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Insight into the true role of hydrogen-carbonate species in CO oxidation over Pd/Al₂O₃ catalyst using SSITKA-transmission IR technique

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ABSTRACT

The powerful combination of steady state isotopic transient kinetic analysis (SSITKA) and operando infrared spectroscopy (IR) was applied to study the CO oxidation reaction on Pd-supported Al₂O₃ catalyst at low temperature. The aim was to reveal the true role of hydrogen-carbonate species in the reaction mechanism. The SSITKA-IR experiments, conducted in presence and absence of CO₂ in the gas feed, have confirmed that hydrogen-carbonate species behave as spectator species in the reaction. Its formation was due to re-adsorption of the CO₂ product itself on the alumina surface as confirmed by the direct ^13CO_2/^12CO_2 exchange observed on the bare support.

1. Introduction

Increasing the level of pollution emitted by combustion engine vehicles has been accompanied by a strictness of global regulations to reduce gas emissions. Recently, the European commission has proposed the new Euro-7 standards bringing stricter emission limits for all types of internal combustion engines [1]. Amongst others, carbon monoxide emission standard for gasoline engines has been reduced to half (500 mg per km). In this work, a particular attention was paid to the short trips (5000 mg per trip, <10 km) impacted by cold-start and warm-up driving periods.

In three-way catalytic converters, CO oxidation is mainly catalyzed by supported palladium-based catalytic materials [2-4]. During the last decade, many efforts have been made to acquire a comprehensive knowledge regarding the catalytic chemistry of CO oxidation on Pd [5]. Investigating the nature of active sites and the role of surface adsorbed species formed during the reaction, namely carbonyls, formates, carbonates, and hydrogen-carbonates is critical in order to better understand the mechanism at hand and increase the activity of the catalyst at low temperatures (T < 200 °C). Recently, Murata et al. [6] have demonstrated that linear carbonyls adsorbed on Pd corner sites and Pd (111) facets the latter are highly active sites for CO oxidation at 130 °C. In their infrared spectroscopy investigation, Füttinger et al. [7,8] stated that CO₂ formation can also occur on Pd/alumina via a reaction between CO and hydroxyl groups on the support, whereas CO dissociation does not occur.

The combination of in situ and operando analysis techniques are needed to obtain the most accurate information on catalytic mechanisms. Steady State Isotopic Transient Kinetic Analysis (SSITKA) was developed by Happel [9], Bennett [10] and Biloen [11] proposed the combination of the advantages of both steady-state and transient kinetic regimes to better understand reaction mechanistic pathways. Coupled with infrared spectroscopy (IR), SSITKA-IR allows to determine kinetic parameters, as well as the nature of surface-adsorbed active and inactive species during the reaction [12,13]. This technique was applied successfully to confirm the role of hydrogen-carbonates as inactive species formed during CO oxidation reaction on Pt/Al₂O₃ [14]. The hydrogen-carbonates could then be formed by either spillover of adsorbed CO₂ on the alumina surface or by re-adsorption of gaseous CO₂ at the surface of the alumina support.

In this study, SSITKA-transmission IR experiments were performed to investigate the oxidation of CO on Pd/Al₂O₃, trying to elucidate the true involvement of hydrogen-carbonate species in the reaction mechanism at a low temperature.

2. Experimental

2.1. Catalyst preparation and characterizations

The Pd/Al₂O₃ catalyst was prepared by the impregnation of γ-Al₂O₃ support (supplied by the Centre de Recherches de SOLAIZE TOTAL Energies) with Palladium (II) acetylacetonate (Fluka) precursor salt to
obtain 1 wt% metal loading. The catalyst was then dried overnight at 85 °C, calcined 4 h under air gas flow at 500 °C, and finally reduced with H₂(g) at 400 °C/2.5 h. Final Pd loading was determined by ICP-OES 5110 (Agilent Technologies).

Nitrogen physisorption measurements were performed on a Micromeritics TriStar II instrument. The experiments were conducted using 0.2 g of powder sample, previously outgazed at 150 °C under vacuum for 2 h. Brunauer-Emmett-Teller (BET) theory was applied to calculate the specific surface area. Pore size distribution was obtained from the complete isotherms based on the Barret-Joyner-Halenda (BJH) theory.

H₂ chemisorption measurements were carried out at 0 °C in a Micromeritics Autochem II instrument using a pulse technique. Prior to the adsorption, the catalyst (50 mg) was pretreated under 5 vol% H₂/Ar at 400 °C for 5 h. The temperature of the catalyst sample was then reduced to 0 °C under Ar gas atmosphere in order to perform the chemisorption measurements during which H₂ pulses (5 vol% H₂ in Ar) were injected until saturation. The Pd dispersion was calculated based on the total volume of gas adsorption, assuming a Pd₃H ratio of 1.

2.2. SSITKA-IR experiments

Operando SSITKA-IR measurements were performed using a 7.5 mg square pellet (1.21 cm²) of Pd/Al₂O₃ catalyst. The system is described in the Supporting Information (Fig. S1). Briefly, the pellet was placed in the homemade transmission IR cell (dead volume < 0.5 cm³, KBr windows) connected upstream to an automatic pilot system for gas flow and switching valves control.

The catalyst was firstly reduced at 400 °C/5 h under 5% H₂/He (30 mL min⁻¹). Then, the SSITKA-IR study was performed by feeding 12 mL min⁻¹ of 2000 ppm CO/10% O₂/He gas mixture to the catalyst. The temperature was increased from 50 °C to 116 °C (1 °C/min). The occurrence of a hysteresis phenomenon is usually reported on noble metals during CO oxidation [15]. Thus, in order to reach steady-state reaction conditions, a 30-min period of isotherm equilibration was first carried out at 116 °C until CO conversion remained stable at 8.5%. The ¹²CO reactant was then replaced by 2000 ppm of its labeled counterpart ¹³CO at isobaric and isothermal conditions. Three isotopic ¹²CO → ¹³CO exchange gas switches were repeated to record the concentration evolution of reversibly adsorbed intermediates, and also check reproducibility (a true steady-state was achieved). The complete SSITKA step-gas switching procedure was as follows: 2000 ppm ¹²CO/10% O₂/He → 2000 ppm ¹²CO/10% O₂/5000 ppm Kr/7900 ppm CH₄/He. Kr and CH₄ were used as inert gas for kinetic analysis (MS) and direct comparison of MS (gas phase) and IR (adsorbed species) signals, respectively. SSITKA-IR analysis was also carried out at the same CO conversion, ca. 8.5%, at 131 °C in the presence of CO₂. In the latter case, the same experimental procedure was used with the only difference the addition of 2000 ppm of CO₂ in both feed gas streams. Moreover, a CO₂ SSITKA-IR was performed directly on the bare alumina support applying the following step-gas switch: 2000 ppm ¹²CO/10% O₂/He → 2000 ppm ¹³CO/10% O₂/5000 ppm Kr/7900 ppm CH₄/He at 116 and 131 °C.

Gas phase analysis of the gas composition towards the inlet and from the outlet of IR cell was performed by micro-gas chromatography (Varian 490) and quadrupole mass spectrometry (MS Pfeiffer QMS 200, 70 eV positive electronic impact). The catalytic surface was analyzed by Fourier transformed transmission IR spectroscopy (Thermo Scientific Nicolet 6700) using the rapid-scan mode (1 scan, resolution ~ 2 cm⁻¹) and a N₂-cooled MCT detector.

The CO conversion (χCO) was calculated as the percentage ratio between the amount of CO consumed at the considered temperature and the initial CO feed concentration. The surface residence times of surfaces adsorbed species, τ₀(CO) (CO-s) and τ₅ (active carbon-containing intermediates leading to CO₂) were determined from SSITKA-MS transient response curves by calculating the integral difference between the ¹²CO and CH₄ response curves, and that between the ¹³CO₂ and ¹³CO response curves, respectively. The concentration of reversibly adsorbed CO (N(CO)), their respective surface coverage (θ(CO)), and the concentration of C-containing intermediates (N(C)) were calculated using the following Eqs. (1)–(3) based on material balances [16]:

\[ N_{CO} (\mu \text{mol g}^{-1}) = \frac{F_T \cdot Y_{CO} \cdot (1 - \chi_{CO})}{W} \tau_{CO} \]  

\[ N_{C} (\mu \text{mol g}^{-1}) = \frac{F_T \cdot Y_{NCO}}{W} \tau_C \]  

\[ \theta_{CO} = \frac{N_{CO}}{N_{pd, surf}} \]  

where \( F_T \) is the total molar flow rate (μmol s⁻¹) of the feed gas stream, \( Y_{CO} \) is the mole fraction of CO in the feed, \( Y_{NCO} \) is the mole fraction of ¹³CO₂ at the outlet of the reactor in steady-state, \( W \) is the total mass of the catalyst pellet (7.5 × 10⁻³ g), and \( N_{pd, surf} \) is the molar amount of surface palladium per gram of catalyst (μmol Pd g⁻¹pd) calculated from the Pd dispersion and loading values. The steady-state turnover frequency of CO conversion, namely TOFₜₐₜ (μmol g⁻¹ s⁻¹), was calculated under the CO/O₂/He feed gas stream before the SSITKA experiment, according to Eq. (4) [17].

\[ \text{TOF}_{CO} = \frac{r_{CO}}{N_{pd, surf}} \]  

where \( r_{CO} \) is the steady-state rate of CO conversion in μmol g⁻¹ s⁻¹ (the calculation of \( r_{CO} \) is detailed in Supporting Information).

3. Results and discussion

3.1. Fresh catalyst

The final mass loading of Pd deposited on the alumina support was determined by ICP-OES analysis and was found to be 0.87 wt% Pd. BET analysis showed that the Pd/Al₂O₃ catalyst developed a high surface area, ca. 235 m² g⁻¹ and a pore volume of 0.79 cm³ g⁻¹, which are comparable to the area and pore volume of the support itself (γ-Al₂O₃), i.e. 225 m² g⁻¹ and 0.76 cm³ g⁻¹, respectively. Metal dispersion measured by H₂ chemisorption showed a Pd dispersion value of 71.1% corresponding to a mean Pd particle size of ~1.54 nm (see estimation details in the Supporting Information).

3.2. SSITKA-IR for CO oxidation

3.2.1. CO oxidation without CO₂ in the feed gas stream on Pd/Al₂O₃

The CO oxidation reaction was first performed without the presence of CO₂ in the feed gas mixture. The transient normalized concentration response curves of CH₄ (tracer gas), ¹²CO and ¹³CO₂ obtained during the SSITKA switch ¹²CO/O₂/He → ¹³CO/O₂/CH₄/Kr/He at 116 °C are presented in Fig. 1. The CO conversion here was limited to 8.5% to maintain differential reactor conditions, and the TOFₜₐₜ was estimated to be 0.035 s⁻¹. It should be mentioned that the CO conversion is equal to the CO₂ yield, suggesting the absence of coke formation. The results show that the ¹²CO transient response curve lags behind that of CH₄ due to the CO converted but also to a detectable concentration of reversibly adsorbed carbonyls (CO-s) formed during the reaction. In the same manner, the transient response curve of ¹³CO₂ lags behind that of ¹²CO response curve and this is attributed to the slower exchange of ¹²C-containing active reaction intermediates formed for their ¹³C-labeled counterparts before they lead to the formation of ¹³CO₂ (the product of reaction). It is worth to note that both isotopic transient response curves cross each other at the expected normalized value of 0.5, ensuring the right isobaric condition of the exchange (Fig. S2).

According to the SSITKA theory, the areas between the response of CH₄ tracer and ¹²CO₂, and that between the ¹³CO and ¹³CO₂ correspond to the mean surface residence time \( \tau \) of reversibly adsorbed CO (\( \tau_{CO} = \) \( \frac{F_T \cdot Y_{CO} \cdot (1 - \chi_{CO})}{W} \)).
Mean residence times, \( \tau \), concentrations \( N \), surface coverages, \( \theta \) of adsorbed CO and C-containing intermediates calculated from the SSITKA CO oxidation experiments on Pd/Al2O3 with and without the presence of CO in the feed.

<table>
<thead>
<tr>
<th>T (°C)</th>
<th>( \tau ) (s)</th>
<th>( N ) (μmol g(^{-1}))</th>
<th>( \theta ) (on Pd)</th>
<th>( \tau ) (s)</th>
<th>( N ) (μmol g(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Without CO2</td>
<td>116</td>
<td>8.6</td>
<td>17.4</td>
<td>0.30</td>
<td>55.5</td>
</tr>
<tr>
<td>With CO2</td>
<td>131</td>
<td>8.2</td>
<td>16.5</td>
<td>0.28</td>
<td>18.6</td>
</tr>
</tbody>
</table>

8.6 s) and active C-containing intermediates (\( \tau = 55.5 \) s) leading to CO\(_2\), respectively. The surface concentrations and surface coverages of these adsorbed species can be estimated using Eqs. (1)-(3). The estimated values are reported in Table 1. A value for the concentration \( N_{\text{CO}} \) of adsorbed CO-s \( \sim 17.4 \) μmol per gram of catalyst was obtained, corresponding to a surface coverage, \( \theta_{\text{CO}} \) (based on Pd) of \( \sim 30\% \). This value can be compared to the expected CO surface coverage based on a heat of CO adsorption of 29 kcal/mol reported by Szanyi et al. [18]. The theoretical CO surface coverage was estimated to be \( \sim 95\% \) at 116 °C on a bare Pd (100) surface. The lower value of \( \theta_{\text{CO}} \) on Pd found here could be partly explained by the competitive adsorption of oxygen, introduced in large amount, on the Pd surface sites. In addition, the concentration of C-containing intermediates, \( N_C \), was \( \sim 10.4 \) μmol per gram of catalyst.

IR transmission spectra were recorded before and after the step-gas switch to determine the chemical nature of adsorbed species during the CO oxidation reaction on Pd/Al2O3. Fig. 2 presents two spectra showing the composition of adsorbed phase on the surface of Pd/Al2O3 at 116 °C under the reaction mixtures of \( \text{12CO} + \text{2O} \) (in black) and \( \text{13CO} + \text{2O} \) (in red) at the steady state reaction conditions.

In the carbonyl region (2200–1700 cm\(^{-1}\)) shown in Fig. 2a, the IR band observed at 2088 cm\(^{-1}\) (black signal) before the isotopic switch is attributed to linear \( ^{12}\text{CO} \) adsorbed on Pd. According to the literature [19,20], linear CO are more likely to be adsorbed on metallic Pd\(^{6}\) particle corner sites or (111) facets. The larger IR signal observed between 2000 and 1800 cm\(^{-1}\) can be decomposed into several overlapped infrared bands corresponding to bridged and multi-bonded \( ^{12}\text{CO} \) adsorbed on palladium [6,21]. After applying the isotopic exchange (Fig. 2a, red curve), a complete red shift of the linear and bridged/multi-bonded CO bands adsorbed on Pd was observed, giving rise to new \( ^{13}\text{CO} \) IR bands at 2039 and 1950–1750 cm\(^{-1}\), respectively.

The formation of hydrogen-carbonate species on the alumina support is detected in the carbonate region (1700–1200 cm\(^{-1}\)) shown in Fig. 2b. The IR bands at 1657 and 1439 cm\(^{-1}\) seen in the black spectrum, were attributed to the asymmetric and symmetric stretching \( \nu(O\text{CO}) \) vibrations of hydrocarbon-species, respectively [7,22,23]. The latter are accompanied by a small band near 1231 cm\(^{-1}\) corresponding to the bending \( \delta(O\text{CO}) \) vibration. Unfortunately, due to the low extinction coefficient in the stretching OH region, the expected \( \nu(O\text{H}) \) band of these species was not clearly observed, as well as the negative OH band due to the consumption of a surface hydroxyl group present on the \( \gamma \)-alumina surface. Upon exchange with \( ^{13}\text{CO} \), the hydrogen-carbonate bands recorded at 1231, 1439 and 1657 cm\(^{-1}\) are in large part red-shifted providing the IR bands centered at 1224, 1400 and 1613 cm\(^{-1}\), respectively. Another IR band was detected at 1384 cm\(^{-1}\) in both spectra as shown in Fig. 2b, which does not appear to be impacted by the isotopic \( ^{13}\text{CO} \rightarrow ^{12}\text{CO} \) exchange, evidencing its spectator nature. In addition, gaseous methane was also detected after the transient exchange with \( ^{13}\text{CO} \) (red spectrum) by the appearance of an extremely weak absorbance at 1305 cm\(^{-1}\).

3.2.2. CO oxidation with CO2 present in the feed gas mixture on Pd/Al2O3

After adding 2000 ppm of \(^{13}\text{CO} \) in both reaction feeds, the SSITKA \( ^{12}\text{CO}/^{13}\text{CO}/^{12}\text{O}_2/\text{Kr}/\text{CH}_4/\text{He} \) exchange on Al2O3 was performed on Pd/Al2O3 to investigate again the CO oxidation and the role of co-fed CO\(_2\). The normalized transient response curves of CH\(_4\), \(^{13}\text{CO} \) and \(^{13}\text{CO}_2\) following the reactant isotopic switch are plotted as a function of time in Fig. 3. A similar CO conversion as in the previous experiment was maintained (\( \chi_{\text{CO}} = 8.8\% \)) by increasing the temperature to 131 °C. It is important to note that this increase in temperature will have an impact on the calculated kinetic parameters since the rate constant also increases. This apparent inhibitor effect of CO\(_2\) besides the experimental error, is surprising but it might be related to a change in the proportion of carbonyl species at the Pd surface. Such a change could be attributed to hysteresis-like phenomena [15]. Compared to the previous experiment, a similar value of TOF\(_{\text{CO}} \) was obtained in the presence of CO\(_2\) in the feed gas stream, ca. 0.035 s\(^{-1}\).

As a direct effect of adding CO\(_2\) in the feed gas stream, the area between the two \(^{13}\text{CO} \) and \(^{13}\text{CO}_2\) response curves decreased significantly, corresponding to a mean residence time (\( \tau_{\text{CO}} \)) of 18.6 s (three times lower than the value obtained without CO\(_2\) co-addition in the reaction feed gas mixture). The concentration of \( ^{13}\text{C} \)-containing intermediates (\( N_C \)) has decreased to the same extent, ca. 3.7 μmol per gram of catalyst. It should be noted that the total number of C-containing intermediates should not change, except for the variation due to the temperature effect, since the pool of C-containing intermediates is simultaneously exchanged under the \(^{13}\text{CO}_2\) co-feeding SSITKA experiment, which is not taken into account in this case. The different kinetic parameters obtained in the presence and absence of CO\(_2\) are summarized in Table 1.

On the contrary, the presence of CO\(_2\) did not have a significant impact on the mean surface residence time and the concentration of reversibly adsorbed CO-s. Only a small decrease of \( \tau_{\text{CO}} = 8.2 \) s and \( N_{\text{CO}} = 16.5 \) μmol g\(^{-1}\) was observed in the presence of CO\(_2\) due to the very slight increase in temperature (change in reaction rate constant, \( k \)) and CO conversion (mainly effect of partial pressure of CO).

The evolution of adsorbed species on the catalytic surface has been followed by transmission IR during the isotopic exchange in order to observe the effect of the presence of CO\(_2\). The IR spectra recorded at 131 °C in both \(^{13}\text{CO} + \text{2O} \) and \(^{12}\text{CO} + \text{2O} \) and \(^{13}\text{CO} + \text{2O} + ^{12}\text{CO}_2 \) steady states are presented in Fig. 4.

Fig. 4a shows the red isotopic shift of the IR bands corresponding to linear, bridged and multi-bonded carboxyls on Pd, ca. from 2088 cm\(^{-1}\) and 2090–1800 cm\(^{-1}\) to 2039 cm\(^{-1}\) and 1950–1750 cm\(^{-1}\) range, respectively. In contrast, in the carbonate region, IR bands at 1657, 1439 and 1231 cm\(^{-1}\) (Fig. 4b) previously assigned to hydrogen-carbonates species, were only \(^{13}\text{C} \rightarrow ^{12}\text{C} \) shifted to a very small extent (<1.5%) in the presence of CO\(_2\) in the reactant feed gas mixture. Again, no shift was observed for the small IR band appeared at 1384 cm\(^{-1}\).

3.2.3. CO exchange on Al2O3 support

In order to further confirm the formation route of
hydrogen-carbonate species, a $^{12}\text{CO}_2/^{13}\text{CO}_2$ SSITKA-IR exchange was performed directly on the bare Al$_2$O$_3$ support. The transient normalized concentration response curves of CH$_4$ and $^{13}\text{CO}_2$ obtained on the alumina following the $^{12}\text{CO}_2$/O$_2$/He → $^{13}\text{CO}_2$/O$_2$/Kr/CH$_4$/He switch at 116 and 131 °C are presented in Fig. S3 (Supporting Information). The comparison with previous results of the CO oxidation on Pd/Al$_2$O$_3$ in the presence of CO$_2$ (Fig. 3) are shown in Fig. 5 a. A clear match between both $^{13}\text{CO}_2$ profiles (blue and orange curves) was observed, concomitant with the overlapping methane profiles (black and green curves). Similar surface residence times between CH$_4$ and $^{13}\text{CO}_2$ were estimated in both cases, namely: $\tau_{\text{CO}_2}=28.8$ s on Al$_2$O$_3$, and $\tau_{\text{CO}_2}=26.7$ s on Pd/Al$_2$O$_3$.

The evolution of surface adsorbed species has also been followed on bare Al$_2$O$_3$ by transmission IR during the $^{12}\text{CO}_2/^{13}\text{CO}_2$ isotopic exchange. The IR spectra recorded in the 1700–1200 cm$^{-1}$ range in both $^{12}\text{CO}_2$/O$_2$ and $^{13}\text{CO}_2$/O$_2$ steady states are presented in Fig. 5 b. As previously, $^{12}$C-containing hydrogen-carbonate species formation is detected at 1655, 1442 and 1231 cm$^{-1}$ (black spectrum) and which is attributed to the $\nu$(OCO)$_{\text{sym}}$, $\nu$(OCO)$_{\text{sym}}$ and $\delta$(OH) vibrations respectively. Hydrogen-carbonates (HCO$_3$(ads)) are formed following the Eq. (5), where an OH surface group on the alumina surface interacts with CO$_2$.

$$\text{CO}_2(\text{g}) + \text{OH}_{(\text{surf})} \rightarrow \text{HCO}_3(\text{ads})$$

(5)

Upon exchange with $^{13}$CO$_2$, the latter bands were red-shifted towards the bands at 1608, 1402 and 1224 cm$^{-1}$ (red spectrum),
respectively, corresponding to $^{13}$C-containing hydrogen-carbonate species. Difference spectrum was produced and shown in Fig. S3b, which clearly showed the partial disappearance of $^{13}$C-containing hydrogen-carbonate vibrations (negative bands) towards the appearance of their $^{13}$C counterpart. Another IR band appeared at 1474 cm$^{-1}$ in both spectra was attributed to undetectable carbonate species. Its intensity decreases after the $^{13}$CO$_2$ switch but this band doesn’t seem to be isotopically shifted towards lower wavenumbers (Fig. S3b, SI). Similar trends were also observed at 116 °C ($t_{CO2} = 26.3$ s).

3.3. General comments

In this study, the SSITKA technique was successfully coupled to transmission IR spectroscopy in order to elucidate the role of adsorbed species formed on Pd/Al$_2$O$_3$ during the CO oxidation reaction. The measurement of kinetic parameters, such as the mean residence times and concentrations of surface adsorbed species on Pd/Al$_2$O$_3$ have been combined with the determination of their nature. Mainly linear, bridged, and multi-bonded carbonyl-type species were formed during the CO adsorption on metallic Pd$^0$ at low temperature. The corresponding $\nu(13C=O)$ IR bands observed at 2088 cm$^{-1}$ and in the 2000–1800 cm$^{-1}$ range (Fig. 2a) under $^{12}$CO reactant feed disappeared completely after ~2 min after the isotopic transient switch towards $^{13}$CO, giving rise to the formation of new $\nu(13C=O)$ bands at 2039 cm$^{-1}$ and 1950–1750 cm$^{-1}$. This trend is consistent with the gas phase evolution of $^{13}$CO observed by mass spectrometry (Fig. S4), where it reached its maximum concentration after 150 s. The fast $^{12}$C $\rightarrow$ $^{13}$C isotopic exchange of carbonyls observed in SSITKA-IR high the exchange of molecularly adsorbed $^{12}$COs with gaseous $^{13}$CO and their reactivity towards subsequent reaction intermediates formed in the mechanism of CO oxidation on Pd/Al$_2$O$_3$.

Initially, the normalized concentrations of $^{13}$CO and $^{13}$CO$_2$ response curves showed a significant time lag (Fig. 1), indicating that in theory the formation of CO$_2$ product passes through various active intermediates after CO adsorption on Pd. The operando IR results (Fig. 2b) reveal that hydrogen-carbonate species are formed on the alumina surface and follow a $^{12}$C $\rightarrow$ $^{13}$C isotopic red-shift exchange during the experiment. Other weak absorption bands were also observed by IR on the catalyst surface in the carbonate region but these were clearly identified as spectator species in the CO oxidation at low temperatures due to their inability to follow the isotopic exchange. It is worth to note that the $^{12}$C $\rightarrow$ $^{13}$C exchange of the hydrogen-carbonate species observed by IR on alumina is very slow compared to surface carbonyl exchange, and even incomplete after 5 min of reaction (Fig. S4 –1640 cm$^{-1}$ curve). Moreover, this trend is not consistent with the transient response curve recorded for $^{13}$CO$_2$ product formation by mass spectrometry.

The addition of 2000 ppm of CO$_2$ in the feed gas stream proved unequivocally against the previous SSITKA-IR conclusion about the active role of hydrogen-carbonate species in CO oxidation reaction on Pd/Al$_2$O$_3$. Indeed, the presence of CO$_2$ has led to a drastic reduction (three times) of the amount of $^{13}$C-containing intermediate species (Nc) compared to the experiment performed without CO$_2$ at similar CO conversion. Also, the IR spectra recorded in the presence of CO$_2$ in the carbonate region (Fig. 4b) show negligible $^{12}$C $\rightarrow$ $^{13}$C isotopic shift of the IR bands attributed to hydrogen-carbonates. This result confirms clearly in coherence with the kinetic calculation that the hydrogen-carbonate species formed during CO oxidation reaction is only a spectator species, and this does not participate in the reaction mechanism of CO$_2$ formation. This conclusion is also supported by the lag observed in Fig. 5 between the relative rate of exchange of $^{13}$CO$_2$-MS signal and the 1640 cm$^{-1}$ IR band intensity evolution curve of hydrogen-carbonate species. Therefore, most of the significant time lag firstly observed between $^{13}$CO- and $^{13}$CO$_2$-MS signals during the SSITKA experiment without CO$_2$ in the feed (Fig. 1) could be attributed to the re-adsorption of CO$_2$ produced by the CO oxidation, on the surface of the alumina support rather than to the formation of some active intermediate species towards CO$_2$ formation. The overestimation of SSITKA C-pool due to CO$_2$ re-adsorption effect was previously reported by Efstathiou [24] in the case of the water-gas shift reaction at high temperature. Similar conclusion was previously obtained for CO oxidation reaction on a Pt-supported alumina catalyst using the same experimental procedure [14]. Linear carbonates were predominant on the Pt surface in contrast to Pd, where bridged carbonyls were mainly observed. The hydrogen-carbonate species behaved similarly on both Pt/Al$_2$O$_3$ and Pd/Al$_2$O$_3$ catalysts suggesting the re-adsorption of CO$_2$ on the alumina support rather than a direct spillover from the metal particles. This hypothesis was confirmed in this work by performing $^{12}$CO$_2$/$^{13}$CO$_2$ isotopic exchange experiments on bare alumina (Fig. 5). A direct exchange was observed between $^{13}$C-containing hydrogen-carbonates and their $^{13}$C-labeled counterpart on bare Al$_2$O$_3$ highlighting the lack of metal contribution in this process.

It is also important to note that the value of N$_c$ did not drop to 0 with the pre-saturation of the surface by CO$_2$ (N$_c$ = 3.7 μmol g$^{-1}$). This observation means either a partial saturation of alumina with carbonate
species by the use of 2000 ppm of $^{12}$CO$_2$ which would be complemented by the formation and re-adsorption of $^{13}$CO during the experiment, or that other C-containing intermediate species could be involved in the CO oxidation mechanism. In the second case, NC must then be carefully described as consisting of two pools, namely: (i) spectator hydrogen-carbonates, and (ii) active C-containing intermediate species not of the nature of hydrogen-carbonates.

Carbonyls are likely the main active reaction intermediates (CO-s) identified by IR spectroscopic means. Various types of carbonyls can be identified in this work such as linear or bridged carbynols. Even if all species follow a similar $^{13}$CO/$^{12}$CO isotopic exchange, they are not necessarily involved in the reaction pathway. Indeed, one of them can be a spectator with a fast conversion into the reactive species. This work paves the way for further analysis to discriminate the individual role(s) of each carbonyl species according to their adsorption sites during the CO oxidation reaction, as well as their deactivation process. This will be the aim of further study.

4. Conclusions

A full understanding of CO oxidation mechanism on PGM supported catalysts at low temperature (< 200 °C) is of great importance in order to improve catalyst efficiency and decrease atmospheric pollution levels. The use of operando SSITKA-IR in the current study has provided valuable information into the role of the carbonyl and carbonate species formed at the surface of Pd/Al$_2$O$_3$ catalyst during the CO oxidation reaction. The dynamic exchange between $^{12}$CO $+$ O$_2$ and $^{13}$CO $+$ O$_2$ in the presence and absence of CO$_2$ in the reaction gas mixture revealed a fast isotopic shift of the IR band signals related to various types of carbonyl species adsorbed on metallic Pd, suggesting that at least one of them could be an active intermediate in the CO$_2$ formation. By contrast, hydrogen-carbonate species were finally found to behave like spectator species since only negligible isotopic exchange of corresponding IR absorption bands were observed in the presence of CO$_2$ in the feed gas stream. The lag between the $^{12}$CO and $^{13}$CO observed on SSITKA normalized concentration curves (no CO$_2$ was added in the feed) was then assigned to the re-adsorption of the CO$_2$ product on the bare alumina support by interaction with a surface hydroxyl group. The direct $^{12}$CO$_2$/$^{13}$CO$_2$ exchange observed on alumina clearly supports this conclusion, indicating the absence of the metal in the formation path of hydrogen-carbonate species.

Statements and declarations

The authors have no financial or non-financial interest to disclose. The data generated during the current study are available from the corresponding author on reasonable request.

CRediT authorship contribution statement

Ibrahim Hatoum: Conceptualization, Data curation, Methodology, Investigation, Writing – original draft. Méliandre Richard: Conceptualization, Funding acquisition, Methodology, Investigation, Supervision, Validation, Writing – review & editing. Christophe Dujardin: Conceptualization, Funding acquisition, Methodology, Investigation, Supervision, Validation, Writing – review & editing.

Declaration of Competing Interest

None.

Data availability

Data will be made available on request.

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Appendix A. Supplementary data

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References