

Effect of CO₂, H₂O and SO₂ in the ceria-catalyzed combustion of soot under simulated diesel exhaust conditions.

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Abstract

The effect of CO₂, H₂O and SO₂ in the Ce_{0.73}Zr_{0.27}O₂ and Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ catalyzed combustion of soot with NO_x + O₂ has been studied. Combustion experiments performed in a fix-bed reactor with soot-catalyst mixtures prepared in loose contact mode showed that CO₂, H₂O and SO₂ lower the activity of both catalysts, and the inhibiting effect follows the trend SO₂ > H₂O > CO₂. Regardless the gas mixture composition, the catalytic activity for soot combustion of Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ is equal or higher to that of Ce_{0.73}Zr_{0.27}O₂ because Nd³⁺ doping seems to promote the participation of the active oxygen mechanism together with the NO₂-assisted mechanism in the catalytic combustion of soot. The maximum soot combustion rate achieved during a Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂-catalyzed reaction in NO_x/O₂/CO₂/H₂O/N₂ is about three times higher than that of the uncatalyzed combustion, and this catalyst also improves the CO₂ selectivity.

In situ DRIFTS experiments showed that CO₂, H₂O and SO₂ compete with NO_x for the adsorption sites on the catalysts' surface. CO₂ partially impedes the catalytic oxidation of NO to NO₂, affecting much more to the Nd³⁺-containing catalyst; however, the contribution of the active oxygen mechanism seems to remain relevant in this case. H₂O also hinders the catalytic oxidation of NO to NO₂ on both catalysts, and therefore the catalytic combustion of soot, because delays the formation of nitrogen reaction intermediates on the catalysts' surface and favors the formation of more stable nitrogen surface species than in a H₂O-free gas stream. For both catalysts, SO₂ chemisorption (with sulfate formation) is even able to remove nitrogen surface groups previously formed by NO_x chemisorption, which significantly inhibits the catalytic oxidation of NO to NO₂ and the catalytic combustion of soot.

Keywords: Diesel soot; soot combustion; ceria-zirconia catalyst; neodymium-ceria catalyst; nitrogen oxides.

38 **1.- Introduction.**

39

40 Catalytic combustion of diesel soot is a topic of ongoing research [1-4],
41 and different catalysts have been investigated for this purpose at laboratory
42 scale. These catalysts include formulations with noble metals, alkali metals,
43 alkali earth metals, and transition/internal transition metals that can accomplish
44 redox cycles (V, Mn, Co, Cu, Fe, Ce, Pr, etc) [5].

45 Platinum catalysts seem the most interesting for practical applications
46 because they combine high activity and stability [6-8], but due to the high price
47 of noble metals, cheaper catalysts are being investigated. Considering the
48 laboratory results obtained by several research groups [4, 5, 9-11], ceria-based
49 catalysts have been proposed as promising substitutes of platinum catalysts.
50 However, extrapolation of laboratory results to real conditions must be done
51 having in mind that, to obtain laboratory results with practical relevance, several
52 aspects must be taken into account.

53 Attention must be paid to the soot-catalyst contact, and it is usually
54 accepted that “loose contact” mixtures of soot and powder catalysts (soot and
55 catalyst are mixed with a spatula, for instance) simulate the poor contact
56 attained in a real diesel particulate filter.

57 Thermal stability of catalysts is also a major issue to pay attention to in
58 laboratory studies. Usually, the exhaust temperature in a diesel vehicle is lower
59 than 500 °C, but hot spots are formed during the regeneration of the diesel
60 particulate filters where the temperature can increase out of control.

61 The presence of NO_x in the gas mixture also plays a key role in the
62 catalytic combustion of soot, and laboratory experiments with practical interest

63 are usually performed with a gas mixture of NO + O₂. NO₂ can be produced in
64 these mixtures, which reacts with soot much faster than NO and O₂.

65 All these aspects have been taken into account in many articles devoted
66 to the study of soot combustion ceria-based catalysts at laboratory [9], but not
67 so attention has been paid to the effect of CO₂, H₂O and SO₂ in the catalytic
68 performance of ceria catalysts, despite these gases are typically present in a
69 diesel exhaust gas.

70 Some authors have reported the poisoning effect of SO₂ on ceria-
71 containing soot combustion catalysts [12-15]. Weng et al. [13] studied the
72 catalytic combustion of soot in NO_x + O₂ with Cu-K/CeO₂ catalysts that were
73 previously treated in a 400 ppm SO₂ stream at 400 °C, and a decrease on the
74 activity was noticed. This deactivation was attributed to potassium sulfate
75 formation, but the effect SO₂ on the ceria behavior was not discussed.

76 The effect of H₂O and SO₂ on the stability of Ba,K/CeO₂ catalysts during
77 diesel soot combustion was also studied [14], and it was concluded that these
78 catalysts had good tolerance to water at low temperatures (e.g., 400 °C) while
79 high concentrations of SO₂ lead to rapid deactivation. The formation of
80 potassium, barium and cerium sulfates was reported, but the particular effect of
81 H₂O (if any) and SO₂ on the performance of ceria was not analyzed in detail.

82 MnO_x-CeO₂ mixed oxides were also tested for the low-temperature
83 oxidation of diesel soot in a O₂ + H₂O + NO + N₂ gas stream, and the effect of
84 SO₂ was also studied [15]. The poisoning effect of SO₂ was also reported in this
85 case, but a detailed analysis of its effect on ceria was not provided apart from
86 the evidences obtained by thermogravimetry about sulfates formation.

87 The goal of the current study is to analyze the effect of CO₂, H₂O and
88 SO₂ in the ceria-catalyzed combustion of soot in simulated diesel exhaust
89 conditions. Taking into account our previous results [16, 17], and in order to
90 provide practical relevance to the study, two catalysts with composition
91 Ce_{0.73}Zr_{0.27}O₂ and Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ have been selected for this study, since
92 they combine a good thermal stability and catalytic activity for soot combustion.
93 As previously concluded, Zr⁴⁺-doping is mandatory for a suitable thermal
94 stability of ceria, and Nd³⁺-doping further improves the activity [17]. Not only a
95 descriptive discussion of the CO₂, H₂O and SO₂ effect on the catalytic soot
96 combustion has been done based on fix-bed reactor experiments, but special
97 attention has been paid to the effect of these gases in the soot combustion
98 mechanism. For this purpose, *in situ* DRIFTS experiments have been carried
99 out.

100

101 **2.- Experimental.**

102

103 *2.1. Catalysts used.*

104

105 Details about the preparation and characterization of the Ce_{0.73}Zr_{0.27}O₂
106 and Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ catalysts were previously reported [17]. In brief, the
107 required amounts of Ce(NO₃)₃·6H₂O (Sigma Aldrich, 99%), Nd(NO₃)₃·6H₂O
108 (Aldrich, 99.9%) and/or ZrO(NO₃)₂·xH₂O (Fluka, x ≈ 6) were dissolved in water
109 and co-precipitation was carried out by dropping an ammonia solution to keep
110 the pH at about 9. After filtering, the precipitate was dried at 110 °C in air
111 overnight and calcined in air at 800 °C for 90 min.

112 Formally, the stoichiometric coefficient of oxygen on $\text{Ce}_{0.64}\text{Zr}_{0.27}\text{Nd}_{0.09}\text{O}_2$
113 should be lower than 2, since the tetravalent cation “ Ce^{4+} ” is replaced by a
114 trivalent one (Nd^{3+}). However, the subscript 2 has been maintained for the sake
115 of simplicity.

116

117 *2.2 Catalytic tests in a fix-bed reactor.*

118

119 Catalytic tests were performed in a tubular quartz reactor coupled to
120 specific NDIR-UV gas analyzers for CO, CO₂, NO, NO₂, SO₂ and O₂ monitoring
121 (Fisher–Rosemount, models BINOS 100, 1001 and 1004). 20 mg of soot, 80 mg
122 of catalyst, or 20 mg of soot + 80 mg of catalyst mixed in the so-called loose
123 contact mode [18] were used in these experiments (and also an empty reactor).
124 In all cases, the samples were diluted with 300 mg of SiC to avoid pressure
125 drop and favor heat transfer. The model soot used in this study is a carbon
126 black by Evonik-Degussa GmbH (Printex U).

127 Temperature programmed reactions were performed from room
128 temperature until 700 °C at 10 °C/min with the gas mixtures NO_x/O₂/N₂,
129 NO_x/O₂/CO₂/N₂, NO_x/O₂/H₂O/N₂ and NO_x/O₂/SO₂/N₂. Total flows of 500 ml/min
130 were used (GHSV = 30000 h⁻¹), and the compositions of the different gases in
131 these mixtures were 500 ppm NO (~ 0 ppm NO₂), 5 % O₂, 4 % CO₂, 2 % H₂O
132 and 80 ppm SO₂ with N₂ balance.

133 Two additional soot combustion experiments were performed without any
134 catalyst and with $\text{Ce}_{0.64}\text{Zr}_{0.27}\text{Nd}_{0.09}\text{O}_2$ under a NO_x/O₂/CO₂/H₂O/N₂ gas flow.
135 These experiments consisted of raising the temperature from room temperature
136 until 530 °C at 10 °C/min, keeping the maximum temperature stable for 30 min.

137

138 *2.3 In situ DRIFTS experiments.*

139

140 A FTIR spectrometer (Jasco, model FT/IR-4100) was used for *in situ*
141 DRIFTS experiments, with a reaction cell that allows temperature and gas flow
142 composition control. The reactor is designed to allow the gas to flow through
143 the catalytic bed. The gas mixtures NO_x/O₂/N₂, NO_x/O₂/CO₂/N₂,
144 NO_x/O₂/H₂O/N₂ and NO_x/O₂/SO₂/N₂ were studied, with the same concentration
145 of the individual gases used in the previously described catalytic tests.

146 The experiments consisted of heating the catalysts (without soot) until
147 460 °C under He flow, and a background spectrum was recorded in these
148 conditions for each catalyst. Then, the inert gas was replaced by the selected
149 reactive gas mixture and spectra were recorded after different times. The
150 background spectra have been subtracted in all spectra shown in this article,
151 and therefore, only bands attributed to adsorbed species are observed.

152

153 **3.- Results and discussion.**

154 *3.1. Catalytic tests in the fix-bed reactor.*

155 Figure 1 shows the soot conversion profiles obtained from CO + CO₂
156 evolved in combustion experiments performed with different gas mixtures,
157 where the effect of CO₂ (Figure 1b), H₂O (Figure 1c) and SO₂ (Figure 1d) in
158 soot oxidation under NO_x and O₂ containing gas streams is analyzed.

159 As it was concluded in a previous study [17], both catalysts decrease the
160 soot combustion temperature with regard to the uncatalyzed reaction in the

161 NO_x/O₂/N₂ gas mixture (Figure 1a), and Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ is more active than
162 Ce_{0.73}Zr_{0.27}O₂. The enhanced activity of the Ce-Zr mixed oxide upon Nd³⁺
163 doping was attributed to the creation of oxygen vacancies, which improve the
164 redox properties of the mixed oxide

165 As a general trend, the presence of either CO₂ (Figure 1b), H₂O (Figure
166 1c) or SO₂ (Figure 1d), in addition to NO_x and O₂, decrease the activity of both
167 catalysts with regard to the reaction under NO_x/O₂/N₂ (Figure 1a), while
168 differences in the uncatalyzed reactions are not obvious for the different gas
169 mixtures tested. The strongest inhibitor effect on the catalyzed reactions is
170 attributed to SO₂, followed by H₂O and finally CO₂. On the other hand,
171 regardless the gas mixture used, the catalytic activity for soot combustion of
172 Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ is equal or higher to that of Ce_{0.73}Zr_{0.27}O₂.

173 In a previous study [17] performed with Ce_{0.73}Zr_{0.27}O₂ and
174 Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂, and also in many others studies performed with different
175 soot combustion catalysts [6, 19, 20], the catalytic oxidation of NO to NO₂ was
176 identified as an important step in the catalyzed soot combustion mechanism,
177 since NO₂ is more oxidizing than NO and O₂. This is the so-called NO₂-assisted
178 soot combustion mechanism, where the catalyst accelerates the oxidation of
179 NO to NO₂, NO₂ ignites soot and O₂ is able to continue the combustion once
180 this has started. In order to explore the reasons of the inhibiting effects of CO₂,
181 H₂O and SO₂, the NO₂ profiles obtained in oxidation experiments performed
182 without (Figure 2) and with soot (Figure 3) have been analyzed.

183 All NO₂ profiles obtained in the absence of soot (Figure 2) are
184 qualitatively similar. In spite of thermodynamics predict that NO₂ should be the
185 main NO_x component at low temperature, NO₂ concentration is null at the

186 beginning of the experiments. This is because the experimental set-up used in
187 this study has been designed to minimize the gas phase oxidation of NO to NO₂
188 before the reactor, in order to mimic real diesel exhausts.

189 Both catalysts accelerate the oxidation of NO to NO₂ in all the gas
190 mixtures until the thermodynamic equilibrium is achieved at a certain
191 temperature, which has been defined as T_{max}NO₂, and above this temperature
192 NO₂ decreases following the thermodynamic equilibrium. Both the catalyst
193 nature and the gas mixture affect the onset NO oxidation temperature (which is
194 not shown in all curves of Figure 3 for clarity), the T_{max}NO₂ and the maximum
195 NO₂ level achieved.

196 In accordance with the soot combustion experiments (Figure 1a),
197 Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ accelerates NO₂ production more efficiently than
198 Ce_{0.73}Zr_{0.27}O₂ in the NO_x/O₂/N₂ gas mixture (Figure 2a). On the contrary,
199 Ce_{0.73}Zr_{0.27}O₂ is more active than Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ under NO_x/O₂/CO₂/N₂
200 (Figure 2b) and NO_x/O₂/SO₂/N₂ (Figure 2d), and both catalysts behave equal
201 under NO/O₂/H₂O/N₂ (Figure 2c). This means that there is not a direct
202 relationship between NO₂ production and soot combustion in this case, as it will
203 be discussed in detail afterwards, in spite of the catalytic oxidation of NO to NO₂
204 is one of the reactions involved in the soot combustion mechanism. This is
205 confirmed by the shape of the NO₂ profiles obtained in the presence of soot
206 (Figure 3), which is different to that obtained in the absence of soot because
207 part of the NO₂ produced by catalytic oxidation of NO is consumed by the NO₂-
208 soot reaction. The main nitrogen product of the NO₂-carbon reaction is NO, and
209 therefore net NO_x (= NO + NO₂) removal is quite low. NO_x removal curves are

210 included in Figure 1SM of the supplementary material, where it is shown that
211 NO_x removal level is always lower than 10%.

212 As a summary, the catalytic oxidation of NO to NO₂ is one of the
213 reactions involved in the soot combustion mechanism in all the gas mixtures
214 tested, and the inhibiting effect of the gases tested on both NO₂ production and
215 soot combustion follows the trend: SO₂ > H₂O > CO₂. However, if the behavior
216 of both catalysts towards NO₂ production and soot combustion is compared, it is
217 concluded that soot combustion not only depends on NO₂ production. In order
218 to analyze this in more detail, the relationship between soot combustion
219 (represented by the T50% (°C) temperatures determined from Figure 1) and
220 NO₂ production (represented by the T_{max} NO₂ (°C) determined from Figure 2)
221 is plotted on Figure 4. Quite linear relationships are obtained for both catalysts,
222 but data obtained with Ce_{0.73}Zr_{0.27}O₂ are always above those obtained with
223 Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂. This confirms that NO₂ production is actually a key step in
224 the catalytic combustion of soot. However, for similar NO₂ production,
225 Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ is more effective for soot combustion than Ce_{0.73}Zr_{0.27}O₂,
226 and this suggests the additional participation of the “active oxygen” mechanism
227 in the soot combustion reactions. It is known that ceria-catalysts are able to
228 exchange oxygen with gas phase O₂ molecules [21-23], and deliver highly
229 reaction active oxygen species to soot, and also to NO to be oxidized to NO₂.
230 This additional mechanism would explain the differences between
231 Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ and Ce_{0.73}Zr_{0.27}O₂, since the introduction of Nd³⁺ on the Ce-
232 Zr mixed oxide improves the oxide reducibility and favors vacant sites
233 formation, [17] and these improvements promote active oxygen production.
234 Therefore, Nd³⁺ doping is expected to promote the participation of the active

235 oxygen mechanism besides the NO₂-assisted mechanism in the catalytic
236 combustion of soot.

237 It is worthy paying attention to the lower NO₂ levels detected during the
238 Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂-catalysed soot combustion experiments in all the gas
239 mixtures tested (see Figures 3a, 3b and 3c), except in NO_x/O₂/SO₂/N₂. These
240 NO₂ levels are lower than those obtained with Ce_{0.73}Zr_{0.27}O₂, regardless the
241 NO₂ production capacity measured in the absence of soot (Figure 2). This
242 means that NO₂ is used more efficiently during the Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂-
243 catalysed soot combustion than when Ce_{0.73}Zr_{0.27}O₂ catalyzes the reaction,
244 which is also related to the important role of the “active oxygen” mechanism.
245 Once the active oxygen of Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ reacts with soot, the remaining
246 oxidizing gases in the mixture, mainly NO₂ and O₂, react more easily, and this
247 produces a synergetic effect between the high oxidation capacities of NO₂ and
248 active oxygen. This would explain why NO₂ is used more efficiently by
249 Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂.

250 Finally, soot combustion experiments were performed in a gas mixture
251 that simulates a real diesel engine exhaust, including NO_x, O₂, H₂O, CO₂ and
252 N₂ (Figure 5 and Table 1). SO₂ has not been included in this gas mixture
253 because the poisoning effect of this gas inhibits almost completely the activity of
254 the catalysts studied (see Figure 1d), and therefore, it is assumed that the ceria-
255 based catalysts can be only used for catalytic combustion of soot on engines
256 running with sulfur-free diesel fuel. These additional soot combustion
257 experiments have been performed with Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂, which is the most
258 active ceria catalyst tested in this study, and without catalyst. The maximum
259 temperature achieved is 530 °C, which is about the maximum temperature that

260 can be reached in a real diesel exhaust using high engine loading and fitting the
261 soot filter just at the exit of the engine [24]. As it was previously mentioned,
262 higher temperatures can be reached in hot spots inside the filters during
263 regeneration, but simulating these transient conditions is out the scope of this
264 study. The only precaution taken in the current study was to calcine catalysts at
265 800 °C in order to provide thermal stability and practical interest to the oxides
266 used.

267 Both the uncatalyzed and catalyzed reactions started at 450 °C in this
268 complex gas stream. However, the uncatalyzed reaction stopped after 23 %
269 soot combustion, while not the catalyzed combustion. This can be explained by
270 the fact that the structure of the model soot used (and also of typical real soots)
271 consists of a turbulent core surrounded by spherical graphitic carbon layers
272 forming an onion-like structure [25-30]. The surface of the soot particles is more
273 reactive than the more internal well-ordered graphitic layers due to the presence
274 of adsorbed hydrocarbons (the volatile matter of the model soot used in this
275 study amounts to 8%). The O₂ molecule, which is the main responsible of the
276 uncatalyzed combustion, is only able to oxidize part of the soot sample and the
277 reaction stops once the most reactive part of the soot particles is consumed. On
278 the contrary, both NO₂ and catalyst active oxygen assist O₂ during the
279 Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂-catalysed soot combustion, and the higher reactivity of
280 these species allows further soot combustion. Also, the maximum soot
281 combustion rate achieved during the catalyzed reaction is about three times
282 higher than that of the uncatalyzed combustion, and Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ also
283 improves the CO₂ selectivity (see data on Table 1). The improved CO₂

284 selectivity can be attributed either to the selective formation of this gas as
285 primary gas product or to the catalytic oxidation of the CO first evolved.

286

287 3.2. *In situ* DRIFTS experiments.

288

289 In order to analyze the reasons of the partial or total inhibiting effect of
290 H₂O, CO₂ and SO₂, *in situ* DRIFTS experiments have been performed at 460
291 °C. The nature of the species adsorbed on the catalysts' surface has been
292 analyzed, and due to this reason, these experiments have been carried out
293 without soot (only with the catalysts). Representative spectra obtained with both
294 catalysts under the different gas mixtures evaluated are included in the Figure
295 2SP (supplementary material), covering the entire range of wavenumbers
296 measured (4000-500 cm⁻¹). Most relevant bands for the purpose of this study
297 appear in the 1800-1000 cm⁻¹ range, and the further analysis and discussion of
298 the spectra is focused on this range of wavenumbers.

299 Figure 6 shows the spectra obtained under the NO_x/O₂/N₂ atmosphere
300 with Ce_{0.73}Zr_{0.27}O₂ (Figure 6a) and Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ (Figure 6b) in the 1800-
301 1000 cm⁻¹ range, where bands of nitrogen-containing surface groups appear.
302 These spectra (and some others) were included and discussed in detail in a
303 previous study, and a complete interpretation was there reported [17]. However,
304 it seems convenient to include also a brief discussion of the behavior of these
305 catalysts in the NO_x/O₂/N₂ atmosphere in this article for further comparison with
306 more complex gas mixtures. Two intense adsorption bands that grow with time
307 are observed on Ce_{0.73}Zr_{0.27}O₂ (Figure 6a) and Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ (Figure 6b)

308 spectra. These bands are located in the 1650-1450 cm^{-1} and 1350-1150 cm^{-1}
309 ranges, and are consistent with the presence of nitrates and nitrites + nitrates,
310 respectively [31]. Several well-defined peaks appear at particular wavenumbers
311 within the two main bands of the $\text{Ce}_{0.73}\text{Zr}_{0.27}\text{O}_2$ and $\text{Ce}_{0.64}\text{Zr}_{0.27}\text{Nd}_{0.09}\text{O}_2$ spectra,
312 and this is attributed to the presence of nitrites and/or nitrates with different
313 configuration (monodentate, bidentate and/or bridged) adsorbed on different
314 surface sites of the heterogeneous surface of these catalysts. In both figures
315 (Figures 6a and 6b), the band which is unambiguously assigned to nitrate
316 species (1650-1450 cm^{-1} range) is the most intense, and the relative intensity of
317 the band at 1350-1150 cm^{-1} with regard to that at 1650-1450 cm^{-1} can be
318 related with the presence of nitrites. The relative intensity of the band at 1350-
319 1150 cm^{-1} is higher for the $\text{Ce}_{0.64}\text{Zr}_{0.27}\text{Nd}_{0.09}\text{O}_2$ catalyst (Figure 6b) than for
320 $\text{Ce}_{0.73}\text{Zr}_{0.27}\text{O}_2$ (Figure 6a), suggesting that Nd^{3+} doping favors the formation of
321 nitrites. Nitrites are less oxidized species than nitrates, and its formation on the
322 Nd^{3+} -containing oxide is consistent with the creation of oxygen vacancies due to
323 the substitution of tetravalent Ce^{4+} cations by trivalent Nd^{3+} cations. Also, nitrites
324 have inferior thermostability than nitrates and decompose more easily [32], and
325 this would explain why Nd^{3+} doping slightly improves the NO_2 production
326 capacity of the Ce-Zr mixed oxide in the $\text{NO}_x/\text{O}_2/\text{N}_2$ atmosphere (see Figure
327 2a). Nd^{3+} promotes the formation of less stable surface nitrogen species and
328 provides an alternative and faster NO_2 production pathway.

329

330 *3.2.1. CO₂ effect.*

331

332 The bands attributed to nitrates ($1650\text{-}1450\text{ cm}^{-1}$) and nitrites + nitrates
333 ($1350\text{-}1150\text{ cm}^{-1}$) [31] are also observed in the spectra obtained with the gas
334 mixture $\text{NO}_x/\text{O}_2/\text{CO}_2/\text{N}_2$, both for $\text{Ce}_{0.73}\text{Zr}_{0.27}\text{O}_2$ (Figure 7a) and
335 $\text{Ce}_{0.64}\text{Zr}_{0.27}\text{Nd}_{0.09}\text{O}_2$ (Figure 7b). However, an additional signal appears around
336 1400 cm^{-1} , between both bands assigned to nitrogen compounds.

337 Various carbonate-type species have been reported to present
338 adsorption modes in the range $1200\text{-}1600\text{ cm}^{-1}$. A band at ca. 1478 cm^{-1} has
339 been proposed to correspond to the antisymmetric stretching of the terminal C-
340 O bonds in poly or monodentate carbonates, and the corresponding symmetric
341 mode would appear at ca. 1350 cm^{-1} [33, 34]. Bands of hydrogen carbonate (at
342 1214 , $1399\text{-}1410$ and ca. 1600 cm^{-1}) and bidentate carbonate (bands at ca.
343 1583 and 1297 cm^{-1}) would appear in this range of wavenumbers as well [33-
344 35].

345 The spectra obtained including CO_2 in the gas mixture (Figure 7)
346 evidence the formation of carbonate species on both catalysts, but a detailed
347 identification of the type of carbonate specie formed is not possible because
348 some of the bands overlap with those of the nitrogen surface groups. However,
349 it seems reasonable to rule out the formation of hydrogen carbonates, because
350 this would require the participation of hydroxyl groups (or water). Hydroxyl
351 groups may be detected by DRIFTS bands in the $4000\text{-}3000\text{ cm}^{-1}$ range, and
352 their potential consumption upon CO_2 chemisorption and hydrogen carbonates
353 formation may be evidenced by negative bands in this range of wavenumbers.
354 These evidences have not been observed in our spectra (see Figure 2SM of the
355 supplementary material). The absence of hydroxyl groups on the catalysts
356 studied must be attributed to the high calcination temperature used ($800\text{ }^\circ\text{C}$). It

357 is therefore presumed the formation of carbonate species on both catalysts
358 during the *in situ* DRIFTS experiments performed under NO_x/O₂/CO₂/N₂.

359 The chemisorption of CO₂ on the catalysts modifies the nature of the
360 nitrogen surface species, mainly affecting the formation of nitrites on
361 Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ (Figure 7b). For this catalyst, the ratio between the DRIFT
362 signal at 1547 cm⁻¹, which is attributed to nitrates, and that at 1242 cm⁻¹, which
363 is assigned to nitrites + nitrates, after 60' on the gas streams increases from 2.0
364 for NO_x/O₂/N₂ to 2.5 for NO_x/O₂/CO₂/N₂. This indicates that CO₂ chemisorption
365 hinders the formation of nitrites on Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂, and nitrites were
366 proposed to be reaction intermediates of the fast catalytic oxidation of NO to
367 NO₂ on Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂. This would explain the important decrease of the
368 NO₂ formation under NO_x/O₂/CO₂/N₂ with regard to NO_x/O₂/N₂ for experiments
369 performed with Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ (compare Figures 2a and 2b). However, the
370 effect of CO₂ on the nature of the nitrogen surface species is not so relevant for
371 the Ce_{0.73}Zr_{0.27}O₂ catalyst (compare Figures 6a and 7a), and neither is on the
372 Ce_{0.73}Zr_{0.27}O₂-catalysed oxidation of NO to NO₂ (compare Figures 2a and 2b).

373 In conclusion, CO₂ and NO_x compete for the adsorption sites on the
374 catalysts' surface, and mainly for the Nd³⁺-containing catalyst, this affects the
375 catalytic oxidation of NO to NO₂. However, the soot combustion experiments
376 (Figure 1b) evidence that, although the Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂-catalysed NO
377 oxidation is impeded, the catalytic combustion of soot remains faster than with
378 Ce_{0.73}Zr_{0.27}O₂. This supports the hypothesis of the important contribution of the
379 active oxygen mechanism during the Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂-catalysed soot
380 combustion, which seems to remain relevant even after the partial inhibition of
381 the NO₂-assisted soot combustion mechanism by CO₂ chemisorption.

382

383 *3.2.2. H₂O effect.*

384

385 The DRIFT spectra obtained under the NO_x/O₂/H₂O/N₂ gas stream with
386 the Ce_{0.73}Zr_{0.27}O₂ and Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ catalysts are compiled in Figures 8a
387 and 8b, respectively. The bands attributed to nitrates (1650-1450 cm⁻¹) and
388 nitrites + nitrates (1350-1150 cm⁻¹) [31] are also observed in the spectra
389 obtained with this gas mixture. The presence of water in the gas mixture delays
390 the formation of the nitrogen surface groups (compare Figures 6 (NO_x/O₂/N₂)
391 and 8 (NO_x/O₂/H₂O/N₂)), and this delay affects both catalysts. For instance,
392 bands of significant intensity appear after 1 minute under reaction conditions in
393 the absence of H₂O (Figure 6), while not in the presence of H₂O (Figure 8). The
394 reason of this delay is expected to be the competition between H₂O and NO_x
395 for the adsorption sites on the catalysts' surface. H₂O chemisorption should
396 form hydroxyl groups, but evidences of this type of groups are not observed in
397 the 4000-3000 cm⁻¹ region (see Figure 2SM in supplementary material). This is
398 because hydroxyl groups are suitable sites for NO_x chemisorption, as
399 previously observed [32], and it seems they are depleted by NO_x chemisorption
400 as soon as they are formed. According to this interpretation H₂O would force the
401 catalytic oxidation of NO to NO₂ to progress through a different and slower
402 reaction pathway than that occurring in the absence of H₂O. This would explain
403 why the presence of H₂O in the gas mixture diminishes the catalytic oxidation of
404 NO to NO₂ (compare Figures 2a (NO_x/O₂/N₂) and 2c (NO_x/O₂/H₂O/N₂)) and the
405 catalytic combustion of soot (compare Figures 1a (NO_x/O₂/N₂) and 1c
406 (NO_x/O₂/H₂O/N₂)), but not inhibits completely the reactions as SO₂ almost does.

407 An additional argument to support this interpretation is obtained by the
408 detailed analysis of the spectra in the 1650-1450 cm^{-1} range (Figure 8), which is
409 attributed to nitrates. Nitrates can be adsorbed on a solid surface in different
410 configurations of different stability, and the actual position of the adsorption
411 bands of each type of nitrate is different. These configurations are, in increasing
412 order of stability: monodentate nitrates, bidentate nitrates and bridging nitrates,
413 and it has been reported they present adsorption bands in the 1530–1480 cm^{-1} ,
414 1565–1500 cm^{-1} and 1650–1600 cm^{-1} ranges, respectively [31]. As deduced
415 from this assignation, the higher the stability of a surface nitrate specie, the
416 higher the absorption band wavenumber in this interval.

417 Different contributions to the bands in the 1650-1450 cm^{-1} range are
418 observed on Figures 6 ($\text{NO}_x/\text{O}_2/\text{N}_2$) and 8 ($\text{NO}_x/\text{O}_2/\text{H}_2\text{O}/\text{N}_2$) due to the
419 presence of different surface nitrate species. For instance, the main band in this
420 region obtained with the $\text{Ce}_{0.73}\text{Zr}_{0.27}\text{O}_2$ catalyst under $\text{NO}_x/\text{O}_2/\text{N}_2$ appears at
421 1527 cm^{-1} (see Figure 6a), and therefore, it can be attributed to monodentate or
422 bidentate nitrates. In addition, some other contributions appear with lower
423 intensity at higher wavenumbers (1547 and 1559 cm^{-1}), which can be assigned
424 to bidentate and/or bridging nitrates. Bands at quite similar wavenumbers are
425 also observed in the presence or H_2O with the $\text{Ce}_{0.73}\text{Zr}_{0.27}\text{O}_2$ catalyst (Figure 8a;
426 $\text{NO}_x/\text{O}_2/\text{H}_2\text{O}/\text{N}_2$), but the relative intensity of the most stable species (bidentate
427 and bridging nitrates) is higher, and a new contribution assigned to bridging
428 nitrates appears at 1594 cm^{-1} . This supports that the presence of H_2O favors
429 the formation of more stable nitrate species with regard to those formed in its
430 absence, and this applies to the two catalysts studied.

431 In conclusion, the presence of water in the gas mixture hinders the
432 catalytic oxidation of NO to NO₂, and therefore the catalytic combustion of soot,
433 because it delays the formation of reaction intermediates on the catalyst surface
434 and favors the formation of more stable nitrogen surface groups than in a H₂O-
435 free gas stream.

436

437 3.2.3. SO₂ effect.

438

439 Finally, the effect of SO₂ has also been studied by DRIFTS, and the
440 spectra are included on Figure 9 for Ce_{0.73}Zr_{0.27}O₂ (Figure 9a) and
441 Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ (Figure 9b).

442 Important differences have been noticed in these experiments performed
443 under NO_x/O₂/SO₂/N₂ with regard to the previously studied gas mixtures. In the
444 presence of SO₂, the nitrates bands at 1650-1450 cm⁻¹ grow once the gas
445 mixture is put in contact with the catalysts, but they decrease afterwards. Also
446 the nitrites + nitrates band (1350-1150 cm⁻¹) of the Ce_{0.73}Zr_{0.27}O₂ catalyst
447 (Figure 9a) grows with time, but much more slowly than in the absence of SO₂
448 (compare with Figure 6a). In addition to these changes in the behavior of the
449 nitrogen surface groups, a prominent band grows at 1348 cm⁻¹ on the
450 Ce_{0.73}Zr_{0.27}O₂ catalyst (Figure 9a), which is progressively shifted to 1376 cm⁻¹.
451 This band appears at 1345 cm⁻¹ on the Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ catalyst (Figure 9b),
452 but with much lower intensity. This band has been assigned to a S=O vibration
453 mode of surface sulfates or adsorbed SO₃ [36-38]. A new band at 1293 cm⁻¹
454 grows on the Ce_{0.73}Zr_{0.27}O₂ catalyst (Figure 9a) as well, which has been also
455 assigned to sulfate species [36].

456 These experiments evidence that SO₂ and NO_x compete by the surface
457 adsorption sites, and sulfates formation significantly inhibits the catalytic
458 oxidation of NO to NO₂ (compare Figures 2a (NO_x/O₂/N₂) and 2d
459 (NO_x/O₂/SO₂/N₂)), and at the end, the catalytic combustion of soot (compare
460 Figures 1a (NO_x/O₂/N₂) and 1d (NO_x/O₂/SO₂/N₂)).

461 According to the important differences observed in the DRIFT spectra
462 obtained in the presence of SO₂ for both catalysts (see Figure 9), sulfate
463 formation seems to affect much more to Ce_{0.73}Zr_{0.27}O₂ than to
464 Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂, but the effect on the catalytic oxidation of NO to NO₂ is
465 stronger for Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂ (see Figure 2d). This suggest that the catalytic
466 oxidation of NO to NO₂ is not only related with the nitrogen surface groups
467 observed by DRIFTS but active oxygen, which cannot be observed so far,
468 seems to be also involved in the catalytic oxidation of NO and seems to be also
469 affected by SO₂ chemisorption. This would explain why the Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂
470 catalyst, whose catalytic activity not only depends on NO₂ formation but also on
471 active oxygen production (see Figure 4 and the discussion on the text), suffer
472 the same SO₂ inhibiting effect than Ce_{0.73}Zr_{0.27}O₂ in spite of the important
473 differences on the nitrogen surface groups behavior observed by DRIFTS (see
474 Figure 9).

475

476 **4.- Conclusions.**

477

478 The effect of CO₂, H₂O and SO₂ in the Ce_{0.73}Zr_{0.27}O₂ and
479 Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂-catalyzed combustion of soot in the presence of NO_x and
480 O₂ has been studied, and the main conclusions achieved can be summarized
481 as follows:

482

483 • The presence of CO₂, H₂O or SO₂, in addition to NO_x and O₂, in the gas
484 mixture decreases the activity for soot combustion of both catalysts with
485 regard to the combustion in a NO_x/O₂/N₂ mixture. The inhibiting effect
486 follows the trend SO₂ > H₂O > CO₂.

487

488 • CO₂ partially inhibits the catalytic activity because competes with NO_x for
489 the adsorption sites on the catalysts' surface, and mainly for the Nd³⁺-
490 containing catalyst, this affects the catalytic oxidation of NO to NO₂.
491 However, the Ce_{0.64}Zr_{0.27}Nd_{0.09}O₂-catalysed combustion of soot remains
492 faster, and this is attributed to the contribution of the active oxygen
493 mechanism.

494

495 • The presence of H₂O in the gas mixture hinders the catalytic oxidation of
496 NO to NO₂ on both catalysts, and therefore the catalytic combustion of
497 soot, because it delays the formation of reaction intermediates on the
498 catalyst surface and favors the formation of more stable nitrogen surface
499 groups than in a H₂O -free gas stream.

500

501 • SO₂ and NO_x also compete by the surface adsorption sites on the
502 catalysts, and sulfates formation significantly inhibits the catalytic
503 oxidation of NO to NO₂, and at the end, the catalytic combustion of soot.
504 The high stability of sulfates explains the strongest inhibiting effect of
505 SO₂.

506

507 • Regardless the gas mixture used the catalytic activity for soot
508 combustion of $\text{Ce}_{0.64}\text{Zr}_{0.27}\text{Nd}_{0.09}\text{O}_2$ is equal or higher to that of
509 $\text{Ce}_{0.73}\text{Zr}_{0.27}\text{O}_2$, and it is proposed that this is because Nd^{3+} doping
510 promotes the participation of the active oxygen mechanism together with
511 the NO_2 -assisted mechanism in the catalytic combustion of soot.

512
513 • The maximum soot combustion rate achieved during a
514 $\text{Ce}_{0.64}\text{Zr}_{0.27}\text{Nd}_{0.09}\text{O}_2$ -catalyzed reaction in $\text{NO}_x/\text{O}_2/\text{CO}_2/\text{H}_2\text{O}/\text{N}_2$ is about
515 three times higher than that of the uncatalyzed combustion, and this
516 catalyst also improves the CO_2 selectivity.

517

518

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Table 1. Results of the soot combustion experiments in a NO_x/O₂/H₂O/CO₂/N₂ atmosphere (curves on Figure 5).

Catalyst	Burn off (%)	CO (%) = 100·CO/CO_x	Maximum rate (µgC/s)
None	23	60	4
Ce _{0.64} Zr _{0.27} Nd _{0.09} O ₂	73	19	10.5

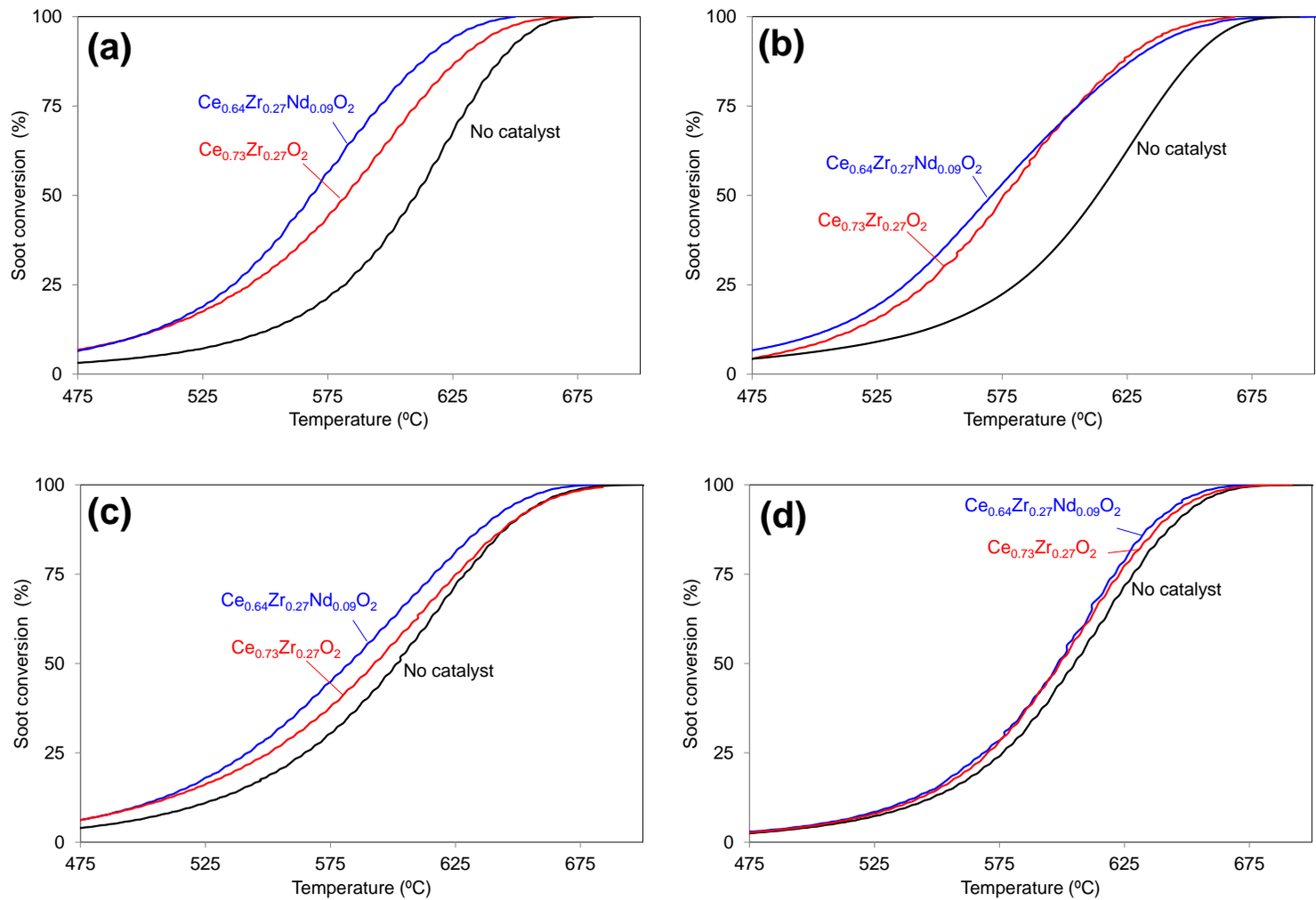


Figure 1. Soot combustion experiments performed under different gas mixtures: (a) $\text{NO}_x/\text{O}_2/\text{N}_2$, (b) $\text{NO}_x/\text{O}_2/\text{CO}_2/\text{N}_2$, (c) $\text{NO}_x/\text{O}_2/\text{H}_2\text{O}/\text{N}_2$, (d) $\text{NO}_x/\text{O}_2/\text{SO}_2/\text{N}_2$.

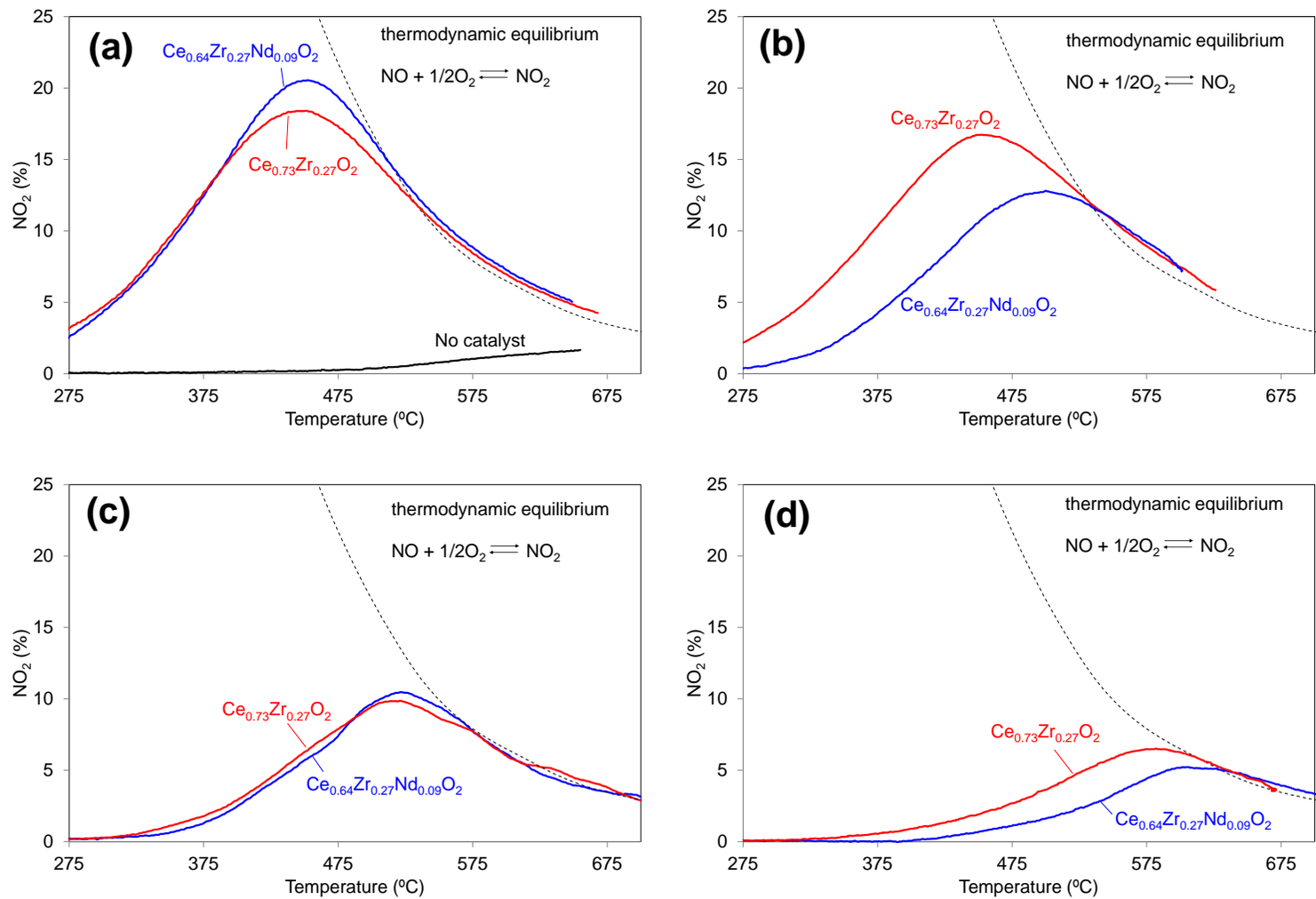


Figure 2. NO₂ formation in oxidation experiments performed without soot under different gas mixtures: (a) NOx/O₂/N₂, (b) NOx/O₂/CO₂/N₂, (c) NOx/O₂/H₂O/N₂, (d) NOx/O₂/SO₂/N₂.

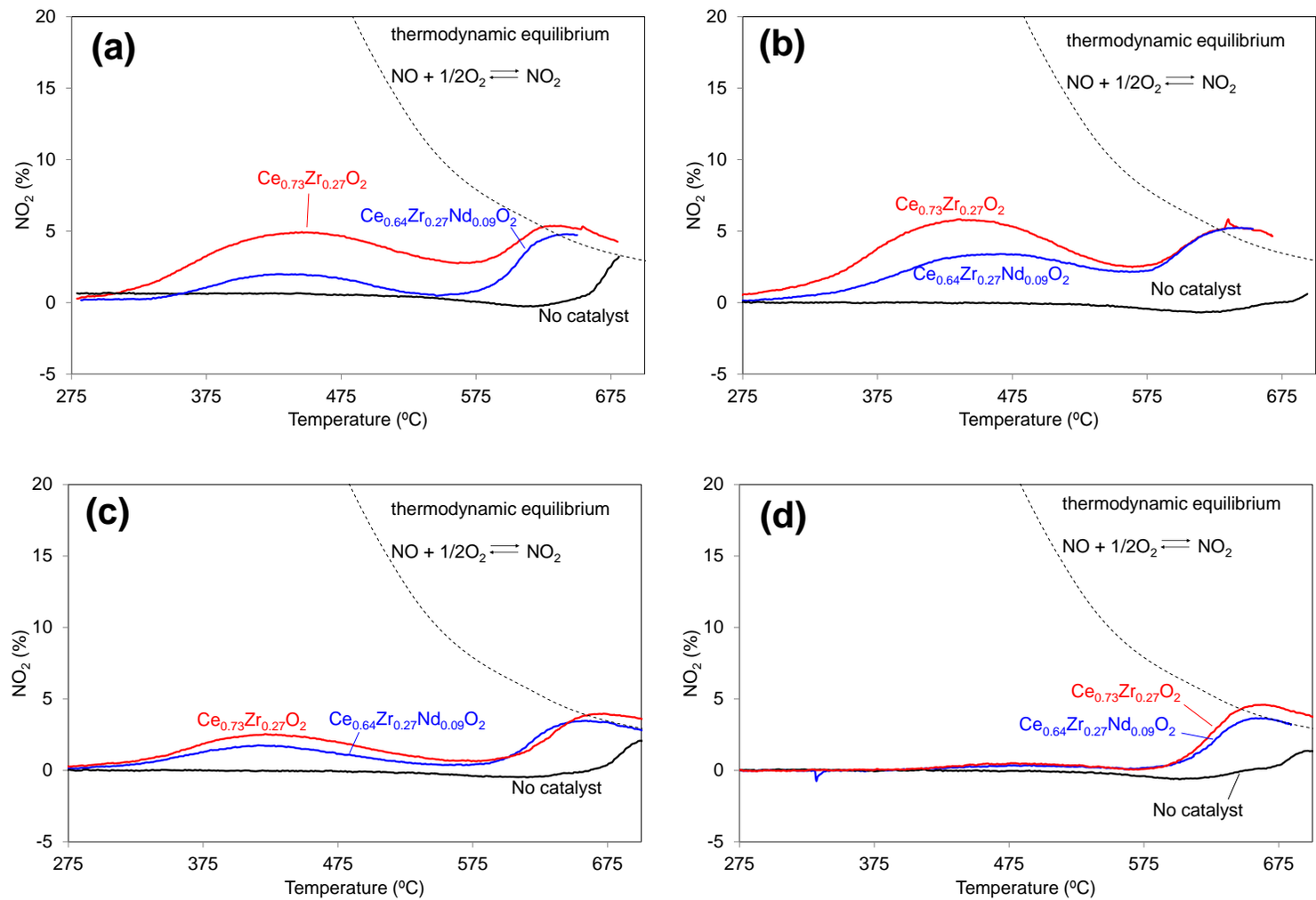


Figure 3. NO₂ formation in oxidation experiments performed with soot under different gas mixtures: (a) NOx/O₂/N₂, (b) NOx/O₂/CO₂/N₂, (c) NOx/O₂/H₂O/N₂, (d) NOx/O₂/SO₂/N₂. (These curves were obtained in the same experiments than soot conversion curves on Figure 1)

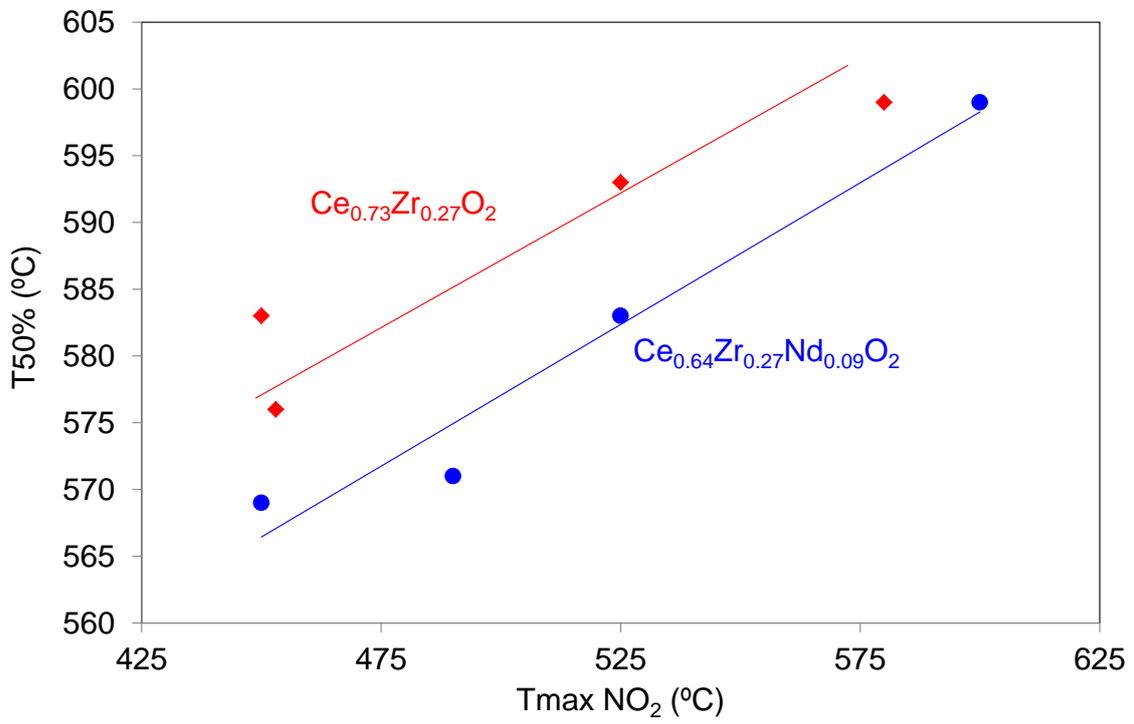


Figure 4. Relationship between soot combustion and NO₂ production. T50% (°C): Temperature for 50% soot combustion; Tmax NO₂ (°C): Temperature of maximum NO₂ production in blank experiments.

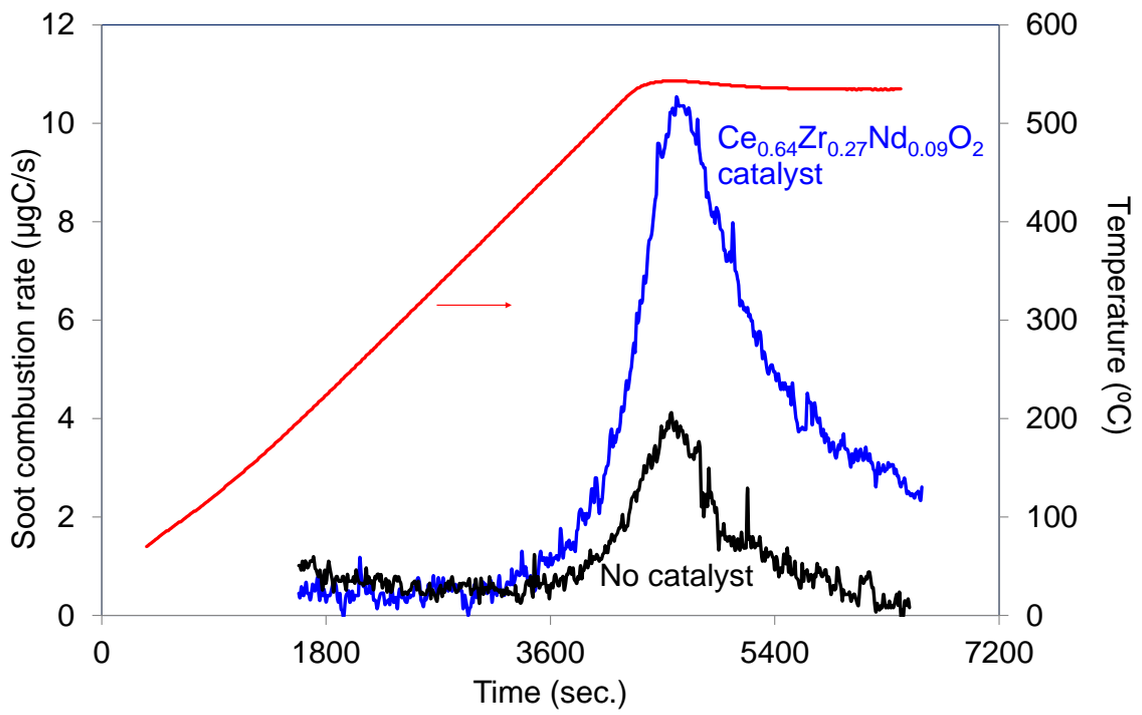


Figure 5. Soot combustion experiments in $\text{NO}_x/\text{O}_2/\text{H}_2\text{O}/\text{CO}_2/\text{N}_2$.

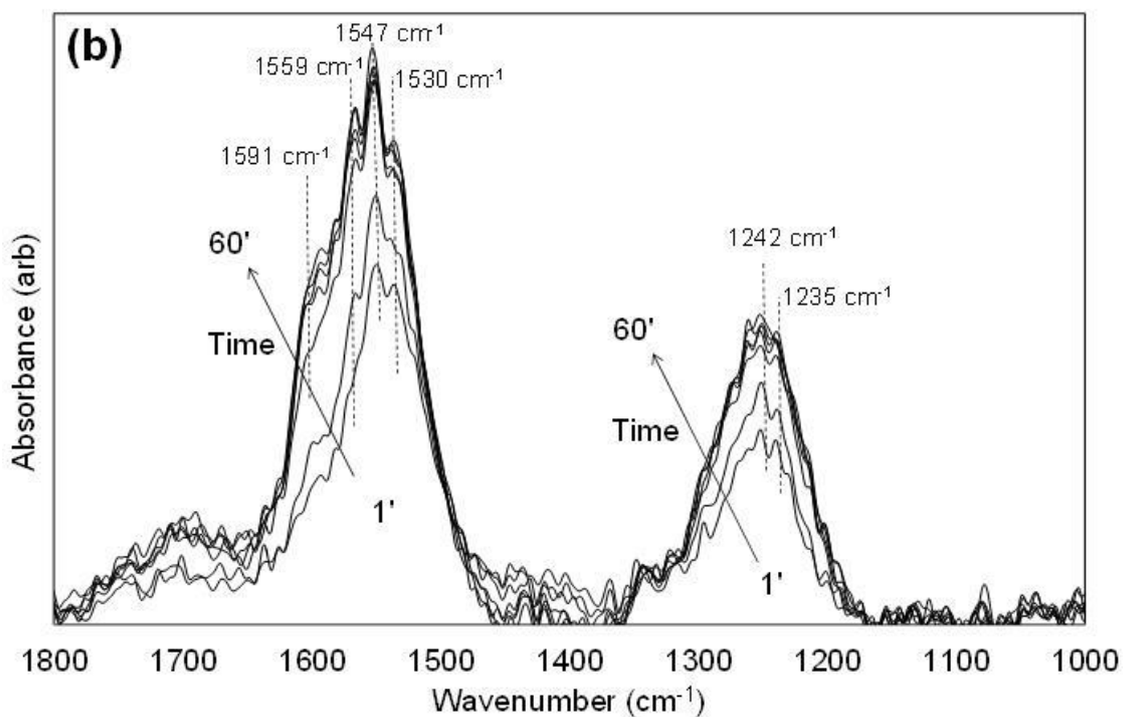
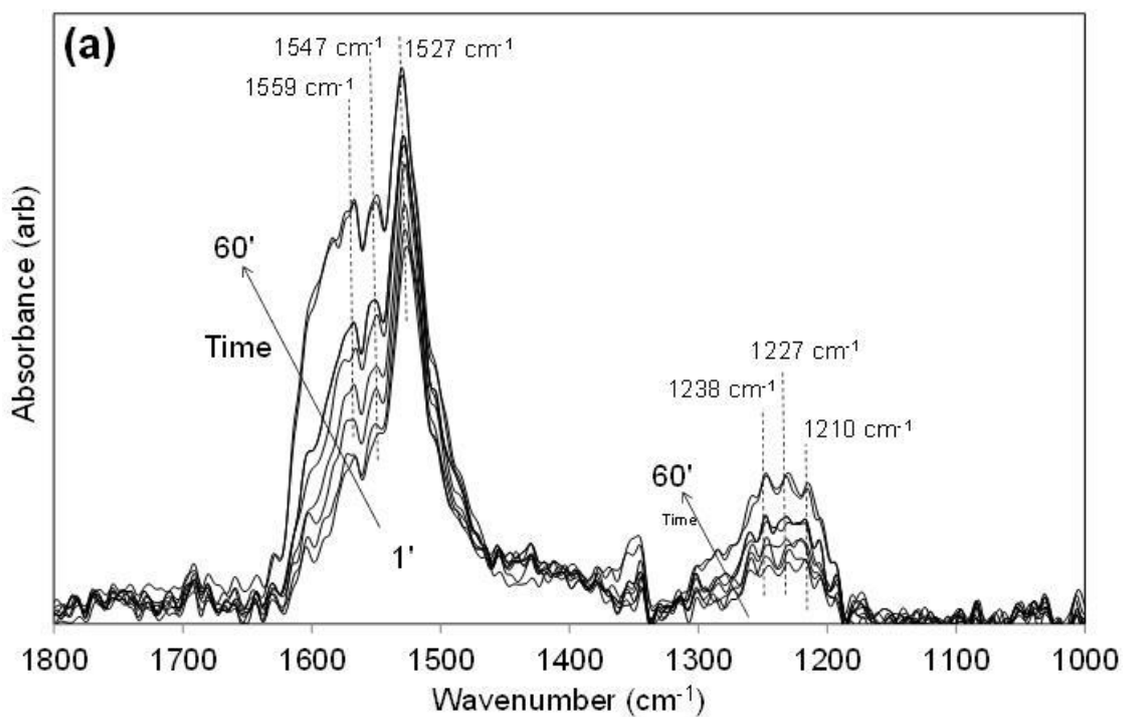


Figure 6. DRIFT spectra obtained in a $\text{NO}_x/\text{O}_2/\text{N}_2$ atmosphere with (a) $\text{Ce}_{0.73}\text{Zr}_{0.27}\text{O}_2$ and (b) $\text{Ce}_{0.64}\text{Zr}_{0.27}\text{Nd}_{0.09}\text{O}_2$. (These Figures have been reprinted from [17] for the sake of clarity).

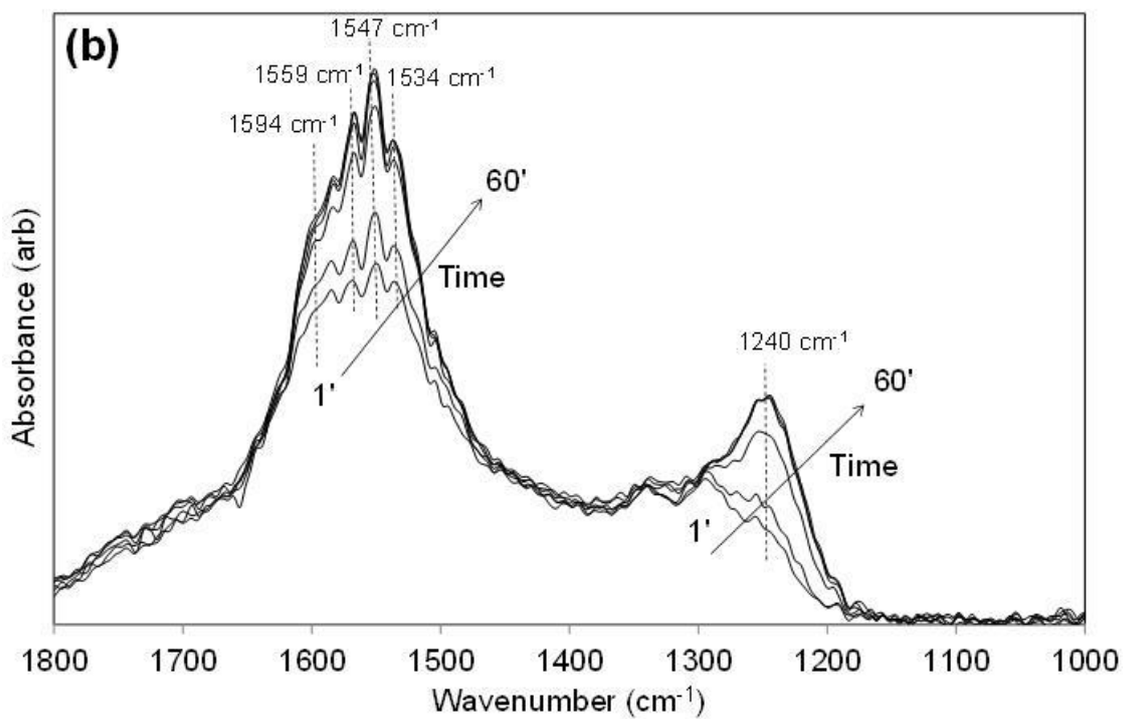
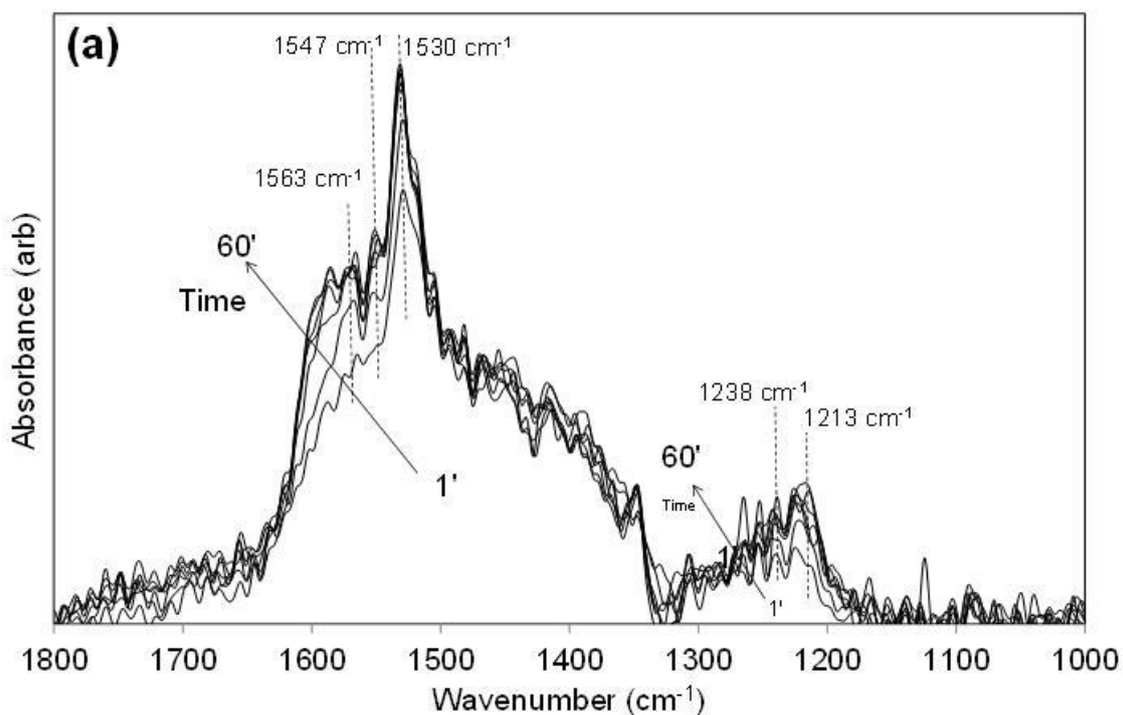


Figure 7. DRIFT spectra obtained in a $\text{NO}_x/\text{O}_2/\text{CO}_2/\text{N}_2$ atmosphere with (a) $\text{Ce}_{0.73}\text{Zr}_{0.27}\text{O}_2$ and (b) $\text{Ce}_{0.64}\text{Zr}_{0.27}\text{Nd}_{0.09}\text{O}_2$.

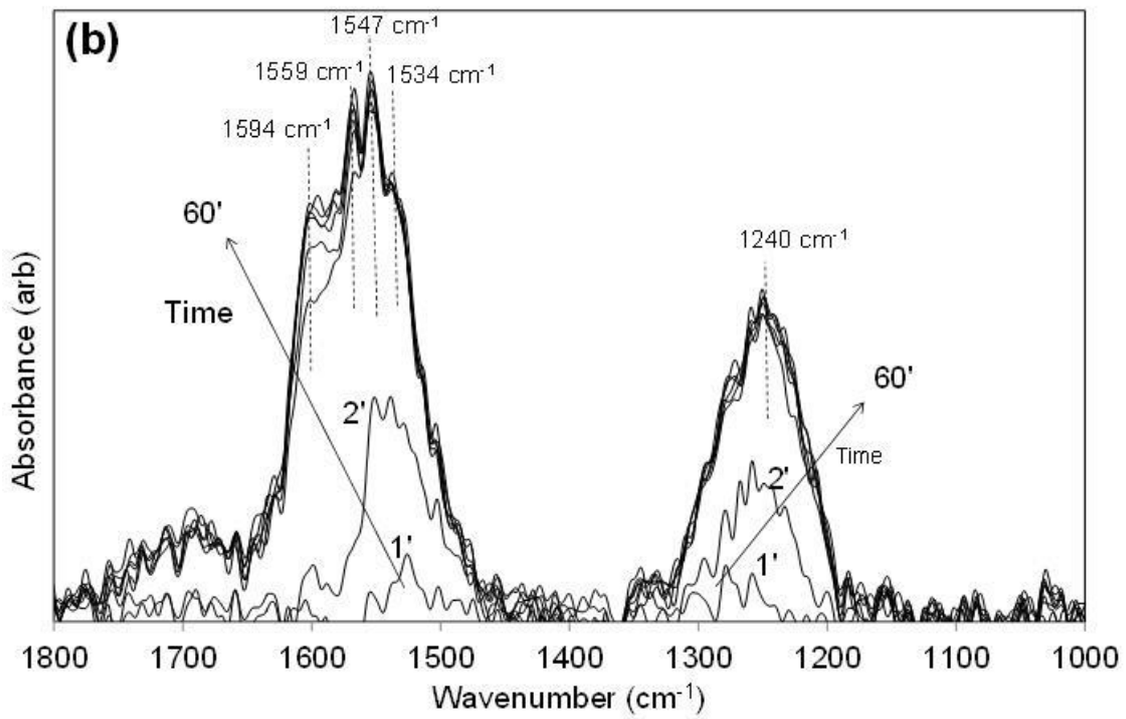
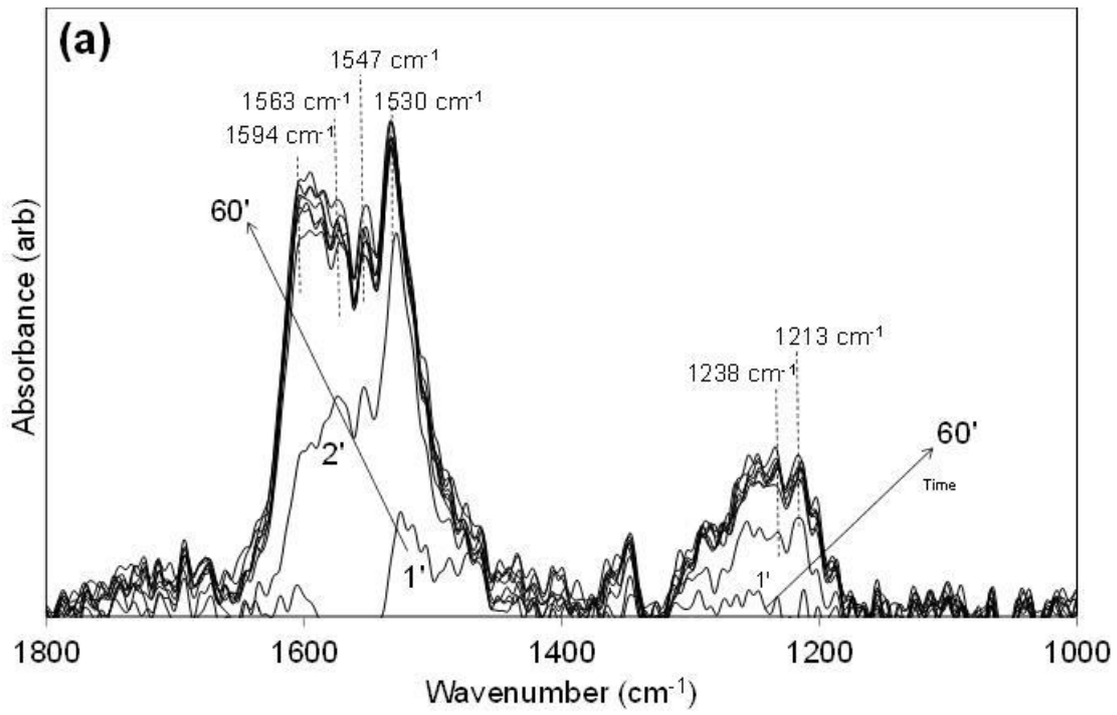


Figure 8. DRIFT spectra obtained in a $\text{NO}_x/\text{O}_2/\text{H}_2\text{O}/\text{N}_2$ atmosphere with (a) $\text{Ce}_{0.73}\text{Zr}_{0.27}\text{O}_2$ and (b) $\text{Ce}_{0.64}\text{Zr}_{0.27}\text{Nd}_{0.09}\text{O}_2$.

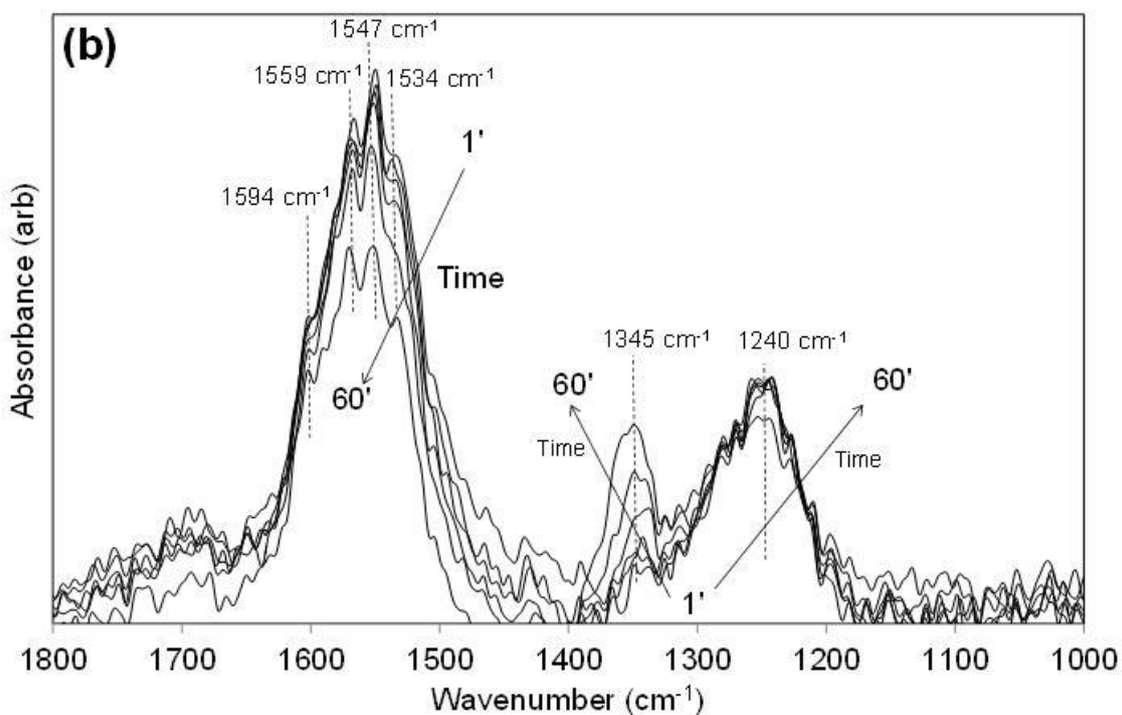
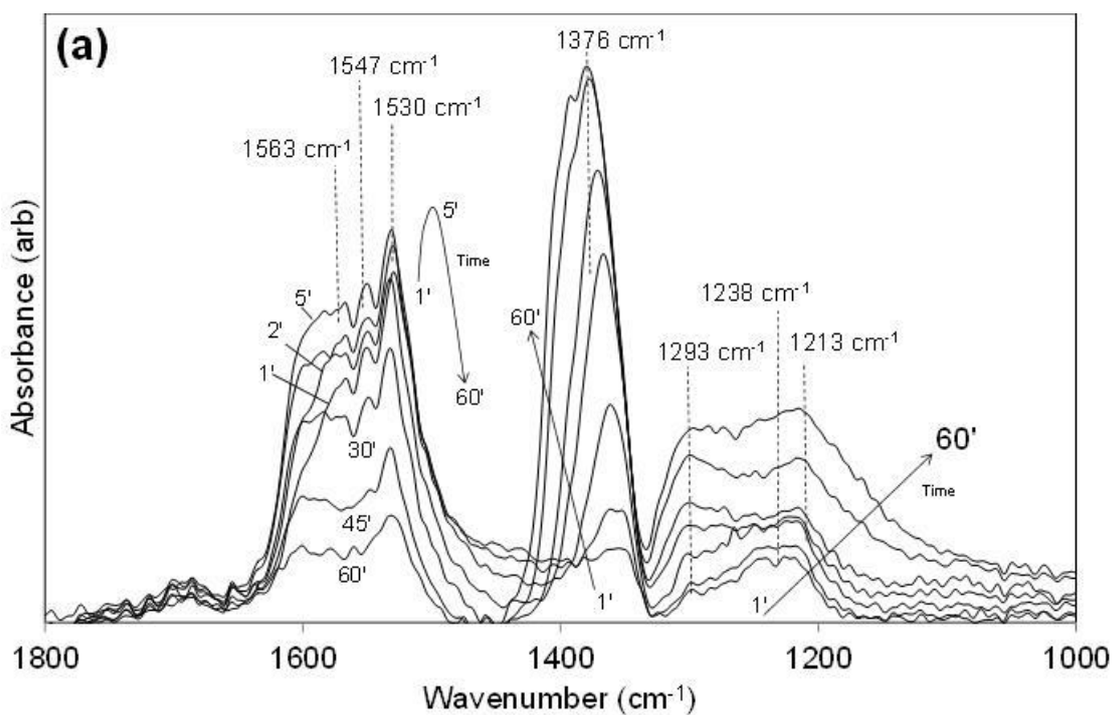


Figure 9. DRIFT spectra obtained in a $\text{NO}_x/\text{O}_2/\text{SO}_2/\text{N}_2$ atmosphere with (a) $\text{Ce}_{0.73}\text{Zr}_{0.27}\text{O}_2$ and (b) $\text{Ce}_{0.64}\text{Zr}_{0.27}\text{Nd}_{0.09}\text{O}_2$.