) to detect H₂ consumption. The sieved catalysts (20–40 mesh, 30 mg) were heated (5 °C·min⁻¹) from room temperature to 400 °C in a 80 20% H₂/Ar (30 mL·min⁻¹) gas mixture. Before measurement, the fresh samples were pretreated in pure O₂ at 300 °C for 30 min.

X-ray absorption fine structure (XAFS): Au L-III absorption edge ($E_0 = 11919$ eV) XAFS spectra were collected ex-situ at BL14W1 beamline of the Shanghai Synchrotron Radiation 85 Facility (SSRF) operated at 3.5 GeV under "top-up" mode with a current of 220 mA. The XAFS data were collected in fluorescence mode with a 32 element Ge Solid State Detector (SSD). The in-situ experiments were conducted at the X18B beamline of the National Synchrotron Light Source (NSLS) at 90 Brookhaven National Laboratory (BNL), operated at 2.8 GeV under "decay" mode with currents of 160-300 mA. The powder sample (~ 25 mg) was loaded into a Kapton tube (O.D. = 0.125 inch) which was attached to an in-situ flow cell. Two small resistance heating wires were installed above and below the tube, 95 and the temperature was monitored with a 0.5 mm chromelalumel thermocouple that was placed inside the tube near the sample. The in-situ CO oxidation reaction (1%CO/16%O₂/83%He, 20 mL·min⁻¹) was carried out under a "steady-state" mode at room temperature (~ 25 °C). Each XAFS 100 spectrum (ca. 15 min collection) was taken under fluorescence

mode with a 4 element Vortex Silicon Drift Detector (SDD). The energy was calibrated for each scan with the first inflection point of the Au L-III-edge in Au metal foil. The X-ray absorption near edge spectroscopy (XANES) and extended X-ray absorption fine 5 structure (EXAFS) data have been analyzed using the Athena and Artemis programs.

X-ray diffraction (XRD): The ex-situ and in-situ XRD experiments were carried out on X7B beamline ($\lambda = 0.3196 \text{ Å}$) of NSLS at BNL. The powder sample (~ 2 mg) was loaded into a 10 quartz tube (I.D. = 0.9 mm, O.D. = 1.0 mm) which was attached to the same flow cell as in the XAFS measurements. One small resistance heating wire was installed right below the tube, and the temperature was monitored with a 0.5 mm chromel-alumel thermocouple that was placed inside the tube near the sample. 15 The in-situ H₂-TPR tests (5%H₂/Ar, 5 mL·min-1) were carried out using a temperature ramp between 25 and 400 °C (5 °C·min⁻¹), and then kept at 400 °C for 20 min. The fresh catalysts were pretreated in 5%O2/He at 300 °C for 30 min before in-situ measurements. Two-dimensional XRD patterns were collected 20 with an image-plate detector (Perkin-Elmer), and the powder rings were integrated using the FIT2D code.

2.3 Catalytic test

CO oxidation activities of gold-iron oxide catalysts were measured in a plug flow reactor by use of 50 mg of sieved (20-40 25 mesh) catalyst in a gas mixture of 1 vol% CO, 20 vol% O₂, and 79 vol% N₂ (from AIR LIQUIDE, 99.997% purity), at a flow rate of 67 mL·min⁻¹, corresponding to a space velocity of 80,000 mL·h⁻¹·g_{cat} -1. Prior to the measurement, the catalysts were pretreated in air at 300 °C for 30 min for activation. After that, $_{30}$ the reactor was cooled down to -50 °C under a flow of pure N_2 gas. The catalytic tests were carried out in the reactant atmosphere by ramping the catalyst temperature (5 °C·min⁻¹) from -50 to 300 °C. The outlet gas compositions of CO and CO₂ were online monitored by a non-dispersive IR spectroscopy 35 (ABB EL 3020). (CO conversion = CO $_{reaction}$ / CO $_{input}$ = CO $_{2}$ $_{output}$ /($CO_{2\ output}$ + $CO\ _{output}$). A typical "steady-state" experiment (30 °C) was conducted in the same gas-mixture at 30 °C for more than 10 h.

3. Results and Discussion

The ICP-AES results (Table 1) show that the experimental Au loading of all four catalysts are close to the target value of 1 wt.%. The XRD data (Figure 1) reveal that the Fe_OH supports in the catalysts are semi-amorphous (see pattern a/b), probably a mixture of several hydroxylated iron oxide phases, even after air-45 calcination at 300 °C. On the other hand, Fe_O obtained by thermal dehydration of Fe_OH, is present as well-defined hematite (α -Fe₂O₃, JCPDS card#: 2-919) in both DP (pattern c) and CD (pattern d) samples, indicating the complete dehydration and crystallization during 400 °C air-calcination. The TEM 50 images (Figures 2 and 3) allow determination of the sizes of Fe_OH and Fe_O supports to ~5 and 10-30 nm, respectively. For the dried DP samples, no gold clusters/particles were found by HRTEM (Figures 2a and 2b). This suggests that the gold atoms are well dispersed on the surfaces of iron-based supports before 55 air-calcination. On the other hand, for the dried CD catalysts, ~ 2 nm Au nanoparticles on Fe_OH and Fe_O surfaces were clearly

observed in TEM (Figures 3a and 3b) due to features of the colloidal deposition preparation, i. e. well-crystallized small-size gold particles already formed before adding the iron-based 60 supports.

Table 1 Characterization of gold catalyst.

Sample	e Au (wt.%)	^a Phase ^b	$D (\text{nm})^c$	$S\left(\mathbf{m}^2\cdot\mathbf{g}^{-1}\right)^d$
DP_Au/Fe_	OH 0.92	Amorphous	s 2.2±0.6	210
DP_Au/Fe	e_O 0.89	α -Fe ₂ O ₃	2.0±0.5	69
CD_Au/Fe_	_OH 0.98	Amorphous	s 2.0±0.6	210
CD_Au/Fe	e_O 0.96	α -Fe ₂ O ₃	2.1±0.6	66

^a Determined by ICP-AES;

^d BET surface areas for calcined catalysts;

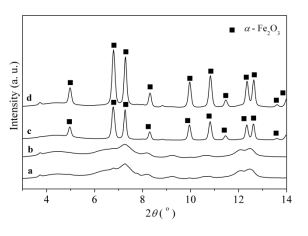


Fig. 1 XRD patterns of calcined Au/FeO_x samples: (a) DP Au/Fe OH; (b) CD_Au/Fe_OH; (c) DP_Au/Fe_O; (d) CD_Au/Fe_O.

In addition to the characterization by electron microscopy, we also carried out XAFS analysis to determine the coordination state of the gold species (Table 2 and Figure 4). The X-ray absorption near edge structure (XANES) spectra reveal the ionic nature of Au species 19 in the dried DP catalysts (Figures 4a and 75 4b), and the related extended XAFS (EXAFS) spectra (Figures 4c and 4d) show the presence of a pure Au-O shell in the dried DP_Au/Fe_OH and DP_Au/Fe_O samples, with distances of the first coordination sphere of ca. 2.0 Å and coordination numbers (CN) of 2.5-2.9 by EXAFS fittings (Table 2), which are 80 consistent with a previous report on fully oxidized Au $^{\delta+}$ species in gold-ceria catalysts. 20 For the CD samples, according to the corresponding XANES (Figure 5a) and EXAFS (Figure 5b) results, gold species were in metallic form with the CN of 10-11 (Table 2), well consistent with the Au⁰ nanoparticle nature.

Upon calcination, the well-dispersed atomic gold species in DP samples were transformed to uniform small (ca. 2 nm) nanoparticles, as seen in the HAADF-STEM images in Figures 2c

^b Identified from XRD patterns for calcined catalysts;

^c Mean Au particle sizes of calcined catalysts, which were calculated from

^{65 &}gt; 100 nanoparticles in HRTEM images;

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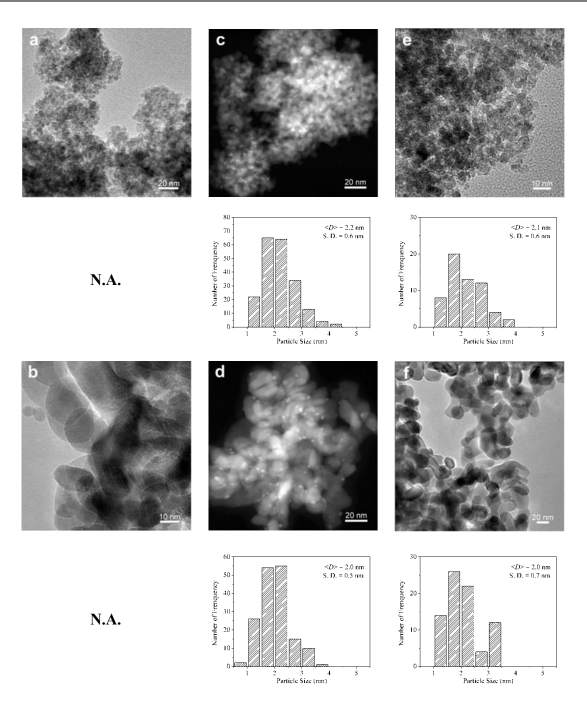


Fig. 2 TEM/HRTEM and HAADF-STEM images of (a) dried DP_Au/Fe_OH; (b) dried DP_Au/Fe_O; (c) calcined DP_Au/Fe_OH; (d) calcined DP_Au/Fe_O; (e) used DP_Au/Fe_OH; (f) used DP_Au/Fe_O.

and 2d (also refer to Table 1). The XANES spectra display distinct metallic gold (Au⁰) features for calcined DP_Au/Fe_OH 5 and DP_Au/Fe_O catalysts (Figures 4a and 4b), and the

corresponding EXAFS fitting results confirm this, since only Au-Au contributions (2.84 Å, CN=10, refer to Table 2, Figures 4c and 4d) are present, instead of the Au-O shell in the dried

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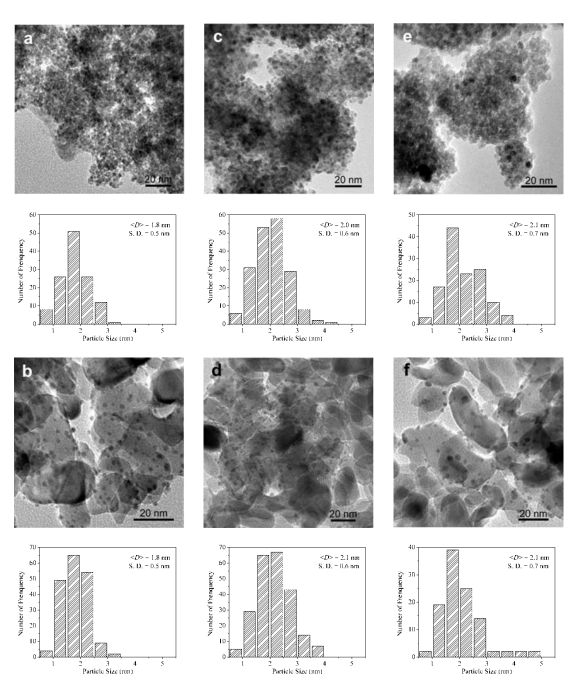


Fig. 3 TEM/HRTEM images of (a) dried CD_Au/Fe_OH; (b) dried CD_Au/Fe_O; (c) calcined CD_Au/Fe_OH; (d) calcined CD_Au/Fe_O; (e) used CD_Au/Fe_OH; (f) used CD_Au/Fe_O.

samples. The formation of gold nanoparticles after 300 °C aircalcination demonstrates the relatively weak interaction between 5 Au atoms and the Fe_OH or Fe_O support in DP samples than the widely studied coprecipitated Au/FeO_x catalysts, i. e. almost no change in the gold nanostructure during oxidative calcination in air at 400 °C or less.21

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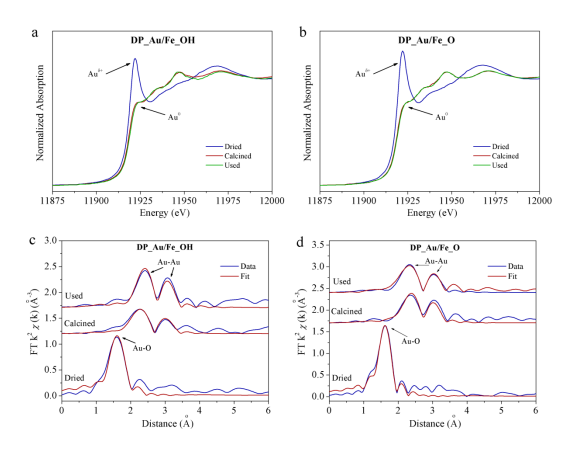


Fig. 4 XANES profiles (a,b) and EXAFS R space fittings (c,d) of Au/FeO_x samples: (a,c) DP_Au/Fe_OH; (b,d) DP_Au/Fe_O.

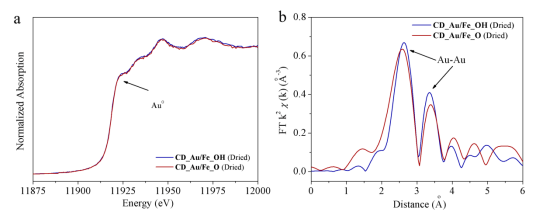


Fig. 5 XANES profiles (a) and EXAFS R space spectra (b) of Au/FeO_x samples (dried CD_Au/Fe_OH and dried CD_Au/Fe_O).

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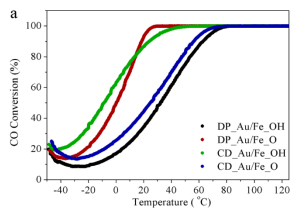
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Table 2 EXAFS fitting results (R: distance; CN: coordination number) of gold catalysts.

G 1	Au-O		Au-Au	
Sample	R (Å)	CN	R (Å)	CN
Au foil	_	_	2.86±0.00	12
DP_Au/Fe_OH (dried)	2.00 ± 0.02	2.5 ± 0.4	_	_
DP_Au/Fe_O (dried)	2.00 ± 0.00	2.9 ± 0.2	_	_
CD_Au/Fe_OH (dried)	_	_	2.84 ± 0.01	11.1±1.4
CD_Au/Fe_O (dried)	_	_	2.83 ± 0.01	10.4 ± 2.2
DP_Au/Fe_OH (calcined)	_	_	2.84 ± 0.02	9.9 ± 2.1
DP_Au/Fe_O (calcined)	_	_	2.84 ± 0.01	9.8±1.5
DP_Au/Fe_OH (used)	_	_	2.87 ± 0.01	10.1±1.6
DP_Au/Fe_O (used)	_	_	2.84 ± 0.01	10.5±1.5

The catalytic performance of the Au/FeO_x catalysts was 5 evaluated for low-temperature CO oxidation. The transient profiles in Figure 6a reveal the higher activity of Au on Fe_O, compared to Au on Fe_OH (T_{90} = 18 °C vs 63 °C) for the DP samples. In contrast, for CD synthesis, the_Au/Fe_OH was superior to CD_Au/Fe_O ($T_{90} = 24$ °C vs 56 °C). At constant 10 temperature of 30 °C (Figure 6b), the final CO conversions after 10 h on stream were around 38%, 70%, 80% and 40% for sample DP_Au/Fe_O, CD_Au/Fe_OH DP_Au/Fe_OH, CD_Au/Fe_O, respectively. This deactivation can be ascribed as the carbonate accumulation. The reactivity of different gold 15 catalysts thus follows the sequence: DP_Au/Fe_O ≈ CD Au/Fe OH > DP Au/Fe OH ≈ CD Au/Fe O. The activities of the DP_Au/Fe_O and CD_Au/Fe_OH catalysts described here are comparable to those of commonly coprecipitated gold-iron oxide catalysts with even higher gold loading of ~ 5 wt.%.

Gold sizes in the used DP catalysts were investigated by TEM/HRTEM. Figure 2 shows that the particles sizes did not change and were around 2 nm before and after the CO oxidation reaction. The related XANES (Figures 4a and 4b) and EXAFS (Figures 4c and 4d) spectra give additional evidence on the 25 metallic nature of the nanoparticles. Similar observations with respect to particles size hold for the CD catalysts: the TEM/HRTEM images of the CD catalysts confirm that gold particle sizes on both Fe_OH and Fe_O supports are constant at ca. 2 nm for dried, calcined and used samples (Figure 3). It 30 reveals that in CD catalysts the interaction between gold and the iron oxide support does not have a significant influence. The major structure is isolated Au nanoparticles randomly dispersed on Fe_OH or Fe_O surface, which can be confirmed by the related XAFS results on dried samples (Table 2 and Figure 5) and 35 very similar to the gold-silica system. 22



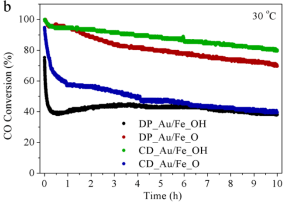
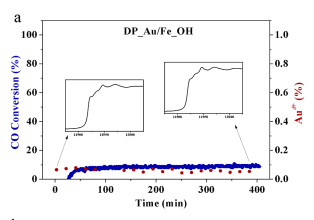


Fig. 6 CO conversions of Au/FeO_x samples measured under (a) transient conditions and (b) at a constant temperature of 30 °C $(1\%CO/20\%O_2/79\%N_2, 80,000 \text{ mL}\cdot\text{h}^{-1}\cdot\text{g}_{\text{cat}}^{-1}).$

Therefore, CD_Au/Fe_OH and CD_Au/Fe_O can be regarded as reference catalysts for the analysis of the support effect, i. e. ~ 2 nm Au on hydroxylated (Fe_OH) and dehydrated (Fe_O). Very interestingly, an inverse order of catalytic activity can be clearly seen from the related experimental data between 45 DP (Au/Fe_OH < Au/Fe_O) and CD (Au/Fe_OH > Au/Fe_O). Recently, it has been reported that the surface hydroxyl groups of the support can enhance the activity of supported gold catalyst for CO oxidation. 23-27 For CD-prepared catalysts, it seems that surface hydroxyl effect plays the dominant role in determining 50 the activity, so that hydroxylated Au/Fe_OH is much more active than Au/Fe_O. However, it has an inverse effect for DP-prepared catalyst. This suggests that other factors should be more significant than surface hydroxyl effect in controlling the reactivity of gold-iron oxide catalysts prepared by the DP method. 55 Since the development of the gold particles follows a much more complex pathway in DP, this is not highly surprising.



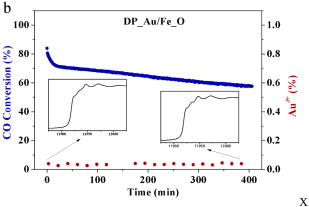


Fig. 7 CO conversions (blue) and Au oxidation states (red) of (a) DP Au/Fe OH and (b) DP Au/Fe O as function of reaction time (1%CO/16%O₂/83%He, 20 mL·min⁻¹, 25 mg, at 25 °C). Inserts are corresponding XANES spectra taken at the start and end points.

First, we focused on the investigation of structural changes of the gold species. Figure 7 exhibits the in-situ XANES results on both DP_Au/Fe_OH and DP_Au/Fe_O. The corresponding fraction of oxidized/reduced gold was calculated by the linear 10 combination method, ²⁸ and the CO conversion was monitored by mass spectroscopy simultaneously. Although the starting Au nanostructures were nearly the same (~ 2 nm metallic particles), the CO oxidation activity of gold on the oxide support was much higher than that on the hydroxylated support (CO conversion at 15 the end of measurement: 58% vs. 10%), in good agreement with Figure 6b. However, the oxidation states of Au were dominantly metallic (Au^0) and the ionic ($Au^{\delta+}$) contributions were below 10% in both samples. Besides, the EXAFS fitting results (Table 2 and Figure 4) further prove the predominance of reduced gold 20 nanoparticles in the calcined (before reaction) and used (after reaction) DP catalysts. This demonstrates that the oxidation state is not the dominant factor governing the activity of gold-iron oxide catalysts obtained via the DP route.

Table 3 Hydrogen consumption (H_2 -consump.) of gold catalysts

	Reduction peak (°C)	Experimental	Theoretical
Sample		H_2 -consump. $(\mu \text{mol} \cdot \text{g}^{-1})$	H_2 -consump. $(\mu \text{mol} \cdot \text{g}^{-1})^a$
DP_Au/Fe_OH	128	437	1300
DP_Au/Fe_O	67, 232	1444	1510
CD_Au/Fe_OH	188, 340	235	1300
CD_Au/Fe_O	257	1042	1510

^{25 &}lt;sup>a</sup> Calculated according to Fe_OH→Fe₃O₄ or Fe₂O₃→Fe₃O₄.

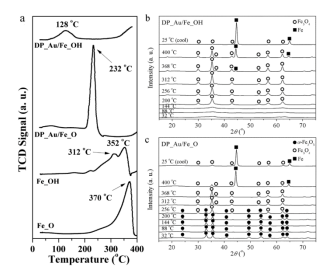


Fig. 8 H₂-TPR profiles (a) and XRD patterns (b,c) under the in-situ conditions of TPR test: (b) DP_Au/Fe_OH; (c) DP_Au/Fe_O.

Another factor which could lead to different catalytic activities of the different DP catalysts could be the transformation of the isolated Au atoms or small clusters (Figures 2a and 2b) into Au particles (Figures 2c and 2d), which takes place during aircalcination. In order to reveal possible differences, hydrogen 35 temperature-programmed reduction (H₂-TPR) was applied to investigate the interaction between metal (Au) and the supports of Fe OH or Fe O (Figure 8 and Table 3). Figure 8a displays the results: for the pure iron-based support, the main reduction peaks are located between 300 and 400 °C, due to the reduction of α - $_{40}$ Fe₂O₃ (hematite) \rightarrow Fe₃O₄ (magnetite). The introduction of gold led to a decrease of this temperature to about 128 and 232 °C, indicating a pronounced Au-O-Fe interaction.²⁷ Figures 8b and 8c show the XRD patterns obtained under in-situ conditions similar to the TPR conditions. For DP_Au/Fe_OH, the starting 45 amorphous hydrated iron oxide was reduced and started to crystallize to Fe₃O₄ at ca. 200 °C, and was further reduced to Fe metal above 368 °C (Figure 8b). For DP_Au/Fe_O, the transformation of the iron oxide from α -Fe₂O₃ \rightarrow Fe₃O₄ occurs at temperatures above 200 °C, the final point of the experiment is 50 metallic Fe and traces of Fe₃O₄ at 400 °C (Figure 8c).

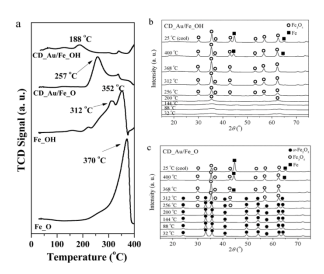


Fig. 9 H₂-TPR profiles (a) and XRD patterns (b,c) under the in-situ conditions of TPR test: (b) CD Au/Fe OH; (c) CD Au/Fe O

For the DP_Au/Fe_O catalyst, there is an intense reduction 5 peak centered at 232 °C. The H₂ consumption related to this reduction peak is 1444 µmol·g⁻¹ which is very close to the theoretical value (1510 µmol·g⁻¹, Table 3), indicating the full reduction of most of the α-Fe₂O₃ to Fe₃O₄. However, for the DP_Au/Fe_OH, a broad reduction peak centered at 128 °C with a $_{10}$ low H_2 consumption value of 437 $\mu mol \cdot g^{-1}$ is observed. The theoretical value of H₂ consumption related to a full transformation of Fe_OH→Fe₃O₄ is 1300 µmol·g⁻¹, which is much higher than that of the reduction peak centered at 128 °C for DP_Au/Fe_OH. Therefore, only part of the Fe_OH was 15 reduced to Fe₃O₄ at a temperature of 200 °C (the end of the broad peak), in agreement with the in situ XRD pattern at 200 °C which covers both Fe₃O₄ and some amorphous phases, as shown in Figure 8b. The TPR results confirm an more effective interaction^{20,27} of gold species with iron oxide than with 20 hydroxylated amorphous iron oxide, which explains why the gold on the dehydrated support (Fe_O) is significantly more active than gold on the hydroxylated support (Fe_OH). It also suggests that the presence of the hydroxylated support prevents the creation of active Au sites during the crystallization of both gold 25 and the iron-oxide support.

On the other hand, for the CD catalysts, the 2 nm Au nanoparticles are already formed before calcination, and the overall interaction between gold and iron-oxide support is less effective than for the DP samples (refer to the lower reduction 30 peak-area for CD_Au/Fe_O in Figure 9a, and the less hydrogen consumption in Table 3). Based on the related in-situ XRD results, the phase transformations of iron-oxide support in CD samples (Figures 9b and 9c) were very similar to those synthesized by DP (Figures 8b and 8c). Here, hydroxylated 35 surface is the dominant factor on tuning the Au dispersion/stabilization, and thus the hydroxylated catalyst (CD_Au/Fe_OH) was more active than the dehydrated sample (CD_Au/Fe_O). However, also the surface hydroxyl groups may have an effect, since this type of support brings the extra activity 40 enhancement for the gold catalyst systems. 23-27

From the results described it can be concluded that the preparation method (DP or CD), respectively, the formation

mechanism of the gold particle formation (locally generated or externally introduced) are crucial in determining the gold 45 reactivity. Deposition-precipitation, or local generation, benefits from the dehydrated support (Fe_O) only, while colloidaldeposition is better suited for the hydroxylated support (Fe_OH). For the DP route, the gold particles and their interaction with the support develop during thermal treatment. Here it is clear that the 50 more strongly interacting gold species lead to more active catalysts. In the case of colloidal deposition, the beneficial effect of the hydroxyl groups on the catalyst surface dominates the performance, and thus the gold on the Fe_OH is the more active system for this synthetic pathway.

55 4. Conclusion

In summary, we have prepared active gold-iron oxide catalysts with uniform size of ~ 2 nm for gold nanoparticles by both deposition-precipitation and colloidal-deposition methods, and further investigated the main factors in governing the 60 catalytic activity of Au. An inverse order of catalytic activity was found between deposition-precipitation (Au/Fe_OH < Au/Fe_O) and colloidal deposition (Au/Fe_OH > Au/Fe_O) system. Higher level of effectiveness of interaction between gold and the oxide support is suggested to account for the superiority of the oxide 65 support (Fe_O) in the deposition-precipitation system, while surface hydroxyls induce the high reactivity of gold on the hydrated iron oxide support (Fe_OH) for the colloidalprecipitation approach. Thus, the formation mechanism of the gold species, which is directly related to the synthesis strategy, is 70 a key factor which contributes to the activity of iron oxide supported gold catalysts, in addition to structural effects.

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- ‡ Footnotes should appear here. These might include comments relevant to but not central to the matter under discussion, limited experimental and 10 spectral data, and crystallographic data.

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