The negative effect of ceria on the propene selectivity for isopropanol decomposition over phosphated and phosphate-free ceria/alumina catalysts

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Abstract
Consequences of loading level of ceria (5, 10 and 20-wt%) on the structural, textural and surface acid–base properties as well as the impacts on catalytic activity towards isopropanol decomposition on pure and phosphated alumina is described. The obtained results indicated that composites mainly kept γ–alumina structure and the addition of ceria and/or phosphate has no effect on the bulk structure of alumina. Textural characteristics show that the phosphated composites have higher surface area than phosphate free samples. Pyridine adsorption as followed by FTIR indicated the presence of Lewis acid sites with different strengths. Moreover, results of gas phase decomposition of isopropanol show a systematic inhibition in propene selectivity based on ceria ratio.

Keywords: Texture; Mesoporous; Acidity; Activity; Decomposition

Introduction
Active alumina, as one of the most common catalyst support, has a good performance in catalysis applications due to its physical and chemical properties (Levy et al. 1968). Alumina has various properties such as high specific surface area, surface acidity and defects in their crystalline structure that are the key factor in the wide range of applications. One of the most important factors in alumina utilization is thermal stability, which is a very important issue for processes running at high temperature (Chen et al. 2001). Therefore, the addition of some additives can improve the thermal stability of alumina catalysts, hold the surface area and improve the surface acidity (Shinjoh 2006; Ozawa 2006; Ozawa & Nishio 2004; Ersoy & Gunay 2004). Phosphates have been claimed to act as a support stabilizer (Gishti et al. 1984; Abbattista et al. 1990; Khalaf et al. 2007). Gishti et al. (1984) evidenced a stable effect of phosphate by studying the phase transitions of alumina in the range 673–1323 K. However, the surface area decreased with the increasing content of phosphate in alumina, even less than the blank sample after calcinated at 1073 K. Khalaf et al. 2007 also proved no surface area stabilization effect from phosphates in the case of the transition phase (spinel) aluminas.

Ceria possesses versatile acid–base properties, depending on the nature and temperature of the pretreatment. It may have high number of basic sites of weak or medium strength (Binet et al. 1999). Martin and Duprez found the following scale for the density of basic sites of some metal oxides CeO$_2$ > MgO > ZrO$_2$ > Al$_2$O$_3$ > SiO$_2$ (Martin & Duprez 1997). Binet et al. (1999) also observed that ceria can chemisorb CO or pyridine, but the band positions strongly suggest that the Lewis acidity of ceria is significantly lower than that of zirconia or titania. In contrast to Lewis basicity, the Lewis acidity would decrease upon reduction of ceria.

Ceria and ceria-based materials has been widely employed as automotive three-way catalysts (TWC) for reducing the exhaust pollutants, fuel cell processes, oxygen permeation membrane systems, exhaust combustion catalysts, and catalytic wet oxidation (Yao & Yao 1984; Dos Santos et al. 2008; Bera et al. 2003; Khalil 2007). Cerium oxide can maintain large amounts of oxygen through the easy transition between Ce(IV) and Ce(III). This property is very useful in catalysis especially in oxidation reactions
Nevertheless, pure ceria is poorly thermostable and undergoes rapid sintering under high temperature conditions, which leads to loss of oxygen buffer capacity and deactivation of the catalysts. The combination of two metals in an oxide can lead to novel structural and electronic properties of the final oxide, consequently modifying its catalytic activity and selectivity. Ceria-alumina mixed oxides are widely used as catalytic materials (Walter & Oldfield 1989; Niu et al. 1999; Sanchez-Valente et al. 2004).

Isopropanol conversion is widely investigated and lately it is used to characterize acid–base or redox properties of catalysts (Khalaf et al. 2007). The conversion of isopropanol is known to occur through two competitive pathways namely dehydrogenation and dehydration. Ai (1977) has assumed that the dehydration of isopropanol probes acid sites, whereas the dehydrogenation probes acid and base sites functioning in a concerted fashion. Isopropanol usually dehydrates to propene over acidic catalysts and dehydrogenates to acetone over basic catalysts (Scheme 1). The acid–base pairs responsible for acetone formation are strongly activated in the presence of oxygen that indicates that they are related to redox properties of the material (Abdellah et al. 1993).

In the present work, ceria-alumina as well as phosphate alumina modified with ceria have been prepared and characterized by several tools to gain an overview on the surface properties of these catalysts. In addition, the catalytic activities towards isopropanol decomposition for these composites have been explored.

### Results and discussion

#### Thermal analysis

The thermal behaviors of the pure alumina gel and phosphate alumina has been studied in previous paper (Khalaf et al. 2007). Figure 1 (a and b) shows the thermal events of pure ceria, phosphate and phosphate-free ceria-alumina samples. Thermal analysis results for pure ceria (Ce) exhibited the loss of physisorbed water and partial surface dehydroxylation through two steps. The first step, 4.3% mass loss, appeared at 370 K due to the loss of physisorbed water, and the other mass loss step (ca 72.4%) is the main decomposition process that occurred in a narrow temperature region (440 – 570 K). The addition of phosphate by 6 wt% into ceria gel (CeP) does not have a significant change in the amount of mass loss. DTA curve for, Ce and CeP, shows an endothermic mass loss at 340 K, corresponding to the elimination of water absorbed by crystalline ceria. Another endothermic peak around 480 K, correlated to a mass loss, that must be considered as due to the crystallization of the residual amorphous phase.

The thermal events of ceria-alumina, xCeAl, and phosphate ceria-alumina, xCeAlP, samples are shown in Figure 1. It is clear that, the increasing in the ceria loading levels resulted in increasing in the mass loss % (39.3, 50.0 and 54.0% for 5CeAl, 10CeAl and 20CeAl samples, respectively). This is logically, due to ceria losses mass higher than alumina. As for phosphated samples, one can noticed that there is no change in the mass loss % by the comparison between phosphated and phosphate-free samples. Thus, the phosphate ions have slightly effect on the mass loss % of phosphate samples. Only the main difference between xCeAl and xCeAlP is in the wider extended region for phosphated samples.

DTA results for all samples show two endothermic peaks, the first one appeared at low temperature (about 330 K) due to the elimination of adsorbed water, and the other one appeared at temperature higher than 480 K due to the decomposition of samples.

#### X-ray diffraction (XRD)

XRD patterns for tested samples were shown in Figure 2. These diffractograms indicate that the coprecipitation of ceria/alumina with different loading levels followed by calcinations causes a modification of γ-Al₂O₃ structure.

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**Scheme 1** Catalytic decomposition of isopropanol.
Many researchers (Khalaf et al. 2007; López et al. 2010; Marcu et al. 2010) had investigated the crystalline structure of pure ceria and alumina. The results of alumina and phosphate alumina have been investigated in the literature (Khalaf et al. 2007). For ceria, the XRD diffractogram (Figure 2a) displayed sharp and intense peaks corresponding to cubic CeO$_2$ fluorite structure as matched with the database in JCPDS (file number 04-0593) (JCPDS, International Centre for Diffraction Data, PCPDFWIN 1995). For other samples, phosphate and nonphosphated, inspecting the diffractograms and matching with the relevant ASTM standards indicate...
that all samples xCeAl and xCeAlP assume the γ-
structure of alumina as prominent and fluorite structure
for ceria as a little. These results are similar to those pre-
viously reported (Khalil 2007; Silversand et al. 1997;
Johnson 1990). The cubic pattern of ceria (fluorite struc-
ture) was slightly noticed in higher loading level (20%) than
other samples, but a second phase of γ-Al2O3 was
also prominent at all samples. Furthermore, increasing
the loading of ceria from w = 5, 10 to 20%, resulted in
more increase in the degree of crystallization (higher
peak intensities). The introduction of phosphate ions
from ammonium salt into gel causes a slight decrease in
the degree of crystallization of these samples. The crys-
tallite size was estimated from the Scherrer equation
(Klug & Alexander 1970), and the results are cited in
Table 1. The smallest particle size (4.3 nm) between the
present two groups of composites (xCeAl and xCeAlP)
detected for the 5CeAlP composite. For 10CeAlP
and 20CeAlP samples, particle sizes were estimated as
5.8 and 7.0 nm, respectively. Particle sizes observed for
the 5CeAl, 10CeAl and 20CeAl composites were 5.4, 8.1,
and 12.2 nm, respectively, which is a higher than the
value obtained for phosphated samples. Moreover, the
obtained diffractograms for the samples at low levels re-
vealed that fluorite-structured CeO2 crystallites are well
dispersed on alumina surfaces. These data are agreed
with that obtained from nitrogen sorption isotherms.

Textural properties
Nitrogen adsorption/desorption isotherms at 77 K for pure
and modified samples are shown in (Figure 3a and b). From
this figure, all isotherms showed Type IV profile according
to BET classification (Brunauer et al. 1938), which is char-
acteristic of porous adsorbents (Gregg & Sing 1982). The
hysteresis loop can be classified as a mixed H2 and H3
types of hysteresis loops according to International Union
of Pure and Applied Chemistry (IUPAC) classification,
which associated with capillary condensation in mesopores
materials due to the textural of inter-particle mesoporosity
(Sing et al. 1985). All the samples have a close closure point
at P/P0 = 0.4. This may actually mean that the complete
monolayer formation takes place slowly and there is an ef-
efective contribution of micropores to the adsorption on
the samples. This is confirmed by the pore size distribution
(PSD) curves that were obtained from the desorption iso-
therms; see Figure 4. It also means that the capillary con-
densation might start from the pore size at about 3 nm.

Data obtained from isotherms and BET plots of the
samples under testing are cited in Table 1. From these
data, it is clear that pure alumina sample has higher spe-
cific surface area ($S_{BET} = 187 \text{ m}^2\text{g}^{-1}$) than pure ceria
($S_{BET} = 12 \text{ m}^2\text{g}^{-1}$) which agree with previous data (Gishti et
al. 1984). The addition of ceria into alumina, xCeAl,
resulted in a gradual decrease in the surface area for the
samples ($S_{BET} = 117, 53$ and $50 \text{ m}^2\text{g}^{-1}$ for $5\text{CeAl}$, $10\text{CeAl}$
and $20\text{CeAl}$, respectively). For phosphated samples,
there is a slightly effect of phosphate on the value of spe-
cific surface area in which the $S_{BET}$ values for xCeAlP
are higher than that of xCeAl. These data agreed with
those obtained from XRD results, which complied in
Table 1, in which the variation in crystallite size has an
higher value of specific surface area. The noticeable increase in the $S_{BET}$
values for xCeAlP after phosphate addition could be attributed to the de-
crease in the crystallite size rather than any modification
in the pore structure of the pure alumina. This result
agrees with what has been reported in the literature
(Larese et al. 2004).

The t-plots, constructed using the appropriate standard
t-curve (Mikhail & Sheb 1970), are shown in Figure 5.
From the resulting curves, one can deduce that all samples
show a positive (upward) deviation in the region corre-
sponding to capillary condensation and hence indicating
the presence of mesoporosity (Mikhail & Sheb 1970). The
existence of micropores was indicated by the marginal
downward deviation of the corresponding t-plots in the
multilayer region (Ismail & Hussein 1996). This can actu-
ally find some support from the pore size distribution
curves (Figure 4). The good agreement between the $S_{BET}$
and $S_{t}$ values (Table 1) for all samples, reflects the higher
accuracy of the BET-C determination and, consequently,
the appropriateness of the reference Va-t curves (Gregg &
Sing 1982).

PSD curves of the samples under study are illustrated
in Figure 4. Examining these curves reveals that the
addition of ceria and/or phosphate on to alumina causes
a development in the porosity. In which, all samples are
in micro-meso range. Moreover, the amount of meso-
pores in the phosphate-free samples is higher than that
of phosphate samples.

| Table 1 Nitrogen adsorption-desorption data |
| Sample  | $S_{BET}$ m$^2$g$^{-1}$ | $C_{BET}$ | $S_t$ m$^2$g$^{-1}$ | Aver $r_p^{(a)}$Å | Crystallite size(b) (nm) |
|---------|------------------------|-----------|------------------|-----------------|------------------------|
| Al      | 187                    | 128       | 193              | 36.3            | 3.3                    |
| Ce      | 12                     | 6.9       | 11.8             | 15.1            | 44.4                   |
| 5CeAl   | 117                    | 98.0      | 114.3            | 20.8            | 5.4                    |
| 10CeAl  | 53                     | 74.0      | 51.2             | 19.9            | 8.1                    |
| 20CeAl  | 50                     | 58.7      | 48               | 19.3            | 9.2                    |
| AIP     | 144                    | 129       | 143              | 42.1            | 4.2                    |
| CeP     | 12                     | 3.5       | 12.1             | 12.1            | 36.2                   |
| 5CeAlP  | 144                    | 129       | 144.2            | 19.8            | 4.3                    |
| 10CeAlP | 105                    | 60.4      | 104.2            | 19.5            | 5.8                    |
| 20CeAlP | 98                     | 70.4      | 99               | 19.5            | 7.0                    |
|         |                        |           |                  |                 |                        |

(a) Mean pore radius at the peak of the distribution curves, (b) Determined using Equation 1.
Figure 3 Nitrogen sorption isotherms for (a): non-phosphated and (b): phosphated ceria/alumina catalysts, $V_a$ is measured at STP.

Figure 4 Pore size distribution curves (a): non-phosphated and (b): phosphated ceria/alumina catalysts in which $r_p$ true radius and $\Delta V_p \cdot \Delta r_p^{-1}$ / cm² g⁻¹ Å⁻¹ is the ratio between the volume decrements ($\Delta V_p$) in cm² g⁻¹ and the decrement in pore radius ($\Delta r_p$) in Å⁻¹.
FTIR spectra of adsorbed pyridine

FTIR spectrum of pyridine (Py) adsorption on pure alumina (Al) is shown (Figure 6) to display five bands at 1614, 1595, 1580, 1490 and 1440 cm\(^{-1}\). These bands are assigned to hydrogen bonded and coordinately bonded Py. The band at ~1614 cm\(^{-1}\) is due to Py coordinately bonded to Lewis acid sites of moderate strength assigned them to tetrahedral aluminum vacancies (Khalaf et al. 2007; Mekhemer et al. 2000). While the band at 1595 cm\(^{-1}\) is due to Py coordinately bonded to octahedral aluminum.
Lewis acid sites (Khalaf et al. 2007; Mekhemer et al. 2000). The other bands: 1580, 1490 and 1440 cm\(^{-1}\) is due to Py species coordinated to Lewis acid sites. No observable bands around 1540–1550 cm\(^{-1}\) indicating the absence of Brönsted acid sites on alumina surface. On pure ceria (Ce), Py adsorption has given rise to five bands at 1625, 1598, 1575, 1490 and 1440 cm\(^{-1}\). Accordingly, these bands are assignable to LPy species. The occurrence of the two bands at two different frequency values (1623 and 1595 cm\(^{-1}\)) may indicate that Lewis acid sites involved assume two different acidity strengths (Zaki et al. 1989).

Spectra taken from Py adsorption on 5CeAl, 10CeAl and 20CeAl samples at 300 K, show common feature of a gradual weakness in the intensity of the bands characteristic of the pure alumina. Py adsorption over phosphated alumina modified with ceria with different loading levels show similar spectra with phosphate free samples, so we only mention only one (xCeAl samples) to prevent the recurrence.

Isopropanol decomposition
Preliminary experiments showed that the decomposition of isopropanol on the studied oxide catalysts proceeds through simultaneous dehydration and dehydrogenation reactions (at ≥ 470 K). Thus, the decomposition of isopropanol over the catalysts under study was carried out at 470 K. The obtained results are listed in Table 2 in terms of conversion%, acetone and propene selectivity%.

The results obtained from decomposition of isopropanol on γ-Al\(_2\)O\(_3\) (Table 2) display mainly conversion of isopropanol to propene, whereas the formation of acetone is found to be very low (3.5%). The obtained results indicated that the selectivity of propene is 96.5%, thus, alumina surface have strong acidic sites, which is evident from the higher dehydration process than dehydrogenation, these results agree with that in literature (Khalaf et al. 2007). Alumina surface is known to acquire strong Lewis acid sites, which explain the high activity towards the dehydrogenation pathway (Waqif et al. 1992). The results for pure ceria show that the dehydrogenation (87.5%) of isopropanol to acetone is higher than the dehydration (12.5%) process indicating the basicity of ceria surface. Ceria modified alumina samples, xCeAl, show that the incorporation of ceria into alumina resulted in a decrease in the propene selectivity from 96.5% for alumina to 74.4% for 20CeAl. This descending in propene selectivity% is analog with ascending of acetone selectivity% and could be attributed to the basicity of ceria. (Zaki et al. 1997) reported that ceria surface has Lewis base sites assuming a strong nucleophilic reactivity.

At the same time, the conversion of isopropanol and the selectivity towards propene formation for the phosphate catalysts, xCeAlP, is higher than that for the phosphate free catalysts. Knowing that isopropanol dehydrate to propene over acidic catalysts, this could explain such behavior of the phosphated catalysts. Furthermore, the obvious increase in the catalytic activity (% conversion) of the phosphated catalysts than phosphate free samples is in accordance with the data obtained from surface area measurement, in which the \(S_{BET}\) for xCeAlP > \(S_{BET}\) for xCeAl.

Conclusions
In conclusion, the addition of ceria and/or phosphate onto alumina with different loading levels causes a modification in the surface texture and porosity. The results show also that the ceria added to alumina is well dispersed on the surface of alumina and has no effect on the phase structure of pure alumina (γ-Al\(_2\)O\(_3\) structure). Alumina, ceria and ceria-alumina composite show Lewis acid sites with different acidity strengths. Isopropanol decomposed to mainly propene by pure alumina and the addition of ceria decreases the amount of propene selectivity% and increases the acetone selectivity%, this is due to the basicity of ceria. Moreover, the phosphated samples have higher propene selectivity% than phosphate free samples indicating that the phosphate ions increase the acidity of the samples.

Experimental
Materials
Two series of the catalysts under studying have been prepared. The first series is ceria doped alumina catalysts, abbreviated as xCeAl (where, \(x = 5, 10 \text{ and } 20 \text{ wt\% ceria}\)) were prepared by coprecipitation method. The 1.5 molar solutions of Al(NO\(_3\))\(_3\)∙9H\(_2\)O and Ce(NO\(_3\))\(_3\)∙6H\(_2\)O were prepared separately and then mixed in a volume proportions according to the final desired composition of catalysts. The resulting solution was stirred and heated to 350 K in a round bottom flask. The aqueous 1:1 M
NH₄OH solution was added drop wise to the nitrate solution under vigorous stirring until pH 8 was attained. After ageing for two hours, the excess solution was removed by filtration. Precipitate was washed repeatedly by distilled water followed by drying at 380 K for 12 h. The dried precursor was crushed to fine powder and calcined in the presence of static air at 873 K for 3 h.

The second series, xCeAlP (P refers to phosphate and x is the loading level of ceria, 5, 10 and 20 wt%), was obtained by impregnation of an aqueous solution of the impregnating (NH₄)₂HPO₄ solution and held stirring for 1 h. The solution was adjusted to give 6 Wt% PO₄³⁻ content. The excess water present was removed by evaporation at 380 K. The catalysts were obtained by calcination of the dried samples at 873 K for 3 h.

Apparatus and techniques

Thermal analysis
Thermogravimetric analysis (TGA) and deferential thermal analysis (DTA) were performed between room temperature and 1273 K in a static atmosphere of air, using Linseis STA PT 1600 thermogravimetric analyzer. The rate of heating was standardized at 10 K/min., and small portions (5–15 mg) of the sample were used in TG measurements.

X-ray powder diffractometry
XRD diffractograms were recorded for all samples using a model JSX-60PA JEOL diffractometer (Japan) using Cu Ka radiation (λ = 1.5418 Å). The generator was operated at 35 kV and 20 mA. The samples were scanned in the range of 2θ = 10–70° at a scanning speed of 6° min⁻¹. For identification purposes, diffraction patterns (I/I°) versus d spacing (Å) were matched with the relevant ASTM standards (Frank et al. 1981). The crystallite size D of the samples were calculated using the Scherrer relationship (Klug & Alexander 1970):

\[ D = \frac{K \lambda}{\beta \cos \theta} \]  

where K is the crystallite shape constant (=1), λ the radiation wavelength, β the line breadth (radians) and θ is the Bragg angle.

Nitrogen adsorption isotherm measurement
Full nitrogen adsorption/desorption isotherms at 77 K were obtained using a NOVA 2200, version 6.10 high-speed gas sorption analyzer (Quantachrome Corporation USA). The calcined samples were first out-gassed at 470 K for 1 h. Twenty four-point adsorption and desorption isotherms were obtained, from which BET surface areas were derived using standard and well-established methods (Sing et al. 1985; Webb & Orr 1997).

FTIR spectra for pyridine adsorption
For pyridine (Py) adsorption on test samples a wafer of 15–20 mg/cm² were mounted in a Pyrex vacuum cell fitted with CaF₂ windows. The samples were pretreated at 700 K for 1 h in stream of O₂ followed by evacuation at 700 K for 1 h, then cooled to room temperature to obtain the background IR spectra using Thermo scientific Nicolet 380 FTIR spectrophotometer. Then, 5 Torr Py (1 Torr = 133.32 Pa) were admitted at 300 k for 5 min., degassed for 5 min. at this temperature in order to remove the fraction physisorbed Py, and the spectra were then taken at room temperature.

Catalytic activity (isopropanol decomposition)
The catalytic activity experiments for isopropanol decomposition were carried out in a fluidized bed quartz flow reactor at atmospheric pressure. 0.2 g of the catalyst was activated in-situ at 670 K for 1 h in N₂. isopropanol (Merck, product, 99.9%) was introduced at a flow rate of 15 ml min⁻¹ into carrier gas flow of N₂. The reaction products were analyzed by gas chromatography on 2-m long 1/8'' column packed with 10% Carbosax and Chromm WHP 80/100 using a model 3400 Varian Gas Chromatograph equipped with a flame ionization detector (FID).

Competing interests
The author declares that he has no competing interests.

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