Structure–reactivity relationship in Co3O4 promoted Au/CeO2 catalysts for the CH3OH oxidation reaction revealed by in situ FTIR and operando EXAFS studies

This is the author's manuscript

Original Citation:

Availability:
This version is available http://hdl.handle.net/2318/1633159 since 2017-05-11T19:34:03Z

Published version:
DOI:10.1039/C6TA06442F

Terms of use:
Open Access
Anyone can freely access the full text of works made available as "Open Access". Works made available under a Creative Commons license can be used according to the terms and conditions of said license. Use of all other works requires consent of the right holder (author or publisher) if not exempted from copyright protection by the applicable law.

(Article begins on next page)
The beneficial influence of cobalt oxide on the CO oxidation behaviour of Au/CoOx/CeO2 supported gold nanoparticles attracted significant attention due to their high catalytic activity in various oxidation reactions at low temperatures. The catalytic oxidation of Volatile Organic Compounds (VOCs), such as methanol, is a subject of considerable interest due to their relevance in many industrial applications as well as in the field of sustainable chemistry. These catalytic processes have been considered as some of the most efficient ways to reduce harmful emissions from various chemical industries. During the past two decades gold-based catalysts have been proven to be effective for many reactions of environmental significance.1

Considerable emphasis is placed on the catalyst preparation and on the selection of the support.2 The oxidation activity of gold catalysts is closely related to the size of the Au particles3 and to the ability of the support to provide active oxygen species. For this reason, the selection of the proper support is a critical factor.

Ceria is very attractive, due to its ability to maintain high metal dispersion and to change Ce oxidation state depending on the redox conditions. The rapid change of oxidation state is related to its ability to store and release oxygen, a property measured by the “oxygen storage capacity” (OSC),4 which results in rapid formation and elimination of oxygen vacancy defects.5 These characteristics make CeO2 a very interesting support for oxidation reactions. In principle, the modification of ceria can increase the concentration of oxygen vacancies and a strong influence of the nature of the added oxide (Me= Fe, Mn, Sn) on the catalytic performances was observed recently by some of us.6 For example, it was reported that the modification of CeO2 with different Zr amounts leads to increased catalytic activity in the CO oxidation7, as well as in the Water Gas Shift reaction8 and in the steam reforming of methanol.9 Moreover, gold catalysts supported on iron-modified ceria oxides containing opportune molar percentages of Fe were proved to be more active than unmodified Au/CeO2 for total and preferential CO oxidation10,11 and for Water Gas Shift reaction.12 The beneficial influence of cobalt oxide on the CO oxidation behaviour of Au/CoOx/CeO2-Al2O3 systems was previously shown.13 It was found that the catalytic activity depends on the cobalt loading, which

Abstract
A strong influence of the amount of the Co2O3 promoter on the catalytic performance in methanol oxidation of different gold catalysts supported on ceria was observed. The activity followed the order: Au/10 wt.% Co3O4-doped CeO2> Au/5 wt.% Co3O4-doped CeO2> Au/15 wt.% Co3O4-doped CeO2> Au/CeO2. FTIR measurements of adsorbed CO indicate that oxidized gold sites are initially present on the activated samples and that such species are involved in the methanol reaction. Methanol oxidation performed in static conditions gave rise at 75 °C to mainly formate species on Au/CeO2 and to a large variety of different carbonate species on Au/10 wt.% Co3O4-doped CeO2. FTIR and EXAFS analyses revealed that the active sites present on the best performing Au/CeO2 catalyst added with 10 wt.% Co3O4 are oxidized gold species, close Co sites, at the interface with the support, which are reduced under reaction conditions. These species are able to activate and to react with oxygen giving rise to formate and carbonate species.

Introduction
Supported gold nanoparticles attracted significant attention due to their high catalytic activity in various oxidation reactions at low temperatures. The catalytic oxidation of Volatile Organic Compounds (VOCs), such as methanol, is a subject of considerable interest due to their relevance in many industrial applications as well as in the field of sustainable chemistry. These catalytic processes have been considered as some of the most efficient ways to reduce harmful emissions from various chemical industries. During the past two decades gold-based catalysts have been proven to be effective for many reactions of environmental significance.1

Considerable emphasis is placed on the catalyst preparation and on the selection of the support.2 The oxidation activity of gold catalysts is closely related to the size of the Au particles3 and to the ability of the support to provide active oxygen species. For this reason, the selection of the proper support is a critical factor.

Ceria is very attractive, due to its ability to maintain high metal dispersion and to change Ce oxidation state depending on the redox conditions. The rapid change of oxidation state is related to its ability to store and release oxygen, a property measured by the “oxygen storage capacity” (OSC),4 which results in rapid formation and elimination of oxygen vacancy defects.5 These characteristics make CeO2 a very interesting support for oxidation reactions. In principle, the modification of ceria can increase the concentration of oxygen vacancies and a strong influence of the nature of the added oxide (Me= Fe, Mn, Sn) on the catalytic performances was observed recently by some of us.6 For example, it was reported that the modification of CeO2 with different Zr amounts leads to increased catalytic activity in the CO oxidation7, as well as in the Water Gas Shift reaction8 and in the steam reforming of methanol.9 Moreover, gold catalysts supported on iron-modified ceria oxides containing opportune molar percentages of Fe were proved to be more active than unmodified Au/CeO2 for total and preferential CO oxidation10,11 and for Water Gas Shift reaction.12 The beneficial influence of cobalt oxide on the CO oxidation behaviour of Au/CoOx/CeO2-Al2O3 systems was previously shown.13 It was found that the catalytic activity depends on the cobalt loading, which
provides a submonolayer coverage of the support surface, where a CoO2−CeO2 interaction takes place giving rise to a synergism between Co and Ce redox properties. Indeed, Co3O4 itself was demonstrated to be very active in the sub-ambient CO oxidation reaction depending on the morphology and on the preferential exposition of the [1 1 0] plane in which the highest number of Co(III) sites are present. Three dimensionally ordered macroporous Au/Co3O4−CeO2 catalysts showed enhanced CO preferential oxidation in H2-rich gas mixtures and formaldehyde oxidation. The former paper proposed that the origin of the high CO conversion and CO2 selectivity was the presence of ionic Au in intimate contact with CeO2. In the latter one, a mechanism involving the synergistic effect between CeO2 and Co3O4 supports, which greatly accelerates the surface active oxygen migration and activates the Au species was proposed to explain the improvement in the catalytic HCHO oxidation. It was established that Au/Co3O4−CeO2 catalysts prepared by mechano-chemical activation exhibited much higher activity in the reaction of complete benzene oxidation than that prepared using coprecipitation. The reason of this different catalytic behaviour was the existence of a Co-containing phase with enhanced redox properties that influenced both gold and modified ceria in close vicinity. Therefore, ceria supports with different amount of Co3O4 were prepared by mechano-chemical activation in order to optimize the content of promoter. Recently, we observed a strong influence of the nature of the added oxide on the activity of ceria-based gold catalysts for catalytic abatement of CO in waste gases. In particular, gold catalyst supported on ceria modified by Co3O4 exhibited the best performance: 100% CO conversion degree was achieved at 25 °C. The support with composition 90 wt.% CeO2 and 10 wt.% Co3O4 was beneficial for the nucleation and growth of highly dispersed gold. On this system, a synergy between gold and Co-doped ceria support causes significant enhancement in the reducibility and capability for oxygen activation, which resulted in an improved oxidation activity. However, a clear correlation between the catalytic activity with the nature and structure of the active sites has not been fully established yet.

The correlation existing between catalyst structure and catalytic activity/selectivity is the main goal in research applied to catalysis. In principle, EXAFS/XANES can give relevant complementary information on the structural and electronic properties of such systems. In addition, FTIR spectroscopy represents a powerful technique to investigate the surface sites at an atomic level. In particular, the analysis of FTIR spectra of adsorbed probe molecules allows to understand the nature and the abundance of the exposed active sites as well as to have detailed information on their chemical environment. Therefore, the use of these techniques can assist in the comprehension of the parameters ruling the unique catalytic properties of gold catalysts, as well as to implement the knowledge in the design of new systems. Keeping in mind these purposes, the aim of this work was an accurate EXAFS and FTIR study on the nature of the active sites exposed at the surface of highly dispersed gold catalysts supported on ceria modified by different amounts of Co3O4. Research efforts were focused to achieve information on the local structure of the active sites in terms of position of cobalt atoms in respect to the gold metallic phase.

Experimental

Materials

Cerium hydroxide was synthesized by precipitation of aqueous solutions of Ce(NO3)3·6H2O with K2CO3 at constant pH=9.0 and at a temperature of 60 °C, aging at the same temperature for 1 hour, filtering, washing until removal of NO3 ions and drying in vacuum at 80 °C. Mixed CeO2−Co3O4 supports were prepared by a simple mechano-chemical mixing procedure: a mixture of cerium hydroxide and (5, 10, 15 wt%) of laboratory-made Co3O4 was subjected to mechano-chemical milling for 30 min in a mortar and calcination at 400 °C for 2 hours. Hereafter the supports will be labelled as Ce5Co, Ce10Co and Ce15Co depending on cobalt oxide loading in wt%. Gold catalysts (3 ± 0.05 wt.%) were prepared by the deposition–precipitation method at constant pH 7 and temperature 60 °C. Gold was deposited on a mixed metal oxide support preliminary suspended in water and activated in an ultrasound disintegrator under vigorous stirring. Hereafter catalysts will be labelled as AuCe5Co, AuCe10Co and AuCe15Co depending on cobalt oxide loading in wt%.

The deposition-precipitation was carried out in an automated laboratory reactor under full control of all preparation parameters (pH, temperature, stirring speed, reactant feed flow rates, etc.). After aging at 60 °C within 1 h, filtering and careful washing, the precursors were dried under vacuum at 80 °C and calcined in air at 400 °C for 2 hours. Moreover, unmodified Au/Co3O2 and Au/Co3O4, (hereafter labeled as AuCe and
Au/Co₃O₄, respectively), used as blank samples, were also prepared by the deposition-precipitation method under the conditions described above. All initial salts used were ‘analytical grade’.

**Catalytic activity measurements**

The catalytic activity of the samples in CH₃OH oxidation was measured using an isothermal plug flow reactor within a different temperature range. The following conditions were chosen: catalyst bed volume – 0.5 g (particle size 0.6 – 1.0 mm), inlet CH₃OH concentrations – 2.0 % balanced with air and space velocity 10000 h⁻¹. The tests were carried out by increasing the reaction temperature step-wise until complete CH₃OH oxidation. The degree of conversion was measured at every 20 °C step after stationary conversion was achieved. The behaviour was also studied during decrease of the temperature down to room temperature, again step-wise. The conversion data at increasing and decreasing temperature match well. The duration of catalytic test for each sample was about 8 hours. The reactant and product gases were analysed for methanol, CO, CO₂, O₂ and N₂ by HP 5890 Series II gas-chromatograph, equipped with flame ionization and thermal conductivity detectors and Porapak Q and MS-5A columns.

**Methods**

The amount of gold in all samples was analyzed by atomic absorption spectroscopy (Varian Vista MPX) and the values determined were 3 ± 0.05 wt % Au.

High resolution transmission microscopy (HRTEM) measurements were performed using a side entry Jeol JEM 3010 (300 kV) microscope equipped with a LaB₆ filament and fitted with X-ray EDS analysis by a Link ISIS 200 detector. For analyses, the powdered samples were deposited on a copper grid, coated with a porous carbon film. All digital micrographs were acquired by an Ultrascan 1000 camera and the images were processed by Gatan digital micrograph. A statistically representative number of Au particles (i.e. 250 for each sample) was counted in order to obtain the particle size distributions.

The FTIR spectra were recorded in transmission mode at a resolution of 2.0 cm⁻¹ on a Perkin-Elmer 2000 spectrometer (equipped with a MCT detector) with the samples in self-supporting pellets introduced in an AABSPEC 2000 cell allowing to run the spectra in situ in controlled atmosphere and temperature (from room temperature up to 600 °C).

As for the experiments at low temperature, the pelletised samples were introduced in a cell allowing thermal treatments in controlled atmosphere and spectrum scanning at controlled temperature (from -180 °C up to room temperature). In order to mimic the activation before the catalytic tests and to follow the effect of such activation on the nature of the exposed active sites and of the species formed during methanol oxidation reaction, the catalysts were subjected to an oxidative treatment before the measurements. The thermal treatment included heating from room temperature (r.t.) up to 200 °C under outgassing below 10⁻³ mbar; an inlet of 20 mbar of oxygen and heating up to 400 °C; at 400 °C the oxygen was changed three times (20 mbar for 10 min each run). After that, the sample was cooled down to r.t. in oxygen and finally outgassed at the same temperature. For each spectrum, the spectrum of the sample before the inlet of the probe/reactants (CO, methanol or oxygen) or of the methanol/oxygen mixture was used as background. All reported spectra were background subtracted and normalized to the density of the pellets.

EXAFS experiments were carried out on the BM23 beamline at ESRF facility (Grenoble, France). ¹¹¹ Fluorescence XAS spectra at the Au L₃-edge (11.919 keV) were collected with a 13-element Ge detector. The beam energy were selected by double-crystal Si(111) monochromator, and the third harmonic rejection was performed by Rh coated mirror with an angle of -2.2 mrad.²¹ EXAFS signal were extracted and analyzed by IFEFFIT package.³² The samples were hosted inside a plug flow reactor using 1 mm quartz capillary available on BM23 for in situ/operando measurements.³³ The experiments can be divided into two stages. Stage 1: the samples have been heated from room temperature up to 200 °C by a small furnace, then they have been treated with a mixture O₂/He (20 % O₂-He, 30 cm³/min) up to 400 °C. After oxygen treatment, a CO-O₂- 1:1/He mixture (5% CO, 5 % O₂) was fluxed at r.t.. Stage 2: the samples were heated at 200 °C and treated with a mixture O₂/He (20 % O₂-He, 30 cm³/min) up to 400 °C. Successively the sample was cooled down to r.t. and a vapour of methanol/O₂/He mixture (20 % O₂-He) was fluxed and finally the catalysts were heated up to 100 °C in the reaction mixture.

**Results and discussion**

Catalytic activity in the CH₃OH oxidation
The temperature dependence of CH$_3$OH conversion degree is presented in Figure 1. The catalytic measurements indicated that the amount of promoter has a strong influence on the activity of ceria-based gold catalysts, particularly in defining the onset temperature of the catalytic activity.

In particular, the activity towards the CH$_3$OH oxidation follows the trend: AuCe10Co > AuCe5Co > AuCe15Co > AuCo$_3$O$_4$ > Ce10Co. The light-off curve related to bare ceria doped by 10 wt% Co$_3$O$_4$ appeared at the highest temperature, giving evidence of the key role displayed by gold.

The best performance was observed in the case of the catalyst doped by 10 wt% Co$_3$O$_4$, over which almost 100% CH$_3$OH conversion degree at a temperature as low as 40 °C was obtained.

**Figure 1.** Temperature dependence of CH$_3$OH conversion over gold catalysts on ceria doped by 5, 10 and 15 wt% Co$_3$O$_4$, over unpromoted Au/CeO$_2$, Au/Co$_3$O$_4$ and bare ceria doped by 10 wt% Co$_3$O$_4$.

The comparison with the data recently reported by Scirè et al. (about 95 % methanol conversion at 125 °C over a gold supported on ceria catalyst prepared by deposition-precipitation) puts in evidence the promising application of AuCeO$_2$ catalysts doped by Co$_3$O$_4$. Moreover, a high defectivity of ceria exposed faces, along with the presence of gold nanoparticles with the smallest size and the highest hydrogen consumption during TPR experiments (e.g. highest oxygen mobility) were found over this AuCe10Co sample, according to our previous high resolution TEM characterization.

The light-off curves for CH$_3$OH oxidation of Au/CeO$_2$ catalysts doped by Co$_3$O$_4$ were shifted to lower temperature with respect to that of undoped Au/CeO$_2$ and to much lower temperature if compared to Au/Co$_3$O$_4$, that is also the worst catalyst. This behaviour is better pointed out from the data in Table 1, where the T$_{50}$ conversion values (i.e. the temperature at which 50 % conversion was reached) are contrasted. From the data reported in Table 1, the promoting effect of Co$_3$O$_4$ is evident, together with the fact that the beneficial addition of cobalt oxide increases at low promoter content, reaches a maximum and then declines, in agreement with the fact that gold supported on the pure promoter phase (Au/Co$_3$O$_4$ sample) is the less active catalyst. In addition, it was found that the kinetic regime is not controlled by diffusional limitations.

This statement is based on the experiments performed with two different catalyst pellets, the former by using a pellet with a given weight and the latter with a pellet with a double weight with respect to that employed in the first test.

**Table 1.** Extrapolated T$_{50}$ values for CH$_3$OH oxidation and calculated TOFs (s$^{-1}$) at two different temperatures over the catalysts.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>CH$_3$OH oxidation (°C)</th>
<th>TOF at 20 °C (s$^{-1}$)</th>
<th>TOF at 40 °C (s$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AuCeO$_2$</td>
<td>53</td>
<td>0.00446</td>
<td>0.0167</td>
</tr>
<tr>
<td>AuCe5Co</td>
<td>41</td>
<td>0.00633</td>
<td>0.0253</td>
</tr>
<tr>
<td>AuCe10Co</td>
<td>&lt; 20</td>
<td>0.0484</td>
<td>0.0505</td>
</tr>
<tr>
<td>AuCe15Co</td>
<td>47</td>
<td>0.00458</td>
<td>0.0172</td>
</tr>
</tbody>
</table>
Gold dispersions were calculated for all samples based on the average particle size determined by HRTEM/HAADF and reported in Ref. 19 (AuCe5Co – 2.4 nm, AuCe10Co – 1.8 nm, AuCe15Co – 2.1 nm), assuming the presence of cuboctahedral particles. According to the catalytic trend for the CH3OH conversion (see also Figure 1), the TOF values obtained for the AuCe10Co catalyst are one order of magnitude higher than those related to the other samples. The temperature has a pronounced effect on the conversion: indeed, with the exception of AuCe10Co, the obtained TOF values increase of about four times when the temperature is raised from 20 °C up to 40 °C. A very recent study on Au supported on bare ceria reported the TOF values at two different temperatures.36 The TOF markedly increased from 3.72 x 10^-5 s^-1 at 45 °C to 1.02 x 10^-4 s^-1 at 70 °C, putting in evidence an enhancement even more accentuate, but in agreement with the data reported in Table 1. Moreover, the behaviour of AuCe5Co and AuCe15Co is very similar to the unpromoted AuCe catalyst. On the contrary, in the case of AuCe10Co, the TOF value is almost unchanged, further showing that 10 wt% Co3O4 provides the optimal composition for the catalysis and that for this catalyst the composition plays the major role, even with respect to the temperature.

Effect of the support composition on the gold dispersion

The catalytic activity measurements revealed almost 100% CH3OH conversion degree at 40 °C for the AuCe10Co catalyst. Therefore, the addition of 10 wt% Co3O4 has improved the activity. It was previously found that the most active catalyst was that containing a big amount of Au nanoparticles with average size of 1.8 nm. 19 Since the deposition-precipitation of gold was carried out on the already doped support, an evaluation of the influence of the Co3O4 presence on the metal dispersion and size distribution was done by performing new HRTEM measurements on the most promising AuCe10Co catalyst. The results were compared with those obtained for undoped AuCeO2 in order to put in evidence peculiar features arisen from the different composition of the support on which the same amount of gold has been deposited. All these data are summarized in Figure 2. Both catalysts are supported on cubic CeO2, mainly exposing the (111) face (not shown).

**Figure 2.** HRTEM images collected on AuCeO2 (section a) and AuCe10Co (section b), where the presence of Au nanoparticles is put in evidence by red circles. Au particle size distributions of AuCeO2 and AuCe10Co are reported in section c and d, where n.p.= number of particles. Instrumental magnification: 300000X and 500000X, respectively.

The AuCe catalyst contains gold species with heterogeneous size in the 2-12 nm range (section a, evidenced by red circles), resulting in an average diameter equal to 7.5 ± 4.5 nm (section c). However, the particle size distribution reveals that the large majority of the roundish gold nanoparticles has size around 2 nm (more than 50 %).

On the contrary, the AuCe10Co catalyst exposes gold nanoparticles with smaller size (section b, evidenced by red circles), as indicated by the narrow particle size distribution, being the average size equal to 1.69 ± 0.41 nm (section d). It can be inferred that the presence of 10 wt% Co3O4 in the ceria support significantly affected the metal dispersion, exploiting a benefic effect in the stabilization of the Au nanoparticles during the final calcination at 400 °C and therefore favoring the formation of particles with more uniform size than in the case of AuCeO2. These findings are in agreement with the enhancement in the catalytic activity observed for the catalyst doped with 10 wt% Co3O4.
Nature of the exposed sites and in situ spectra of methanol oxidation reaction

FTIR spectroscopy measurements were carried out on the bare AuCe and on AuCe10Co, i.e. on the most active doped sample. This vibrational study is aimed to investigate the effect of the promoting species on the nature and the abundance of gold exposed sites and to have information on the nature of the surface species formed during methanol oxidation from r.t. up to 100 °C.

In Figure 3a, the FTIR spectra collected upon CO adsorption at r.t. on unpromoted AuCe oxidised at 400 °C are shown.

Besides a shoulder at 2112 cm\(^{-1}\), related to CO on partially oxidized metallic gold,\(^{37}\) a broad absorption at 2138 cm\(^{-1}\) is observed after the inlet of 15 mbar CO at r.t. on AuCe (blue line). Upon CO pressure decreases at the same temperature (black lines) the band at 2138 cm\(^{-1}\) decreases in intensity without changing position, whilst the shoulder at lower wavenumber is totally depleted. The residual absorption is irreversible to the outgassing at r.t. (down to 1.0 \(\times\) 10\(^{-3}\) mbar, red line), giving an anomalous evidence of a strong bond between CO and the involved adsorption sites, since CO is usually reversibly adsorbed on metallic gold. Some of us reported on a similar band some years ago and assigned it to CO on positively charged Au\(_n^+\) clusters, where 4\(\leq n\leq 6\) stabilized on the oxidized surface of ceria.\(^{38}\) This assignment was supported by DFT calculations of Wu et al.\(^{39}\), showing that the adsorption energy of CO on small cationic, neutral, and anionic clusters progressively increases with the electron depletion from the cluster. Also the calculated C-O stretching frequencies supported this assignment as they increased with increasing the positive charge on the clusters. In addition, also the paper by Fielicke et al.\(^{40}\) in which the CO vibrational frequencies of Au\(_n^+\)(CO)\(_m^+\) complexes in the gas phase (3 \(\leq n\leq 10\) and 3 \(\leq m\leq 8\)) have been measured experimentally, corroborated our assignment together with further spectroscopic findings. Cationic gold clusters are still present after methanol oxidation reaction, as signalled by the broad band at 2121 cm\(^{-1}\), whilst no bands related to oxidised gold species exposed at the surface of the nanoparticles have been detected (Figure 3b). The inlet of the CO probe at r.t. on the AuCe10Co catalyst (Figure 3c) produced a band at 2149 cm\(^{-1}\) (red line) that despite the decrease of the CO pressure, firstly increases in intensity and red shifts at 2140 cm\(^{-1}\) (orange line) and then decreases in intensity and moves to 2135 cm\(^{-1}\) (black lines down to violet line). Also in this case the residual absorption is irreversible to the outgassing at r.t. (violet line). These results indicate the absence of partially
oxidized metallic gold (band at 2112 cm\(^{-1}\))\(^{30}\), showing that only cationic gold clusters with the same nuclearity as those detected on AuCe are exposed at the surface of the oxidised AuCe10Co sample before reaction. Moreover, if the spectra are compared with those reported in section a, the amount of these species is higher on the Co-containing sample. The increased intensity of this absorption despite the reduction of the CO pressure is an indication that the small cationic gold clusters are initially covered by adsorbed oxygen and probably are in a higher oxidized state. Indeed, the inlet of the probe at r.t. produced some CO\(_2\) by reaction of CO and oxygen adsorbed on gold clusters and big amounts of mono-, bi- and polydentate carbonate species are observed (see Figure S1). On the contrary, only oxidised gold species exposed at the surface of the nanoparticles are observed after methanol oxidation reaction, as indicated by the presence of a band centred at 2113 cm\(^{-1}\) (Figure 3d). It is worth noting that no CO is adsorbed at r.t. on the Ce10Co sample, indicating that neither cerium nor cobalt cations are able to adsorb the probe in our experimental conditions. However, previous experiments carried out at low temperature demonstrated the presence of Ce\(^{4+}\) at the surface of both samples. Interestingly, the presence of bands related to Co\(^{3+}\) cations was hardly hypothesized on AuCe10Co.\(^{19}\)

The AuCe and AuCe10Co catalysts were contacted with methanol (5 mbar) and then oxygen (5 mbar) at r.t. and then the temperature was gradually increased up to 100°C in order to follow the formation (and the possible evolution) of surface species during methanol oxidation by \textit{in situ} FTIR spectroscopy. Notwithstanding that \textit{in situ} conditions, under static equilibrium pressure of the reactants are far away from the ideal \textit{operando} conditions where reactants are sent in a plug-flow reactor, FTIR spectroscopy can provide insights on both spectators and intermediate species formed on the different catalysts. Firstly, the inlet of methanol at r.t. on AuCe and AuCe10Co (blue lines in Figure S2) produced different amounts of on top, doubly (two species) and triply bridged methoxy species, as summarised in Table 2, where the vibrations related to the methoxy species adsorbed on AuCe and on AuCe10Co catalysts are reported. The evolution of the methoxy species at increasing temperature is shown in Figure S2 (from brown to red lines). On top and different doubly bridged methoxy species have been detected on both samples\(^{41, 42}\), starting from room temperature (blue line) up to 50 °C (brown lines), then a marked increase in intensity of the bands related to on top methoxy species is observed at 75 °C (violet lines). Methoxy species are definitely more abundant on AuCe10Co than on AuCe, in particular the intensity of the absorptions due to the doubly bridged methoxy species with different coordination of Ce cations with oxygen ions (bands at 1053 and 1038 cm\(^{-1}\)) seems enhanced on AuCe10Co, further indicating that the addition of cobalt leads to a lower degree of coordination of the ceria surface sites of the doped sample and as a consequence to an enhanced reactivity towards the same amount of methanol. It has been reported that methoxy species formed upon methanol dissociation over CeO\(_2\) behave both as spectators and intermediate species, depending on their proximity with gold nanoparticles.\(^{36}\) It can be proposed that on AuCe10Co, that is the most active catalyst, the higher is the metal dispersion, the larger is the amount of active methoxy species acting as reaction intermediates.

The above findings are also in agreement with the results of the previous analysis of the HRTEM diffraction fringes related to the ceria-containing phases, indicating an increase of about four times of the spacings due to defective ceria phases for AuCe10Co (37.35%) if compared to AuCe (10.77 %).\(^{19}\)

The presence of more defective ceria may also favour the formation of double bridged reactive methoxy species on the surface of the catalyst. In addition, triply bridged methoxy species are easily distinguishable on AuCe10Co also at 100 °C,\(^{25, 26}\) whilst they are almost depleted at the same temperature on AuCe, as indicated by the weak bands at 1026, 2847 and 2934 cm\(^{-1}\). These features further indicate that, due to the presence of cobalt, AuCe10Co displays a more easy activation of methanol and an enhanced reactivity of methoxy species if compared to AuCe.

The FTIR spectra collected on AuCe and on AuCe10Co in contact with the methanol/oxygen mixture at increasing temperature are reported in Figure 4 section a and b, respectively.

On the AuCe sample, the methanol oxidation was mainly limited to the production of bidentate (bands at 2845, 1572, 1549, 1398, 1371 cm\(^{-1}\)) and monodentate (bands at 1605 and 1250 cm\(^{-1}\)) formate species.\(^{43}\) However, bands related to monodentate (1507 and 1358 cm\(^{-1}\)), bidentate (1564, 1299, 1022, 856 cm\(^{-1}\)) and polydentate (1475, 1053 and 833 cm\(^{-1}\)) carbonate species\(^{27}\) start to increase in intensity at 75 °C, simultaneously with the decrease of the bands related to methoxy species (see Figure 5 and Figure S2, where the methoxy region in the 1150–950 cm\(^{-1}\) region is zoomed).

The FTIR results put in evidence that on both catalysts methanol reacted with the surface giving rise to different methoxy species at r.t. and to formates at 50 °C. These species further evolved to mono-, bi- and
polydentate carbonates by increasing the temperature at 75 °C and up to 100°C. On AuCe10Co larger amounts of carbonates are produced.

**Table 2.** Wavenumbers for vibrations of on top, doubly (two species) and triply bridged methoxy species adsorbed on AuCe and on AuCe10Co catalysts.

<table>
<thead>
<tr>
<th>Vibrational modes</th>
<th>( \nu_2(\text{CH}_2) ) cm(^{-1} )</th>
<th>( 2\delta(\text{CH}_2) ) cm(^{-1} )</th>
<th>( \nu_3(\text{CH}_3) ) cm(^{-1} )</th>
<th>( \nu(\text{OC}) ) cm(^{-1} )</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>On top methoxy species</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AuCe</td>
<td>2911</td>
<td>2884</td>
<td>2801</td>
<td>1105</td>
</tr>
<tr>
<td>AuCe10Co</td>
<td>2911</td>
<td>2881</td>
<td>2800</td>
<td>1105</td>
</tr>
<tr>
<td><strong>Doubly bridged methoxy species</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AuCe</td>
<td>2923</td>
<td>2834</td>
<td>2800</td>
<td>1050</td>
</tr>
<tr>
<td>AuCe10Co</td>
<td>2936</td>
<td>2838</td>
<td>2801</td>
<td>1053</td>
</tr>
<tr>
<td><strong>Doubly bridged methoxy species with different coordinations of Ce cations with oxygen ions</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AuCe</td>
<td>2910</td>
<td>2841</td>
<td>2807</td>
<td>1039</td>
</tr>
<tr>
<td>AuCe10Co</td>
<td>2911</td>
<td>2838</td>
<td>2809</td>
<td>1038</td>
</tr>
<tr>
<td><strong>Triply bridged methoxy species</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AuCe</td>
<td>2934</td>
<td>2847</td>
<td>—</td>
<td>1026</td>
</tr>
<tr>
<td>AuCe10Co</td>
<td>2949</td>
<td>2836</td>
<td>—</td>
<td>1023</td>
</tr>
</tbody>
</table>

**Figure 4.** FTIR difference spectra collected on AuCe (section a) and AuCe10Co (section b) in contact with the methanol/oxygen mixture after 10 min at r.t. (blue lines), at 50 °C (brown lines), 75 °C (violet lines), at 100 °C (orange lines) and after 20 min (red lines) at the same temperature.
Differently from what previously observed in a combined steady state isotopic transient kinetic analysis and *operando* FTIR study, bands related to the presence of carbonate species were also detected due to the adopted experimental conditions (static conditions rather than flow conditions reported in the literature). The presence of carbonates indicates that formate species are converted into CO$_2$, that evolves into surface carbonates in static conditions, whereas they would be flushed away in dynamic flow. The comparison between *in situ* and *operando* experiments indicates that both methoxy and formate species are intermediates and not only spectators.

The thermal methanol decomposition was also performed on both catalysts in order to compare the results with experiments carried out without oxygen: it was observed that on AuCe on top methoxy species disappeared along with the formation of formate species at 100°C, simultaneously to the conversion of doubly bridged methoxy with different coordination into more coordinated doubly bridged methoxy species (Figure S3). Therefore, the methoxy species with different coordinative unsaturation could be related to the formate production. On the contrary, in the case of AuCe10Co no formate species have been observed and only very intense bands due to carbonate-like species are produced (Figure S4), whereas the behaviour of the methoxy species is the same as observed on AuCe for the same reaction. These results demonstrate that in the absence of oxygen as a reactant, the AuCe10Co catalyst is able to efficiently provide oxygen to decompose methanol, further confirming that on this system the addition of Co causes a significant enhancement in the reducibility, which resulted in an improved oxidation activity. However, the same reaction carried out on the bare Ce10Co support did not give rise to any product (Figure S5) indicating that there must be a positive synergy involving gold to enhance oxygen mobility and activation.

It has been reported that promoting supported gold nanoparticles with lanthanum oxide largely increases the hydrogen selectivity in the partial oxidation of methanol, due to the fact that La$_2$O$_3$ was found to facilitate the reduction of initially present oxidic gold species. If compared with the results related to AuCe, the CO adsorption experiments at room temperature revealed the presence of more positivized gold sites on AuCe10Co before methanol oxidation reaction (Figure 3c). These sites could be responsible of the enhanced reactivity displayed by AuCe10Co and appear also to be reduced in higher extent after reaction (Figure 3d), demonstrating that an effect of the addition of cobalt on gold is occurring.

**Insights on the structure of the active sites**

In order to investigate the effect of the Co addition to the ceria substrate on the average local environment of supported gold (5, 10 and 15 wt.%) Au L$_3$ edge EXAFS experiments were performed on the AuCe, AuCe5Co, AuCe10Co and AuCe15Co catalysts previously submitted to a thermal treatment in O$_2$ at 400°C, that is the activation procedure to which the catalyst underwent before the catalytic tests. The K$^\text{3}$-weighted, phase uncorrected, FT of the EXAFS spectra collected on all catalysts are shown in Figure 5, section a. The analogous spectrum collected on Au foil is also reported for comparison (grey line).

All spectra are characterized by two main contributions. The first one, appearing in the 1.0-2.0 Å region and having a maximum at about 1.55 Å, is due to the single scattering Au-O of an Au oxide phase. The second one contributes in the 2.0-3.5 Å region and, independently from the presence of two maxima at 2.39, 2.98 Å, is mainly due to the first shell single scattering Au-Au of supported metal nanoparticles. This assignment is supported by the imaginary parts of the FT (see Figure S6). Moreover, the AuCe10Co catalyst shows the highest intensity of Au-O component due to the interaction between the active gold nanoparticles and the oxygen atoms of the oxidic support. This EXAFS evidence is in accordance with the FTIR characterization results here discussed and with previous high angle annular dark field observations that revealed the smallest average size of gold particles on this catalyst if compared to AuCe5Co and AuCe15Co.

The quantitative results obtained from the EXAFS fits on all the samples are summarized in Table 3. The quantitative results confirm what observed previously on the simple basis of the intensity of the Au-O component at 1.55 Å: the sample with the highest oxidized fraction is AuCe10Co, followed by AuCe5Co and then by AuCe15Co, that is comparable with the undoped AuCe. Conversely, no trend can be put in evidence for the Au-Au contributions that are formally equivalent within the experimental error bars. It should be noted that in this case the intensity of the Au-Au contribution is inversely proportional to the fraction of the oxidized phase discussed above but is also influenced by the different particle size distribution, so that the two effect cannot be directly disentangled in a straightforward manner. On top of these the region between 2.0-3.5 Å is expected to host also the higher shell contributions from oxide phase,
that have been neglected in these refinements. This fact also explains the minor discrepancy between experimental data and fit reported in Figure 5b.

Figure 5. Section (a): modulus of $k^2$-weighted, phase uncorrected, FT of the Au L$_3$-edge EXAFS spectra of the catalysts prepared on differently Co doped ceria substrate and submitted to thermal treatment in oxygen at 400°C. Corresponding imaginary parts are reported in Figure S5. Section (b): Comparison between the experimental (scattered black dots) and the best fit (red full line) for AuCe$_{10}$Co$_{ox}$. Also reported are the imaginary parts of the experimental data (scattered black dots) and the modulus (full lines) and the imaginary parts (dashed lines) of the individual Au-O (blue lines) and Au-Au (orange lines) contributions.

Table 3. Results of the fit performed on oxidized samples. $S_0^2 = 0.85$ fixed from the value extracted from the fit of the Au foil ($S_0^2 = 0.85\pm0.8$); $\Delta k = (3.3–10.0)$ Å$^{-1}$ and $\Delta R = (1.4–3.5)$ Å, resulting in a number of independent points larger than 11.

<table>
<thead>
<tr>
<th></th>
<th>CN</th>
<th>$\sigma^2$ (Å$^2$)</th>
<th>$R$ (Å)</th>
<th>$E_\sigma$ (eV)</th>
<th>$R$ factor</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Au–O</td>
<td>Au–Au</td>
<td>Au–O</td>
<td>Au–Au</td>
</tr>
<tr>
<td>AuCe</td>
<td>1.2±0.3</td>
<td>5.5±1.1</td>
<td>0.006±0.001</td>
<td>0.004±0.002</td>
<td>1.96±0.02</td>
</tr>
<tr>
<td>AuCe5Co</td>
<td>1.7±0.3</td>
<td>4.6±0.5</td>
<td>1.95±0.01</td>
<td>2.848±0.006</td>
<td>0.006</td>
</tr>
<tr>
<td>AuCe10Co</td>
<td>2.2±0.4</td>
<td>5.0±0.6</td>
<td>1.94±0.02</td>
<td>2.852±0.008</td>
<td>0.010</td>
</tr>
<tr>
<td>AuCe15Co</td>
<td>1.2±0.2</td>
<td>4.7±0.3</td>
<td>1.95±0.01</td>
<td>2.850±0.004</td>
<td>0.003</td>
</tr>
</tbody>
</table>

The methanol oxidation reaction was then investigated on Au10CoCe sample, being the most active catalyst: the first set of spectra was collected at room temperature. Furthermore, starting from the catalysts previously oxidized, another set of spectra was recorded under operando conditions, flowing methanol and oxygen, starting from room up to 120 °C. Au/CeO$_2$ was measured as reference sample, too. Three subsequent $\chi(k)$ spectra were measured at each temperature, to evaluate the experimental uncertainty. The data analysis was
performed on the averaged $\chi(k)$ spectrum by non-linear fitting procedures. Figure 6 shows the $k^3$ weighted, phase uncorrected FT (module and imaginary part) of the AuCe10Co sample collected during methanol oxidation reaction at 120°C (black scattered points). The signal due to the Au-O distance at 1.55 Å disappears during reaction\textsuperscript{50} while, a new contribution at about 1.8 Å is observed. The component at high distances (2.0-3.5 Å range) is clearly due to Au-Au contributions, whereas the new component around 1.8 Å is associated to an Au-Co contribution. The EXAFS fit of the experimental data performed assuming such structural model (red line in Figure 6) clearly shows the good agreement with the experimental spectrum, validating the presence of Co in the first coordination shell of a fraction of Au atoms. The quantitative results of the fit are reported in Table 4. The EXAFS study, corroborated by FTIR measurements, performed on all catalysts indicate that oxidized gold species are involved in the methanol reaction (Figure 5 and related data analysis in Table 3). In addition, the EXAFS analysis of the catalyst in working condition (Figure 6) suggests that the active sites present on the most active sample (with 10 wt% Co$_3$O$_4$) are those where gold sites are strongly interacting with Co. On these active sites reactive oxygen species, possibly close to Co sites can be activated and participate to the oxidative dehydrogenation reaction of methanol to formate species\textsuperscript{36}. Moreover, due to the synergic effect between cobalt and gold, the AuCe10Co catalyst is also able to provide reactive oxygen species in reaction conditions: reduced gold species close to oxygen vacancies and in strong interaction with cobalt ions are formed after reaction. Such Au species are those located at the interface with the support. These features can be at the origin of the enhanced catalytic activity of AuCe10Co.

**Table 4.** Result of the fit performed on AuCe10Co$_{met}$ EXAFS spectrum. $S_0^2 = 0.85$ fixed from the value extracted from the fit of the Au foil ($S_0^2 = 0.85\pm0.8$); $\Delta k = (3.3-10.0)$ Å$^{-1}$ and $\Delta R = (1.4-3.5)$ Å, resulting in a number of independent points larger than 11; R factor = 0.005.

<table>
<thead>
<tr>
<th>Paths</th>
<th>CN</th>
<th>$\sigma^2$ (Å$^2$)</th>
<th>$R$ (Å)</th>
<th>$E_0$ (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Au-Co</td>
<td>0.3 ± 0.1</td>
<td>0.010 ± 0.001</td>
<td>2.09 ± 0.03</td>
<td>3.6 ± 0.7</td>
</tr>
<tr>
<td>Au-Au</td>
<td>8.8 ± 0.8</td>
<td></td>
<td>2.835 ± 0.007</td>
<td></td>
</tr>
</tbody>
</table>

**Figure 6.** Modulus of the $k^3$-weighted, phase uncorrected, FT of the Au L$_3$-edge EXAFS spectrum of AuCe10Co collected during methanol oxidation reaction at 120 °C (black scattered dots) compared with the fit (red curve). Also reported are the imaginary parts of the experimental data (black scattered dots) and the modulus (full lines) and the imaginary parts (dashed lines) of the individual Au-Co (purple lines) and Au-Au (orange lines) contributions.

**Promotional role of Co on oxygen mobility**
Spectroscopic evidence of the effect of the support composition on the oxygen mobility is reported in Figure 7, where the evolution of the bands during CO and $^{18}$O$_2$ interaction on all the samples is shown starting from −180°C (orange lines, i.e. immediately after the inlet of $^{18}$O$_2$ on preadsorbed CO), during the heating (black and purple lines) and up to room temperature (blue lines). The inlet of $^{18}$O$_2$ at increasing diffusion times and temperature (black and purple lines up to blue line) over the samples previously saturated by CO at -180 °C (orange line) caused an erosion from the low-frequency side of the band at about 2102-2105 cm$^{-1}$, due to CO on Au particles covered by adsorbed oxygen\textsuperscript{37}, a
gradual decrease in intensity of the component at 2121-2127 cm\(^{-1}\) due to CO adsorbed on cationic gold clusters\(^{38,40}\) and the rapid depletion of the bands related to the support (component at 2150 cm\(^{-1}\) due to CO in interaction OH groups and bands at 2155-2161 cm\(^{-1}\) and at 2171-2173 cm\(^{-1}\) due to CO on different Ce\(^{4+}\) sites\(^{43}\)). Only in the case of AuCe10Co a band at 2183 cm\(^{-1}\), tentatively assigned to CO adsorbed on Ce\(^{3+}\) sites\(^{41}\) is observed during the heating at room temperature. Simultaneously, bands related to CO\(_2\) formation during CO oxidation are produced at higher frequencies (asymmetric CO\(_2\) stretching or \(v_3\) mode\(^{25}\)). Such absorptions have the highest intensity in the case of AuCe10Co (section c) and their intensity follows the same trend as for the catalytic activity: AuCe10Co > AuCe5Co > AuCe15Co> AuCe. More in detail, the three isotopomers of the CO\(_2\) molecule were observed: a growing band at 2326 cm\(^{-1}\), assigned to the C\(^{16}\)O\(^{18}\)O solid-like phase, accompanied by bands at 2340-2342 and at 2353-2357 cm\(^{-1}\), assigned to C\(^{16}\)O\(_2\) and C\(^{18}\)O\(_2\) solid-like phase, respectively\(^{53}\). The spectra showed a rapid exchange between the oxygen of ceria and the \(^{18}\)O molecules coming from the gas phase occurred already at low temperature. Moreover, the high intensity of the band ascribed to C\(^{16}\)O\(_2\) indicated that oxygen participating in the reaction at low temperature mainly comes from the support, putting in evidence the fact that the catalyst promotion with Co led to increased oxygen mobility because of the improved exchange properties of the ceria support.

The results demonstrated the enhanced ability of the Co-promoted-ceria support to supply active lattice oxygen that is beneficial for the reaction and were in good agreement with already published TPR results\(^{17}\) for highest hydrogen consumption and shift of \(T_{\text{max}}\) to lower temperature observed in the profile of AuCe10Co sample. Here we recall that these experiments were performed on catalysts oxidised at 400 °C, i.e. the temperature used for the activation before performing the catalytic tests on CH\(_3\)OH oxidation, see Figure 1, Table 1 and related discussion.

### On the stability of the catalysts

The analysis of the spectra reported in Figure 7 (right) indicated that on all the samples mainly two kinds of carbonate-like species were formed simultaneously to the production of CO\(_2\) (see the spectra reported in the right parts of Figure 7). Such species have different abundance, depending on the composition of the four catalysts. Hydrogen carbonates with peaks at 1600-1612 cm\(^{-1}\), 1392-1399 cm\(^{-1}\) and the component around 1020 cm\(^{-1}\) (\(\nu\text{CO}_3\) modes), at 820-823 cm\(^{-1}\) (\(\pi\text{CO}_3\)) and at 1211-1220 cm\(^{-1}\) (\(\delta\text{OH}\)) were detected\(^{53}\). In addition, bidentate carbonates are also produced, as revealed by the absorptions at 1568-1579 cm\(^{-1}\), at 1285-1298 cm\(^{-1}\) and at 1006-1009 cm\(^{-1}\) (\(\nu\text{CO}_3\) modes) and at 851-854 cm\(^{-1}\) (\(\pi\text{CO}_3\)).\(^{43}\) Please note that approximately in the 1350-1400 cm\(^{-1}\) region there must be the symmetric CO\(_2\) stretching (or \(v_3\) mode) of the CO\(_2\) molecules giving rise to the \(v_3\) mode (complex band at 2320-2360 cm\(^{-1}\) discussed above). For symmetry reason this band is IR inactive for CO\(_2\) molecules in the gas phase, but becomes IR active upon surface adsorption because of symmetry break. However, as the absorption coefficient of the \(v_3\) mode of adsorbed CO\(_2\) is typically one order of magnitude lower than that of the \(v_2\) mode\(^{55}\), meaning that we can neglect their presence in the spectra reported in the right parts of Figure 7.

Summarizing this isotopically labelled IR study, it is found that the presence of another oxide to ceria has a remarkable influence on the intensity of the bands in the carbonate region: on the AuCe catalyst (section a) the highest amount of hydrogen carbonate as well as of bidentate carbonate species is adsorbed on the surface after reaction if compared to the Co-promoted catalysts. Indeed, the intensity of such bands decreases with increasing the amount of produced CO\(_2\), giving evidence that cobalt oxide lowers the basicity of ceria and therefore the carbonate species formation results inhibited, despite the CO\(_2\) formation. AuCe10Co is the most active catalyst, but also the most stable one, due to the highest amount of produced CO\(_2\) and the lowest amount of carbonate-like species present at the surface after reaction. This represents a remarkable achievement for the promotion of the gold on ceria catalyst with Co, as in most of the cases in catalysis an increase of the catalyst activity is accompanied by a faster deactivation time and compromises must be found about these two aspects of the promotion.
Figure 7. Evolution of FTIR difference spectra collected on AuCe (section a), AuCe5Co (section b), AuCe10Co (section c) and AuCe15Co (section d) oxidised at 400 °C immediately after the inlet of $^{18}$O$_2$ at -180 °C on preadsorbed CO (orange lines) and at increasing contact times and temperature (black and purple lines) up to room temperature (blue lines) in the 2000–2400 cm$^{-1}$ (left) and in the 1800–800 cm$^{-1}$ (right) regions. The spectrum collected before CO dosage has been used to obtain the reported difference spectra, that have been normalised to the density of the pellets.

Conclusions
The catalytic performance in methanol oxidation of different gold catalysts supported on ceria modified by the addition of CoO$_4$ was investigated. Au/CeO$_2$ and Au/Co$_3$O$_4$ were also studied as reference samples. A strong influence of the amount of the Co promoter was observed. The activity trend followed the order: AuCe10Co > AuCe5Co > AuCe15Co > Au/CeO$_2$. TOFs have been calculated for all catalysts, and it was found that the TOF values obtained for AuCe10Co are one order of magnitude higher than those related to the other catalysts. Moreover, with the exception of AuCe10Co, for which the TOF value is almost unchanged, the conversion is strongly dependent on the temperature.

Different in situ FTIR experiments were performed. Adsorption of CO on AuCe and AuCe10Co indicate that oxidized gold sites are initially present on both activated samples, suggesting that such species are involved in the methanol reaction. Upon CH$_3$OH dosage, different methoxy groups are formed on both catalysts at r.t. (more abundantly on AuCe10Co). After successive O$_2$ addition they are consumed on both catalysts starting at about 75 °C and giving rise to mainly formate species on AuCe and to big amounts of different carbonate species on AuCe10Co. Moreover, the synergy between gold and Co-promoted ceria support causes significant enhancement of reducibility and capability for oxygen activation, which resulted in an improved oxidation activity, as demonstrated by the FTIR experiments carried out in the absence of oxygen. The capability of AuCe10Co to supply active oxygen species involves the participation of the oxygen vacancies on modified ceria as well as of the surface of separate, highly reducible, CoO$_x$ phase.

As suggested by EXAFS analyses, the active sites present on the best performing AuCe10Co catalyst are oxidized gold species at the interface with the support and close to Co sites, which are able to activate and to
react with oxygen giving rise to the formation of formate species and of carbonate species. The presence of a multiplet of bands in the CO\textsubscript{2} asymmetric stretching region after CO interaction with \textsuperscript{18}O\textsubscript{2} clearly reveals that the promotion of ceria by Co facilitates the exchange reaction with the \textsuperscript{16}O\textsubscript{2} atoms of the support. Moreover, it is demonstrated that the production of carbonate species is lowered in the presence of cobalt oxide, despite the CO\textsubscript{2} formation is increased. Such results give evidence of an improved stability of the Co-promoted catalysts.

The correlation of the catalytic activity and of a well-defined metal dispersion with the findings, provided by \textit{in situ} FTIR and \textit{operando} EXAFS studies about the nature and structure of the active sites allows to bridge the gap of experimental data previously obtained by HAADF, XRD, FTIR, TPR and to clarify the precise nature of the synergy between the gold phase and the Co-promoted ceria support, which resulted in an excellent CH\textsubscript{3}OH oxidation activity.

**Acknowledgements**

M. Manzoli and F. Vindigni gratefully acknowledge the IIT (Istituto Italiano di Tecnologia) for financial support (NANO GOLD Project). The Bulgarian authors gratefully acknowledge the financial support by the Bulgarian National Science Fund (Project DFNI T 02/4). C. Lamberti acknowledges the Mega-grant of the Russian Federation Government to support scientific research at the Southern Federal University, no. 14.Y26.31.0001.

**References**

Structure-reactivity relationship in Co$_3$O$_4$ promoted Au/CeO$_2$ catalysts for the CH$_3$OH oxidation reaction revealed by in situ FTIR and operando EXAFS studies

M. Manzoli*, F. Vindigni, T. Tabakova, C. Lamberti, D. Dimitrov, K. Ivanov, G. Agostini

* Department of Drug Science and Technology and NIS Interdepartmental Centre, University of Torino, 10125 Torino, Italy.

Department of Chemistry and NIS Interdepartmental Centre, University of Torino, 10125 Torino, Italy.

Institute of Catalysis, Bulgarian Academy of Sciences, 1113 Sofia, Bulgaria

Department of Chemistry, Agricultural University, 4000 Plovdiv, Bulgaria

Leibniz Institute for Catalysis at the University of Rostock, Albert-Einstein-Str. 29a, D-18059 Rostock, Germany

IRC “Smart Materials”, Southern Federal University, Zorge Street 5, 344090 Rostov-on-Don, Russia.

European Synchrotron Radiation Facility, 38043 Grenoble, France

mail to: *maela.manzoli@unito.it

Figure S1. FTIR difference spectra collected upon CO inlet (15 mbar) at r.t. and subsequent pressure decreases down to 1.0 x 10^-3 mbar on AuCe10Co before methanol oxidation reaction.
Figure S2. FTIR difference spectra collected on AuCe (section a) and AuCe10Co (section b) upon the adsorption of 5 mbar CH$_3$OH at r.t. (blue lines) and at increasing temperature after the inlet of 5 mbar O$_2$ on preadsorbed methanol: 50 °C (brown lines), 75 °C (violet lines), after 10 min (orange lines) and 20 min (red lines) at 100 °C.
Figure S3. FTIR difference spectra collected on AuCe in contact with 5 mbar methanol after 10 min at r.t. (blue line), at 50 °C (red line), at 75 °C (black line), at 100 °C (cyan line) and after 20 min at the same temperature (orange line).

Figure S4. FTIR difference spectra collected on AuCe10Co in contact with 5 mbar methanol after 10 min at r.t. (blue line), at 50 °C (violet line), at 75 °C (orange line) and at 100 °C (red line).
**Figure S5.** FTIR difference spectra collected on Ce10Co with the methanol/oxygen mixture after 10 min at r.t (black curve), at 50 °C (blue curve), at 75 °C (orange curve), and at 100 °C (red curve).

**Figure S6:** Imaginary part of k$^3$-weighted, phase uncorrected, FT of the Au L$_3$-edge EXAFS spectra of the catalysts prepared on differently Co doped ceria substrate and submitted to thermal treatment in oxygen at 400°C. Same data reported in Figure 5a.