

Insight into the gas-phase glycerol dehydration on transition metal modified aluminium phosphates and zeolites

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Abstract

BACKGROUND: The production of biodiesel has notably increased in recent years, with glycerol as the main by-product, an important raw material in the production of value-added products. Those obtained by dehydration are especially important.

RESULTS: Gas phase dehydration of glycerol has been tested on mesoporous AlPO_4 and modified AlPO_4 , with a small amount (1 wt%) of a transition metal (Co, Cu, Cr, Fe), calcined at 450°C and 650°C, and compared with some commercial zeolites (HZSM and HY). All the catalysts were active in the reaction, the phosphates showing better catalytic behaviour than the zeolites. The main products of the reaction were acrolein and hydroxyacetone.

CONCLUSIONS: The positive effect of metal on the catalytic behaviour of AlPO_4 in the dehydration has been verified, attaining the maximum yield to acrolein (54%) on AlCoPO_650 with 50 mg of catalyst and 280°C of reaction temperature. Furthermore, the phosphates did not suffer deactivation after 22 h reaction, whereas the zeolites suffered a strong loss of activity (around 70%) due to blockage of the micropores by coke. Moreover, the yield to acrolein decreased with time, while the yield to hydroxyacetone increased. An E_2 mechanism for the acrolein formation and E_1 for the hydroxyacetone formation have been proposed.
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Keywords: glycerol; acrolein; hydroxyacetone; metal-aluminium phosphates; zeolites; deactivation

INTRODUCTION

In recent years, the production of biodiesel has notably increased. According to the FAO (Food and Agriculture Organization) this tendency is going to continue in the future, but at a slower rate than in the past.¹ Biodiesel production by transesterification of triglycerides generates glycerol (1,2,3-propanetriol) as the main by-product. This fact has led to a drastic surplus of glycerol in the chemical markets.^{2,3} The composition of the crude glycerol, obtained directly from biodiesel production, generally oscillates between 50 and 80% glycerol, 3–15% water, 5–15% salt, <1–20% methanol, 1–5% fatty acids and 1–5% non-glycerol organic material, by weight.³ This crude glycerol has little value due to its impurities, thus employing it to produce value-added products would increase its value. Among these products, those obtained from glycerol dehydration, propen-2-al or acrolein (ACR) and hydroxyacetone (HA) or acetol, are especially important. ACR can be used as a raw material to produce DL-methionine and acrylic acid and its derivatives, whereas HA is an intermediate for the production of propanediols and can be used as a flavour in the food industry, as dyes, as additives in cosmetics and so on.

Glycerol dehydration reaction has been investigated mainly on heterogeneous catalysts, as has been indicated in recent reviews.^{4,5} Typically, the reaction is carried out in the temperature range 250–350°C using aqueous glycerol as a feedstock, which

is advantageous because glycerol is usually obtained mixed with water, as aforementioned. Furthermore, the reaction in gas phase generally occurs with better results than in liquid phase, as we have previously reported.^{6,7}

The acidity of the catalyst plays an important role not only in its activity and selectivity in glycerol dehydration, but also in its stability or resistance to deactivation. However, there is no agreement in the literature about the nature and strength of the acid sites implied in the formation of the main dehydration products. Thus, Chai *et al.*⁸ based on the study of various solid catalysts with a wide range of acid-base properties reported that acrolein formation is enhanced by strong acid sites, mainly Brønsted acid sites, whereas the basic solids hardly showed any selectivity to acrolein. Likewise, some recent studies have proposed that the formation of acrolein takes place on Brønsted acid sites^{9–12} and that of hydroxyacetone on Lewis acid sites.^{9,12} Foo *et al.*¹⁰ have demonstrated the cooperative role of Brønsted and Lewis acid sites in the

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glycerol dehydration to hydroxyacetone over niobia catalysts, whereas, Wang *et al.*¹³ indicated this cooperative effect but for the acrolein formation. Kinage *et al.*¹⁴ on sodium-doped metal (Ce, Al, Ga, Zr) oxides even reported the participation of the basic sites in the formation of HA, proposing a complicated reaction scheme implicating three steps: glycerol dehydrogenation, dehydration of the obtained compound and rehydrogenation of this latter.

More agreement seems to exist about the participation of the very strong acid sites, mainly Brønsted sites, in the coke formation,^{6–8,12,15} which provokes catalyst deactivation, mainly by pore blocking. The microporous materials being more quickly and widely deactivated than those proving to be meso or macroporous.^{8,16,17}

On this basis, the aim of the present research is to verify the influence of the acid sites (nature and strength) and textural properties of the catalysts on its activity and stability in the formation of glycerol dehydration products in the gas phase, as we previously demonstrated in liquid phase.⁷ Thus, we have studied materials such as a mesoporous aluminium phosphate (AlPO₄) and different zeolites (HZSM and HY). Furthermore, we have also studied the influence of a second metal (Co, Cu, Cr, Fe) on the catalytic behaviour of aluminium phosphate. In previous work,⁶ we demonstrated the favourable effect of a metal with redox properties, such as vanadium, on the activity in the acrolein formation of AlPO₄, whereas Ca did not have this effect.

The solids have been characterized by X-ray diffraction (XRD); scanning electron microscopy (SEM); ³¹P and ²⁷Al magic angle spinning nuclear magnetic resonance (MAS NMR) and diffuse reflectance ultraviolet-visible (DR-UV-Vis) spectroscopy. Their acid properties were determined from temperature programmed desorption (TPD) and diffuse reflectance infrared Fourier transform (DRIFT) of adsorbed pyridine. Furthermore, the test reaction of 2-propanol transformation in the absence of oxygen was also employed in evaluating the acid sites (responsible for dehydration to propene) and the basic/redox sites (responsible for propanone formation) of the solids.

A study on the formation and nature of coke over time on stream, has also been carried out. In this respect, an off-line analysis of the effluent from the reactor was carried out in order to avoid deposits of unidentified compounds, which were responsible for some obstruction of the reaction system lines, as previously reported.⁶

EXPERIMENTAL

Catalyst preparation

Aluminium phosphate (AlPO₄) was prepared by a sol-gel method previously reported.^{18–20} It was synthesized from AlCl₃·6H₂O and H₃PO₄ (85 wt%) by precipitation with aqueous ammonia under stirring at 0°C. The precipitate was allowed to stand at room temperature and the pH value at the 'end point' was 6.7. After filtration the solid was washed with 2-propanol, dried at 120°C for 24 h and calcined in air at 450°C for 3 h. The binary system of Co, Cu, Cr and Fe¹⁸ were prepared from an aqueous solution with the appropriate amounts of AlCl₃·6H₂O, of the corresponding nitrate and of H₃PO₄, in order to obtain a molar ratio P/Al + M = 1 and equivalent to the theoretical molar ratio Al/M = 49, 52, 42 and 50, respectively. The metal percentage added in those systems was 1 wt%. They were calcined in air at 450°C and 650°C for 3 h. These systems will be denoted as AlMPOT, where T indicates the calcination temperature. For comparative purposes, another solid was prepared with Al and Co (Al-CoPOT) changing the order in which the metal salts and H₃PO₄ were added. Thus, AlCl₃·6H₂O was dissolved in water,

H₃PO₄ was then incorporated and finally, Co(NO₃)₂·6H₂O, after stirring. From this point the procedure continued as has been previously described. All the solids were screened at <0.149 mm to avoid internal diffusion limitations in the reactions.

Three commercial zeolites from Zeolyst⁷ were used: two were ZSM-5 in the NH₄⁺ form with a different SiO₂/Al₂O₃ ratio (CBV 5524G and CBV 3024E), and the other was zeolite HY (CBV 600).⁷ The ZSM zeolites were converted into the H form after activation in a stream of dry air for 6 h at 600°C. This same thermal treatment was employed for the HY(5.2) zeolite. These solids will be denoted as H-ZSM-5(50); H-ZSM-5(30) and HY(5.2), where the number in brackets indicates the SiO₂/Al₂O₃ ratio.

Characterization of the catalysts

X-ray diffraction (XRD) patterns were performed on a diffractometer (Siemens D-5000), equipped with an automatic control system. A Ni-filtered, graphite monochromator, Cu K_α radiation (λ = 1.5406 Å) was used, operating at 35 kV and 20 mA. The samples were scanned 2° ≤ 2θ ≤ 80° at a speed of 2° min⁻¹.

Scanning electron microscopy (SEM) studies were carried out in a Jeol apparatus, model JSM 6300, equipped with an Oxford Instrument detector, model Link ISIS which allows the chemical analysis by energy dispersion of X-rays (EDX) of the studied solid surface. The measurements were repeated at least three times on different zones of the solid surface (experimental error ≤ 7%).

Diffuse reflectance spectra in the UV-Vis region (200–800 nm) were collected at room temperature with a Variant Cary 1E UV-Vis spectrophotometer equipped with reflectance attachment using BaCl₂ as reference.

³¹P and ²⁷Al MAS NMR spectra were collected on a Bruker Avance 400WB spectrometer using a 4 mm Bruker double resonance MAS probe. All experiments were performed nominally at room temperature. ²⁷Al and ³¹P MAS NMR were recorded at resonance frequencies, ν₀, of 104.26 and 161.98 MHz and Al(H₂O)₆³⁺ and H₃PO₄ solutions were used as external standard, respectively. The data were processed by Fourier transformation, phase correction and baseline correction using the Dm2005test NMR data processing software.

The textural properties (BET surface area, S_{BET}, pore volume, V_p, and average pore diameter, d_p) were obtained from N₂ adsorption-desorption isotherms, at liquid nitrogen temperature (77 K), using a Micromeritics ASAP 2000 instrument. Before measurements, all samples were degassed to 0.1 Pa. The pore size distribution has been determined using the Barrett, Joyner and Halenda (BJH) method, assuming a cylindrical pore model. The surface area values were calculated by applying the BET equation at relative pressures in the range p/p₀ = 0.05–0.30, assuming a cross-sectional area of 0.162 nm² for the nitrogen molecule.

Total acidity was measured by temperature programmed desorption of pyridine (Py-TPD), using an apparatus from PID Eng&Tech equipped with a TCD (thermal conductivity detector). Before starting the adsorption experiments, the catalyst (30 mg) was pre-treated *in situ* from room temperature to 400°C (rate of 20°C per min). After the catalyst pre-treatment, saturation of the sample with pyridine was carried out at 50°C for 30 min, thereafter, physisorbed pyridine was desorbed at 50°C for 60 min. Then, the measurements were performed in the range of 50–450°C, with a heating rate of 10°C per min to remove the chemisorbed pyridine. The exceptions were the zeolites, in which the maximum temperature in the pretreatment and in the measurements was 500°C and 550°C, respectively.

Diffuse reflectance infrared Fourier transform (DRIFT) spectra of adsorbed Pyridine were recorded on a FTIR instrument (Bomem MB-3000) equipped with an 'environmental chamber' (Spectra Tech) placed in a diffuse reflectance attachment (Spectra Tech, Collector). A resolution of 8 cm^{-1} was used with 256 scans averaged to obtain a spectrum from $4000\text{ to }600\text{ cm}^{-1}$. The reference spectrum of the sample was recorded at 350°C . Pyridine (99.8% from Aldrich) adsorption was carried out at 120°C for a certain period of time (typically 1 h) allowing saturation of the catalyst surface. The samples were subsequently degassed in four steps of 1 h each at 50, 150, 200 and 300°C , respectively.

The 2-propanol conversion was carried out in a continuous-flow fixed bed micro-catalytic reactor (4 mm internal diameter and 90 mm long) at atmospheric pressure. The catalyst (20 mg, $<0.149\text{ nm}$) was pretreated by *in situ* heating under nitrogen (20 mL min^{-1}) for 30 min at 300°C . The 2-propanol (99.5% from Panreac) was fed by means of a microfeeder at 0.6 mL h^{-1} ($2.19\text{ }\mu\text{mol s}^{-1}$) and the flow of nitrogen was 20 mL min^{-1} . The catalytic measurements were performed at 170, 190, 210 and 230°C . The final values reported are the average of six values obtained after the stationary state was attained (15 min). The reaction products (propene, propanone and diisopropyl ether), as well as the 2-propanol unconverted were analysed by gas chromatography (CG) with a flame ionization detector (FID), by using a capillary column ($60\text{ m} \times 0.25\text{ mm} \times 0.25\text{ }\mu\text{m}$) DB-WAX (100%, polyethylene glycol). Under these conditions, no diffusional limitations exist. The estimated experimental error was $<4\%$. The catalysts did not suffer deactivation.

Raman spectra were acquired with a Renishaw Raman instrument (inVia Raman Microscope) by excitation with green laser light (532 nm) and a grating of 1800 lines per mm. A total of 50 scans per spectrum were performed in order to improve the signal-to-noise ratio, with an exposure time of 15 s.

Coke determination was carried out by thermogravimetric analyses (TGA) using a Setaram Setsys 12 thermobalance in the presence of static air with a 20 mg sample. The heating rate was $10^\circ\text{C min}^{-1}$ (temperature range: $30\text{--}1000^\circ\text{C}$).

Catalytic test

Glycerol transformation was carried out in a continuous-flow fixed-bed microreactor (0.35 cm internal diameter and 9 cm long) under atmospheric pressure. Reaction products together with unconverted glycerol were collected in two consecutive ice traps and they were analysed without preliminary extraction or separation of water. The analysis was carried out by gas chromatography (GC) equipped with a flame ionization detector (FID) using a 100% poly(ethylene glycol) ($25\text{ m} \times 0.25\text{ mm} \times 0.25\text{ }\mu\text{m}$) capillary column.

The catalyst (50 mg, $<0.149\text{ }\mu\text{m}$) was pretreated at the reaction temperature 280°C , for 2 h in a N_2 flow ($50\text{ cm}^3\text{ min}^{-1}$). A 36 wt% glycerol (99.5%, Sigma–Aldrich) aqueous solution was fed at 0.3 mL h^{-1} ($1.27 \times 10^{-3}\text{ mol h}^{-1}$), which resulted in a contact time of $2.38\text{ g}_{\text{cat}}\text{ min mmol}^{-1}$. In general, each catalytic test was conducted for 22 h. The products were identified through chromatographic patterns and/or GC-MS (Varian CP 3800, Quadrupole MS 1200) equipped with a Supelcowax 10 (100% poly(ethylene glycol), $30\text{ m} \times 0.25\text{ mm} \times 0.25\text{ }\mu\text{m}$) capillary column. A blank test showed the absence of homogeneous reactions and the inactivity of the reactor in the absence of a catalyst.

In order to avoid the polymerization of acrolein,²¹ hydroquinone was added (0.2 wt%) to the aqueous solution of glycerol in the feed and also to the cold trap where the reaction products were

collected. The hydroquinone also allowed the carbon balance to be estimated. The absence of hydroquinone reactivity has been verified by tests carried out, with or without this compound under reaction conditions.

The glycerol conversion (X_{Gly}) was calculated from $X_{\text{Gly}} = (\text{mol}_{\text{Gly,in}} - \text{mol}_{\text{Gly,out}}) \cdot 100 / \text{mol}_{\text{Gly,in}}$, whereas the selectivity (S_i) and the yield (Y_i) to product i were expressed as mol% on a C atom basis.

RESULTS AND DISCUSSION

Structural and textural characterization of the solids

The XRD patterns (not shown) of the phosphates showed only a very broad hump in the range 2θ between 15 and 40° , a characteristic of amorphous metal phosphates, regardless of the metal added and the calcination temperature. However, the commercial zeolites showed a crystalline character.⁷ Both H-ZSM zeolites exhibited a similar XRD pattern which corresponds with a MFI structure, whereas the HY(5.2) exhibited a Faujasite structure.

The micrographs obtained by SEM (Fig. 1) confirmed the amorphous character of the solids also showing a very unhomogeneous distribution in morphology and particle sizes, independently of the presence of one or two metals on the solid, as well as the calcination temperature. With regard to the shape of the particles, both elongated and rounded particles were observed. In comparison with the theoretical composition [Al/M = 49 (Co), 42 (Cr) and 50 (Fe)], the modified aluminium phosphates showed a superficial enrichment of M according to the results obtained by EDX analysis, Table 1. This fact was emphasized in the solids calcined at 650°C , as well as in Al-CoPO in comparison with AlCoPO.

^{31}P and ^{27}Al MAS NMR spectra of the solids studied are collected in Fig. 2. For all the phosphates, the ^{31}P NMR showed a single peak around -25 ppm , typical of amorphous aluminium phosphates, which can be assigned to P atoms in tetrahedral coordination. Hence, the environments of phosphorus nuclei are mainly $\text{P}(\text{OAl})_4$.^{22,23}

On the other hand, the ^{27}Al NMR of the different phosphates showed a main resonance at approximately $+38\text{ ppm}$, which is typical of tetrahedral aluminium sharing oxygen atoms with four phosphorus tetrahedrals ($\text{Al}(\text{OP})_4$). Other small resonances at $+14$ and -13 ppm are ascribed to Al species partially coordinated with hydroxyl groups in octa- and pentahedral coordination, respectively.^{22–24} With the presence of a second metal in the AlPO_4 a slight increase of both components was observed, especially in Al-CoPO450 and AlFePO450.

The UV-Vis spectra obtained for the solids studied and some reference compounds are shown in Fig. 3. Absorption of the AlPO_4 450 was insignificant in comparison with those of the modified phosphates, therefore we can assume that the absorptions of the modified phosphates are related to the charge transfer between transition metal and oxygen. The oxidation state of each metal was as expected in accordance with the synthesis of modified phosphates. Thus, the aluminium phosphates modified with Co showed similar spectra, with a broad asymmetric band between 530 and 700 nm, with three maxima of absorption that can be assigned to Co^{2+} in a tetrahedral environment.²⁵ The absorption at 473 nm in the cobalt phosphate spectrum is related with Co^{2+} in an octahedral coordination. The spectra presented another much weaker band at 313 nm, that would indicate the presence of some tetrahedral Co^{3+} in the framework of the solid.²⁵ In the spectra of the solids with Cu, a broad band appeared at 235 nm that is ascribed to the charge transfer transition between O^{2-} and Cu^{2+} .²⁶ These samples

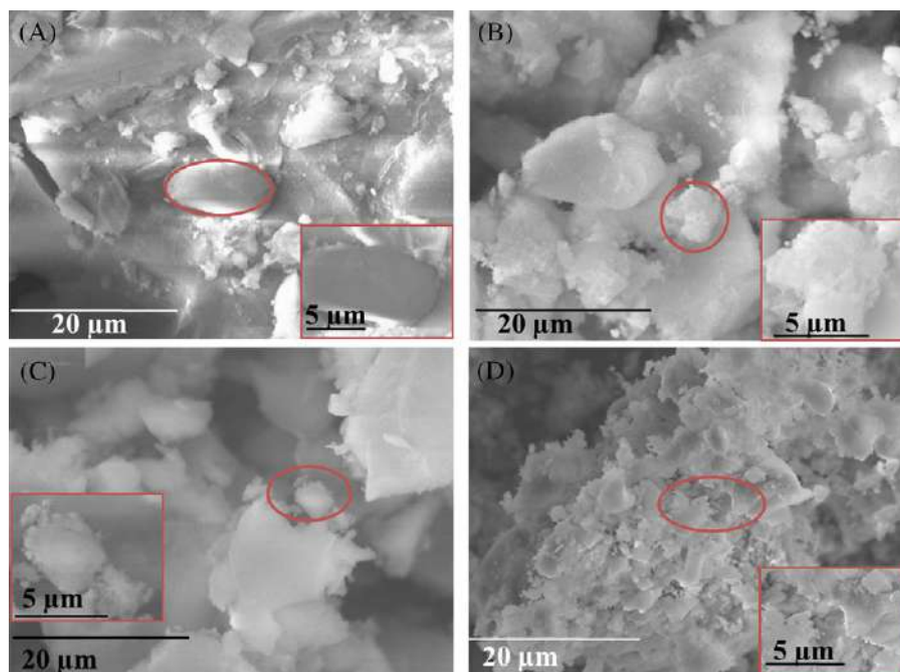


Figure 1. SEM micrographs: (A) AlPO_4 450; (B) Al-CoPO450; (C) AlCrPO450; (D) Al-CoPO650.

Table 1. EDX superficial chemical composition and textural properties of the solids studied								
Catalyst	EDX		S_{BET} ($\text{m}^2 \text{g}^{-1}$)	V_p (mL g^{-1})	d_p (Å)	Pore size distribution (%)		
	Al/M	P/(M + Al)				>500 Å	500-20 Å	<20 Å
AlPO_4 450	-	1.08	199.1 ± 0.8 (137.6 ± 0.4)	0.82 (0.60)	165 (175)	12 (14)	87 (86)	- (-)
AlCoPO450	48	1.03	256.1 ± 0.8 (144.7 ± 0.5)	0.94 (0.43)	146 (120)	13 (22)	87 (78)	- (-)
AlCoPO650	43	1.04	225.5 ± 0.6	0.85	150	14	86	-
Al-CoPO450	42	1.07	262.3 ± 0.6	0.80	122	10	90	-
Al-CoPO650	33	1.25	243.8 ± 0.7	0.77	127	13	87	-
AlCuPO450	n.d. ^a	-	213.4 ± 0.7	0.82	154	5	95	-
AlCuPO650	n.d. ^a	-	201.6 ± 0.7	0.82	162	15	85	-
AlCrPO450	35	1.21	262.6 ± 0.7	0.82	125	9	91	-
AlCrPO650	34	1.22	247.8 ± 0.7	0.79	127	9	91	-
AlFePO450	33	0.88	248 ± 1 (200.1 ± 0.6)	1.04 (0.79)	168 (153)	17 (26)	82 (76)	-
H-ZSM-5(50)	-	-	425^b	$0.27 (0.15)^c$	19.3	20	29	51
H-ZSM-5(30)	-	-	400^b (49.2 ± 0.6)	$0.27 (0.12)^c$ (0.26) (0)	22.6 (215)	23 (79)	35 (20)	42 (1)
HY(5.2)	-	-	660^b (32.4 ± 0.2)	$0.36 (0.20)^c$ (0.12) (0)	23.0 (149)	21 (60)	31 (39)	48 (1)

In brackets, used catalysts after 22 h.
^a Not determined.
^b Data from Zeolyst.
^c Volume of micropores of the fresh catalysts obtained by the t-plot method.

showed a greenish-blue colour, the pyrophosphate being greener than the binary solids. The spectra of the AlCrPO450 showed four bands. The bands at 266 and 349 nm are normally assigned to Cr^{6+} of chromate and/or polychromate species,²⁷ the latter could also be ascribed to the third transition of a pseudo-octahedral Cr^{3+} species.²⁸ The bands at 449 and 618 nm are assigned to d-d transitions of the Cr^{3+} in distorted octahedral coordination.²⁷ Both

samples were yellowish green. The spectrum of the AlFePO₄450 as with that corresponding with an iron phosphate, exhibited a broad band with two maxima at 240 and 310 nm, indicating Fe^{3+} in tetrahedral and octahedral coordination, respectively.¹⁸

According to the textural analysis, the nitrogen isotherms obtained for all the metal phosphates studied were type IV of the Brunauer, Deming, Deming and Teller (BDDT) classification,

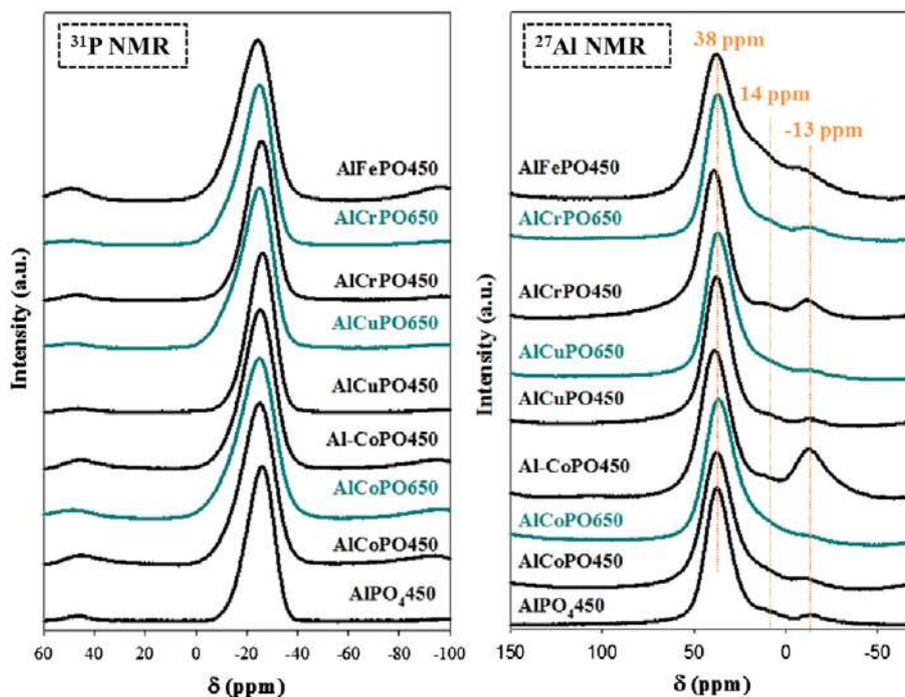


Figure 2. ^{31}P and ^{27}Al MAS NMR spectra for the phosphates studied.

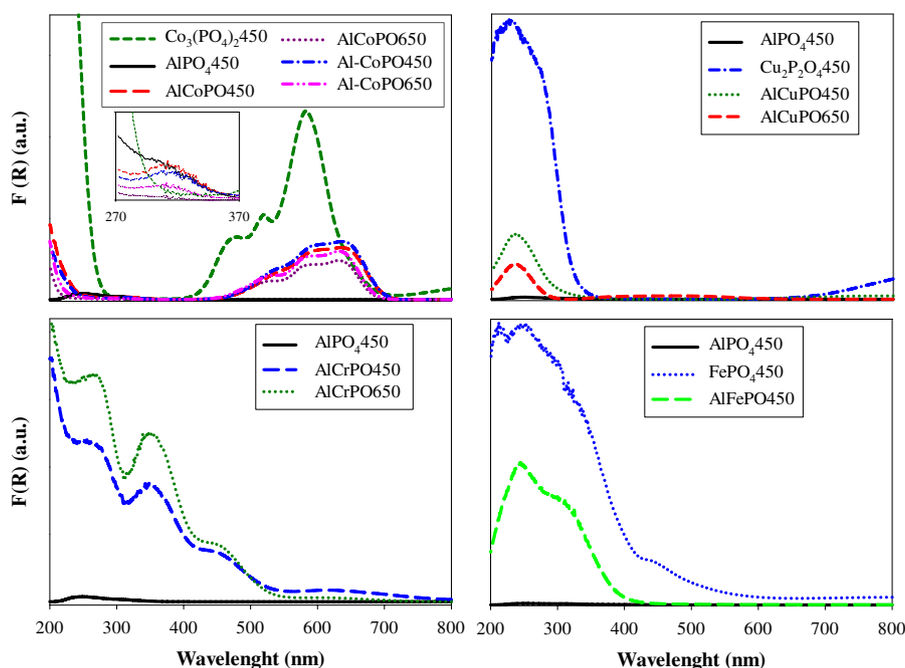


Figure 3. UV-Vis spectra.

exhibiting type H1 hysteresis loops, which corresponded with mesoporous solids. Table 1 compiles the values of S_{BET} , the pore volume, the average pore size, as well as the percentage of macropores (>500 Å), mesopores (20–500 Å) and micropores (<20 Å) for all the phosphates. As can be seen, the presence of a second metal gave rise to an increase in the S_{BET} between 25 and 32%, except for the AlCuPO450 whose increase was less (7%). On the contrary, the d_p value decreased, except for the AlFePO450 which remained almost constant. In addition, the increase of the

calcination temperature produced a decrease in the S_{BET} (between 6 and 12%) and a little increase in the d_p . It is worth noting that none of the phosphates presented micropores, and the percentages of macropores and mesopores in AlPO₄ were not changed significantly by the presence of a second metal. Similar S_{BET} values were obtained for the solids with Co, independently of the synthesis procedure. However, small variations in the V_p and d_p were observed, being lower in the Al-CoPO system. With regard to the zeolites, they exhibited higher S_{BET} and lower d_p and V_p values

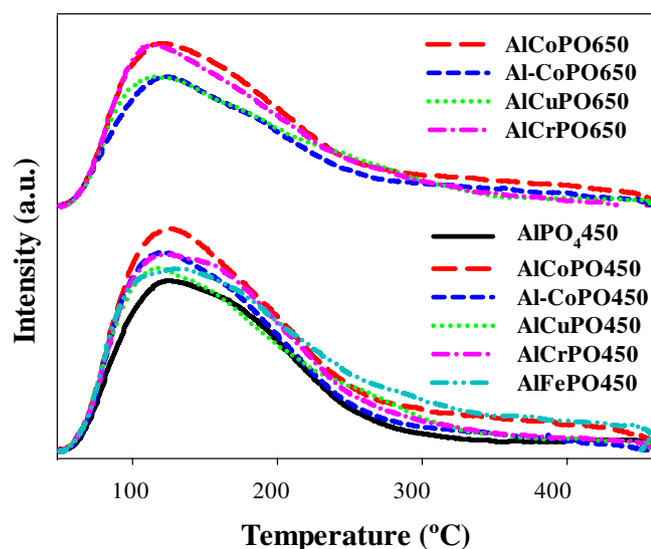


Figure 4. Py-TPD profiles observed for the phosphates studied.

than metal phosphates, showing around 50% micropores, 20% macropores and 30% mesopores, Table 1.

Acid and base properties

As far as the acid characterization is concerned, the Py-TPD profiles of the phosphates, Fig. 4, showed a broad band centered between 100°C and 200°C. As previously reported,⁷ the zeolite profiles also showed one maximum peak at around 120°C and another of less intensity, around 500°C. With the exception of Al-CoPO450, AlCrPO650 and HY(5.2) profiles, the others never reached the baseline.

The values of total acidity and the percentage of acid sites as a function of their strength are collected in Table 2. The total acidity of the phosphates, as happened with the S_{BET} , increased with the presence of a second metal in the solids with a remarkable rise in the AlFePO450 (39%). Likewise, the increase of calcination temperature promoted a decrease of total acidity. In addition, Al-CoPO exhibited lower total acidity than AlCoPO, which could be due to the highest percentage of cobalt on the surface of Al-CoPO. On the other hand, the density of acid sites of all the phosphates oscillated between $0.5 \mu\text{mol m}^{-2}$ (Al-CoPO650) and $0.8 \mu\text{mol m}^{-2}$ (AlFePO450 and AlCuPO450), weak acid sites predominating ($\geq 60\%$) in all the phosphates. It was remarkable that the AlFePO450 and the AlCoPO systems presented the highest percentage and density of strong acid sites. As far as the zeolites are concerned,⁷ both H-ZSM-5 adsorbed approximately the same amount of pyridine, independently of the $\text{SiO}_2/\text{Al}_2\text{O}_3$ ratio, whereas the HY(5.2) adsorbed the highest quantity of pyridine. Furthermore, the zeolites exhibited a total acidity higher than the majority of the binary phosphates, but a lower density of acid sites. Moreover, all the zeolites showed mainly weak acid sites, as occurred with the phosphates, although the percentage of strong acid sites was superior to that of the phosphates. The HY(5.2) zeolite exhibited the highest percentage and quantity ($62 \mu\text{mol g}^{-1}$) of this type of site.

The DRIFT spectroscopy allows distinguishing the nature of acid sites, because adsorbed pyridine shows different bands between 1400 and 1700 cm^{-1} depending on the type of acid sites. The pyridine adsorption bands around 1440–1460 cm^{-1} correspond to pyridine adsorbed on Lewis sites (L) and bands around 1540–1560 cm^{-1} are associated with Brønsted sites (B).^{7,20,29} In general, all

the solids exhibited similar spectra to that shown for AlCoPO650 (Fig. 5(A)), indicating that all the solids analysed showed bands associated with both, B and L. In order to estimate which type of acid sites were more abundant and their strength, the B/(L + B) ratio has been calculated at two temperatures, 150°C and 300°C desorbed pyridine. The results shown in Table 2 indicate that all the phosphates possess more Lewis acid sites than Brønsted acid sites. Furthermore, the presence of a second metal in the solids, as well as the thermal treatment at a higher temperature produced an increase in the percentage of Lewis acid sites. Regarding the strength of the sites, desorption of pyridine at 300°C indicated that Lewis acid sites were stronger than Brønsted acid sites. In the case of the zeolites, the opposite occurred; in general, the zeolites⁷ exhibited a higher percentage of Brønsted acid sites than the phosphates, Lewis acid sites increased as the Al_2O_3 increased. In fact the H-ZSM-5(50) exhibited the highest percentage of Brønsted acid sites.

Figure 5(B) shows the OH stretching vibration region (3000–4000 cm^{-1}) for the AlCoPO650. The band at 3670–3680 cm^{-1} is assigned to isolated germinal O–H attached to the phosphorus atom or to single P–OH groups, which exhibit nearly the same OH stretching frequencies but different combination frequencies, and the broad band with maximum at 3780–3790 cm^{-1} is the characteristic band of surface Al–OH with Al atoms in tetrahedral coordination.^{20,29} The P–OH groups represent the most stable Brønsted acid sites on AlPO_4 surfaces, whereas Al–OH would enhance the acidity of the P–OH groups through H-bonding.^{20,29} All the solids presented the two bands, which gradually decreased with increase of the degassing temperature. In the zeolites spectra,⁷ bands at 3735 cm^{-1} and 3600 cm^{-1} could be assigned to the vibration of terminal silanol groups (Si–OH) located at the external surface and to the O–H vibration of bridging Si(OH)Al groups located at the internal zeolites surface, respectively. The intensity of the second band diminished more slowly than the first one, indicating higher strength of the bridging OH in comparison with the terminal silanol groups.

The transformation of 2-propanol takes place preferably by two reactions: dehydration to propene (C_3H_6), mainly on acid sites, and dehydrogenation to propanone ($\text{C}_3\text{H}_6\text{O}$), mainly on basic/redox sites.²² Furthermore, under certain experimental conditions, such as low temperatures and high partial pressure of 2-propanol, the formation of diisopropyl ether ($\text{C}_6\text{H}_{14}\text{O}$), would be favoured also on acid sites. As can be seen in Table 2, all the catalysts produced propene as the major product (selectivity values $>66\%$) and diisopropyl ether at lower percentage (selectivity values below 16%). Consequently, in all the solids studied the acid sites dominated. Propanone was obtained on all catalysts with the exception of AlPO_4 . This result corroborates that previously reported,²² in the sense that the formation of propanone requires basic sites with redox ability. Unfortunately, the low percentage of transition metal in the AlPO_4 did not allow for detection of H_2 in the TPR (temperature programmed reduction) experiments. Evaluation of the surface acidity and of the basic/redox sites of the solids was carried out based on the selectivity values to propene and propanone, respectively, obtained at isoconversion. This has been made, given that the values of selectivity to propene increased and selectivity to propanone decreased, as the reaction temperature (or the 2-propanol conversion) increased. The values of selectivity (S_i) obtained at isoconversion ($\approx 1 \text{ mol}\%$), together with the values of temperature necessary to get such a conversion value, are collected in Table 2. As can be seen, the modified AlPO_4 were slightly more active than the aluminium phosphate itself, although

Table 2. Acidity measurements from Py-TPD; B/L + B ratio for each catalyst after pyridine degassing for 1 h at 150°C and 300°C; and temperature and selectivity at isoconversion (≈ 1 mol%) for 2-propanol decomposition ($w = 0.02$ g; $F_{C_3H_8O} = 0.6$ mL h $^{-1}$)

Catalyst	Acidity		Strength of acid sites (%)			Acid sites nature		T_{ISO} (°C)	2-propanol conversion		
	($\mu\text{mol g}^{-1}$)	($\mu\text{mol m}^{-2}$)	Weak (80–200°C)	Medium (200–300°C)	Strong (>300°C)	B/L + B (150°C)	B/L + B (300°C)		$S_{C_3H_6}$	$S_{C_3H_6O}$	$S_{C_6H_{14}O}$
AlPO ₄ 450	144 (77) ^a	0.7 (0.6)	72 (79)	20 (15)	8 (6)	0.47	0.38	210	85	-	15
AlCoPO450	186 (133)	0.7 (0.9)	66 (54)	21 (19)	13 (27)	0.37	0.24	170	83	17	-
AlCoPO650	169	0.7	66	21	13	0.19	0.22	170	88	12	-
Al-CoPO450	166	0.6	73	20	7	0.24	0.29	190	75	16	9
Al-CoPO650	127	0.5	68	21	11	0.22	0.15	190	86	4	10
AlCuPO450	162	0.8	69	23	8	0.24	0.20	170	87	13	-
AlCuPO650	133	0.7	67	25	8	0.19	0.13	190	86	11	3
AlCrPO450	179	0.7	70	22	8	0.16	0.15	170	82	18	-
AlCrPO650	151	0.6	72	22	6	0.15	0.15	190	89	5	6
AlFePO450	200 (115)	0.8 (0.6)	60 (50)	25 (27)	15 (23)	0.46	0.38	n.d. ^b	-	-	-
H-ZSM-5(50)	202	0.5	80	3	17	0.82	0.74	170 (1) ^c	83	17	-
H-ZSM-5(30)	187	0.5	81	6	13	0.47	0.61	170 (6) ^c	94	2	4
HY(5.2)	281 (32)	0.4 (1.0)	67 (92)	11 (4)	22 (4)	0.40	0.53	170 (32) ^c	98	1	1

^a In brackets, used catalysts after 22 h.

^b Not determined.

^c Conversion values (mol %) at 170°C (zeolites cannot be compared at isoconversion).

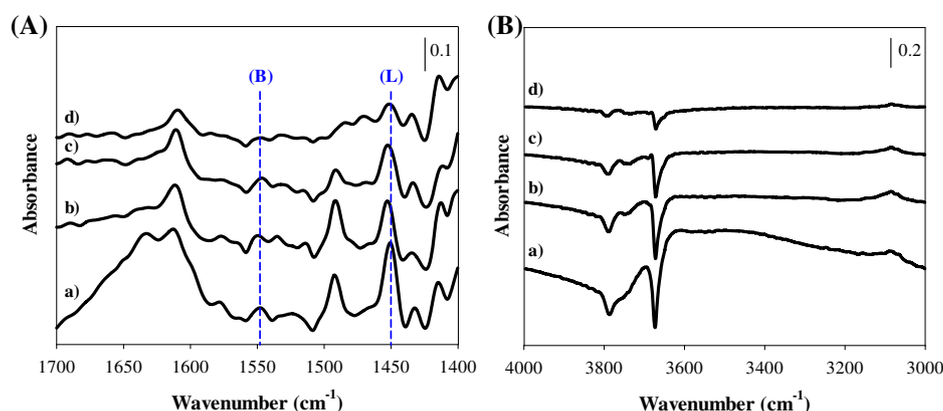


Figure 5. DRIFT spectra of adsorbed pyridine after degassing at four different temperatures (a) 50°C, (b) 150°C, (c) 200°C and (d) 300°C for 60 min for AlCoPO650 in the region (A) 1400–1700 cm $^{-1}$ and (B) 4000–3000 cm $^{-1}$.

the capacity to produce propene, and consequently the acidity, hardly changed, in clear agreement with the results obtained from Py-TPD. The formation of propanone was slightly favoured on the modified phosphates calcined at 450°C. As can also be seen in Table 2, the zeolites exhibited much higher activity than the aluminium phosphates. In fact, only the H-ZSM-5(50) zeolite with the highest SiO₂/Al₂O₃ ratio showed a similar value of conversion (1 mol%) at 170°C, also showing a value of selectivity to propanone (17%) similar to those exhibited by the modified aluminium phosphates calcined at 450°C. The presence of some basic sites in this zeolite has previously been reported.⁷

Glycerol transformation

The main products obtained under the experimental conditions studied were ACR and HA. Other by-products were glycerol formal (GF), glycidol (GD) and ethyl glycolate (EG). The possible routes of formation of these products has been discussed elsewhere.⁶ In addition to these, some unidentified products ('Unknown') were

detected, while phenol (P) was also obtained on some catalysts. CO₂ was never detected.

Catalyst activity as a function of time on stream

The glycerol conversion value increased considerably with the incorporation of a second metal to the AlPO₄, regardless of the method of synthesis and thermal treatment used, Table 3. Thus, the conversion values attained at 3 h of operation on modified phosphate calcined at 450°C, were double that of AlPO₄ (41%), those calcined at 650°C exhibiting higher values (around 100%). This increase in the activity of modified aluminium phosphates in relation to the aluminium phosphate cannot be explained based only on their acidity values, which hardly changed, as previously discussed, Table 2. We have to assume the participation of redox sites in the glycerol transformation, responsible for the propanone formation from 2-propanol, associated with the metal introduced in AlPO₄, in clear accordance with the results previously reported.⁶ Furthermore, the slight metal superficial enrichment

existing on the phosphates calcined at 650°C, in comparison with those calcined at 450°C, would contribute to enhancing the activity. However, the nature of the metal did not seem to play a decisive role in the activity of glycerol, contrary to what was indicated by Suprun^{30,31} and Liu.²⁸ The low percentage of metal employed in the present research, in comparison with that employed by these authors, could explain this fact.

Furthermore, none of the phosphates suffered deactivation after 22 h of operation, Table 3. At this time on stream, the conversion value of AlPO₄ was 76%, whereas those of all the modified phosphates were more than 90% (91–98%), with the exception of AlCuPO650 which exhibited a value of 85%. These changes in the conversion values with respect to those obtained at 3 h, were due to the differences in evolution of the conversion values with time on stream, as can be seen in Table 3. Thus, the conversion values on AlPO₄ appreciably increased from 3 h (41%) to 6 h (69%), remaining practically constant after 22 h (76%). This same tendency, although less pronounced, was observed for modified phosphates calcined at 450°C, whereas the values obtained on those calcined at 650°C hardly underwent any changes. These results could be based on the coke formation. In fact, all the catalysts changed from their natural colour to black after reaction. Furthermore, the coke formation should take place preferably at the initial stage of the reaction, as proved by the identical TGA profiles obtained for samples of phosphates after 3 h, 6 h and 22 h of reaction (Fig. 6). Therefore, the real conversion values at 3 h should be greater than those shown in Table 3 and, consequently similar to those obtained at 6 h, since the glycerol or reaction products transformed into coke were not taken into consideration. In fact, a higher formation of coke generally implies a lower conversion value at 3 h of reaction, the AlPO₄ exhibiting the highest amount of coke (242 mg_C/g_{cat}) and the modified phosphate calcined at 650°C the lowest (61 mg_C/g_{cat}), as shown in Table 3.

As can also be seen in Table 3, in contrast to the phosphates, the zeolites underwent a strong loss of activity with time on stream. Thus, after 6 h of operation the conversion values obtained for the zeolites decreased by half in relation to those at 3 h, showing a value around 15% at 22 h. These results were similar to those reported in the literature; both Chai⁸ and Kim³² used ZSM-5 at 315°C, losing around 70% and 60% of glycerol conversion, respectively, after 10 h of reaction. Furthermore, similarly to that obtained previously in liquid phase⁷ the zeolites were less active than the AlPO₄ and therefore than the modified AlPO₄, at 22 h of operation. However, at 3 h similar activity to AlPO₄ (around 45%) was exhibited, with the exception of H-ZSM-5(30), which showed a higher activity, in a likewise manner to the modified AlPO₄ calcined at 450°C.

The deactivation of the zeolites cannot be explained solely on the basis of the quantity of coke formed, similar to that obtained in the phosphates, but by the different role played by the coke on both types of solid. Thus, the drastic change of the textural properties of the used zeolites, Table 1, with a huge decrease in surface area and pore volume values, suggest that in the zeolites the coke acts by blocking the pores, mainly micropores, leading to the subsequent loss of acid sites, as shown in Table 2 and Fig. S1 in the Supplementary information. In contrast, the slight decrease in surface area, as well as in pore volume and size suffered by the used phosphates, suggests the formation of coke deposits inside the pores, acting as a poison. In this sense, Dalil *et al.*³³ have also reported that coke covered the internal walls of the pores rather than plugging the entrances. Although the number of acid sites decreased in a parallel manner with the loss of surface, the density

of the acid sites remained within the range exhibited by the fresh phosphates.

On the other hand, as a tendency the catalysts with a higher percentage of strong Brönsted acid sites produced a higher amount of coke, which would confirm once again, the participation of this type of acid site in the formation of coke, as indicated in the Introduction. Furthermore, the present results corroborate those previously postulated in liquid phase,⁷ in the sense that not only the strong acid sites have influence in the deactivation of the catalysts, promoting coke formation, but also the textural properties. In this context, the low conversion value shown by the HY(5.2) zeolite at 3 h operation, despite being the most active in the 2-propanol transformation, not only among the studied zeolites but also the phosphates, could be explained by a more extensive blockage of its pores by coke, possessing the highest micropore volume.

Catalyst selectivity as a function of time on stream

Another general result obtained in the catalytic performance of the phosphates as a function of time on stream, in addition to its high resistance to deactivation, was the change in the production of both main products, acrolein and hydroxyacetone. In effect, the yield to acrolein decreased and similarly, the yield to hydroxyacetone increased as the time of operation advanced, Table 3. Suprun *et al.*^{17,31} reported similar profiles of selectivity to both products over microporous silicoaluminophosphates, Al₂O₃ or TiO₂ modified with H₃PO₄ by impregnation and PO₄/Al₂O₃ modified with different transition metal oxides. The values of yield to acrolein at 3 h oscillated between 44% and 54%, and those to hydroxyacetone between 51% and 65% at 22 h on the modified aluminium phosphates, whereas the values on AlPO₄ were 20% and 37%, respectively. These results bring to light once more the positive effect of the modification of AlPO₄ with metals, as well as the little influence the type of metal has on the formation of either of the products. However, the systems calcined at 650°C and modified with Co and Cr are slightly more active in acrolein formation, while those modified with Cu are more active in that of hydroxyacetone. In this sense, catalysts based on Cu have been reported to be especially active in the formation of hydroxyacetone.^{28,31,34,35}

As can be seen in Table 3, the modified aluminium phosphates calcined at 650°C also promoted the formation of phenol, attaining yield values of 9 and 13%, which decreased appreciably with time on stream (final values of 3–4%). In contrast, the yield to 'Unknown' products increased with time on stream, AlPO₄ and modified AlPO₄ calcined at 450°C being the most active (yield values between 9 and 12% at 22 h of operation). In this same way, the formation of glycidol and ethylglycolate also increased, attaining the highest values, 10% and 5%, respectively, at 22 h on AlPO₄. With regard to glycerol formal, its formation did not seem to be affected by the presence of another metal, nor by the time on stream. In either case, the maximum yield obtained did not exceed 8%.

Such a notable change of the main reaction products, acrolein and hydroxyacetone, could be due to an interconversion of Lewis to Brönsted sites, as has previously been reported.^{8–10,20,36} In effect, the water presence in the feed at the reaction temperature, gives rise to a change in the equilibrium between Lewis and Brönsted sites, favouring the formation of Brönsted sites.²⁰ Thus, the Brönsted acid sites become predominant on the catalyst surface with time on stream, which could favour hydroxyacetone formation over acrolein. In fact, as shown in Fig. 7 the increase in the ratio HA/ACR with time of reaction was notably higher for AlPO₄ and AlFePO450, which initially exhibited a balanced percentage of

Table 3. Values of conversion and yield to reaction products at different times on stream, and amount of coke formed ($\text{mg}_C/\text{g}_{\text{cat}}$) after 22 h of operation. Experimental conditions: 36 wt% glycerol; $F_{\text{Gly}} = 0.3 \text{ mL h}^{-1}$; $F_{\text{N}_2} = 50 \text{ mL min}^{-1}$; $w = 0.05 \text{ g}$

Catalyst	X_{Gly} (mol%)			Y_{ACR} (mol%)			Y_{HA} (mol%)			Y_p (mol%)			Y_{Unknown} (mol%)			$\text{mg}_C/\text{g}_{\text{cat}}$
	3 h	6 h	22 h	3 h	6 h	22 h	3 h	6 h	22 h	3 h	6 h	22 h	3 h	6 h	22 h	
AlPO_4 450	41	69	76	20	16	8	9	25	37	1	2	2	6	8	9	242
AlCoPO 450	86	94	95	44	31	22	25	46	60	7	6	2	4	5	6	235
AlCuPO 650	100	100	96	54	42	30	29	44	55	13	8	4	4	4	4	66
Al-CoPO 450	87	86	94	47	30	22	22	37	61	5	4	2	5	7	5	95
Al-CoPO 650	92	90	91	50	26	16	26	45	61	9	6	3	5	10	7	177
AlCuPO 450	77	89	94	46	33	20	16	42	55	6	6	2	4	4	12	211
AlCuPO 650	98	98	85	49	33	22	33	49	51	7	4	2	4	6	4	69
AlCrPO 450	83	92	94	40	36	22	22	40	51	7	4	2	6	5	7	216
AlCrPO 650	100	100	98	53	38	28	28	42	58	13	7	4	5	10	5	61
AlFePO 450	99	99	93	52	37	14	29	46	65	5	5	2	9	9	9	144
H-ZSM-5(50)	43	15	13	16	5	3	16	6	5	0	0	0	2	1	2	75
H-ZSM-5(30)	84	39	17	44	14	3	22	14	8	0	0	0	4	3	2	67
HY(5.2)	48	24	17	18	7	4	23	14	10	0	0	0	2	1	1	142

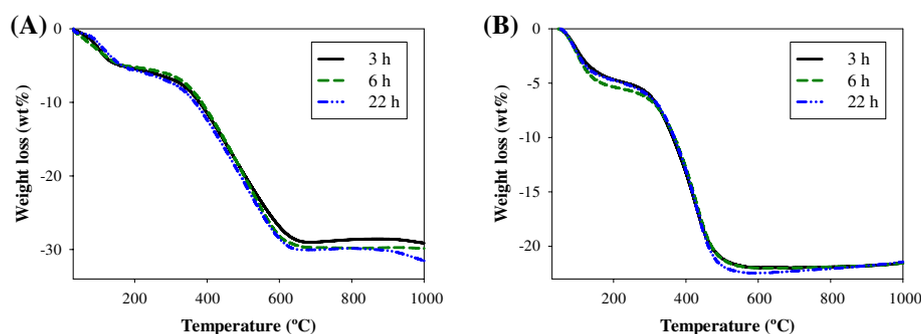


Figure 6. TGA analysis of the used (A) AlPO_4 450 and (B) AlFePO 450 at different times on stream.

Lewis and Brønsted acid sites, than for the rest of the phosphates in which the Lewis acid sites predominated. Therefore, the formation of hydroxyacetone preferably through an E_1 mechanism, Fig. 8, would be doubly favoured in relation to acrolein due to the existence of two primary hydroxyl groups in the glycerol molecule. At 3 h of time on stream, the Lewis sites were predominant and the glycerol dehydration might take place mainly through an E_2 mechanism, involving a pair of Lewis acid and base sites. In this mechanism, the adsorption of a glycerol molecule on a basic site through the hydrogen atoms bonded to primary carbons is twice as likely, Fig. 9, and consequently the formation of 3-hydroxypropanaldehyde, the precursor of acrolein, would be favoured. The transformation of 3-hydroxypropanaldehyde to the more stable aldehyde (acrolein), would easily occur with the participation of either a Brønsted or Lewis acid site. The participation of the acid (or base) sites caused by the presence of the metal transition, in addition to those related to aluminium, would explain the greater activity exhibited by the modified phosphates in respect to AlPO_4 .

Nature of coke

As Fig. 10 shows, the weight loss (Table 3) was accompanied in some solids by two exothermic peaks around 370°C and 510°C, in the differential thermal analysis (DTA) curves. These peaks are due to the combustion of coke on the used catalysts, as has been reported by other authors in the dehydration of glycerol.^{17,36–38} Furthermore, an endothermic peak also appeared around 115°C,

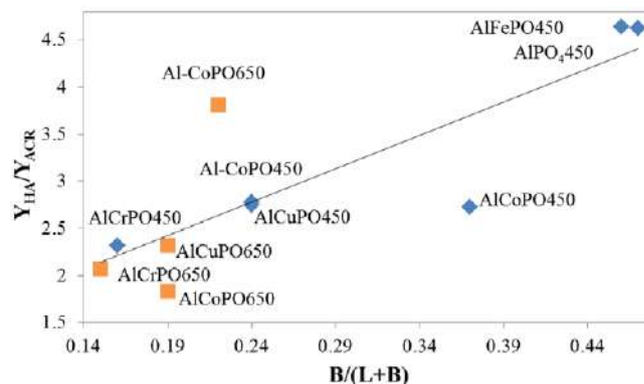


Figure 7. Influence of the nature of acid sites ($B/B + L$ after pyridine degassing at 150°C, Table 2) on the formation of hydroxyacetone and acrolein ($Y_{\text{HA}}/Y_{\text{ACR}}$ ratio at 22 h of time on stream, Table 3).

corresponding with the adsorbed water. The first exothermic peak could be assigned to the combustion of a highly hydrogenated carbonaceous species that are easily oxidized, such as aliphatic compounds, while the second peak would be related to the combustion of unsaturated or aromatic compounds, which are more deficient in hydrogen and therefore difficult to burn, as has been previously published.^{9,36,38} The weight loss of the first peak was considerably less than that corresponding to unsaturated or aromatic species. It was remarkable that both peaks appeared in

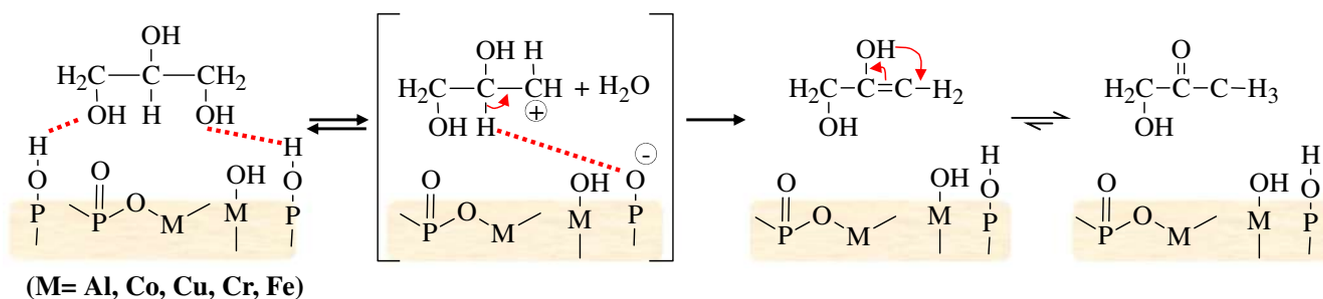


Figure 8. Proposed mechanism of hydroxyacetone formation on Brønsted acid sites.

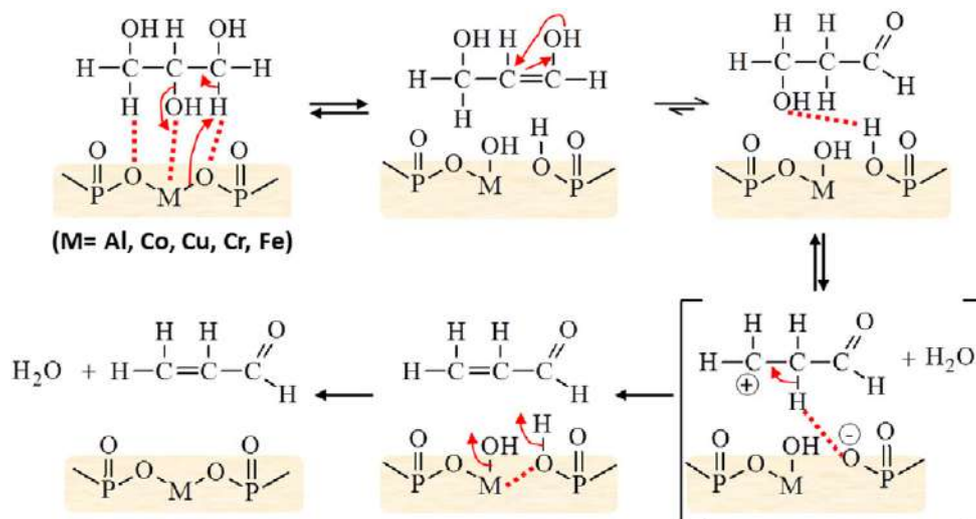


Figure 9. Proposed mechanism of acrolein formation on a pair of Lewis acid and base sites.

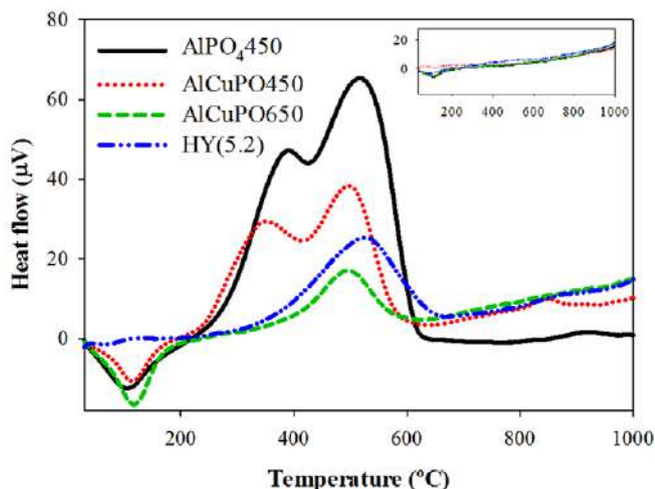


Figure 10. DTA profiles obtained for used catalysts. Those of the fresh catalysts appear in the top right-hand corner.

the solids calcined at 450°C, with the exception of the Al-CoPO450, while in the profiles corresponding to the phosphates calcined at 650°C, as well as those of the zeolites, only the peak at the highest temperature appeared.

The existence of saturated and aromatic compounds in the coke has been confirmed by Raman and IR spectroscopy, as shown in Fig. 11 for the used AlPO₄450. No noticeable XRD peaks representing crystalline carbon were observed for the used catalysts,

suggesting that the carbon deposits existed as amorphous carbon species on the catalyst surface; Raman spectra (Fig. 11(A)) showed the band for single-crystal graphite (at 1580 cm⁻¹) and the band for amorphous graphite (1350 cm⁻¹).¹² The band around 800 cm⁻¹ was related to the deformation vibration of C–H bond in aromatic compounds. This information is complemented by that obtained by IR spectra, Fig. 11(B), that were recorded after sample dilution to 15 wt% in KBr and equilibrated for at least 1 h at 280°C in a N₂ flow of 50 cm³ min⁻¹. Thus, the band that appeared at 1700 cm⁻¹ is assigned to C=O stretch, whereas those between 1400 and 1600 cm⁻¹ to aromatic C=C stretch. In the range 3000–2840 cm⁻¹ the bands appeared due to the vibration of the C–H bond of aliphatic compounds, and between 3080 and 3030 cm⁻¹ those due to C–H in aromatic compounds. The band in the region of the vibration of OH (3500–3800 cm⁻¹) assigned to isolated P–OH groups present in the spectra of the fresh solids, is transformed to a broad band in those of the spent catalysts, indicating H-bonded hydroxyl groups.

CONCLUSIONS

The presence of a second metal (1 wt% of Co, Cu, Cr, Fe) in the precipitation medium of AlPO₄ did not affect its amorphous character and mesoporosity even after being calcined at 450°C and 650°C.

The modified aluminium phosphates calcined at 650°C, exhibiting a superficial enrichment of metal, showed the highest activity (near 100%) in the glycerol transformation, regardless of the

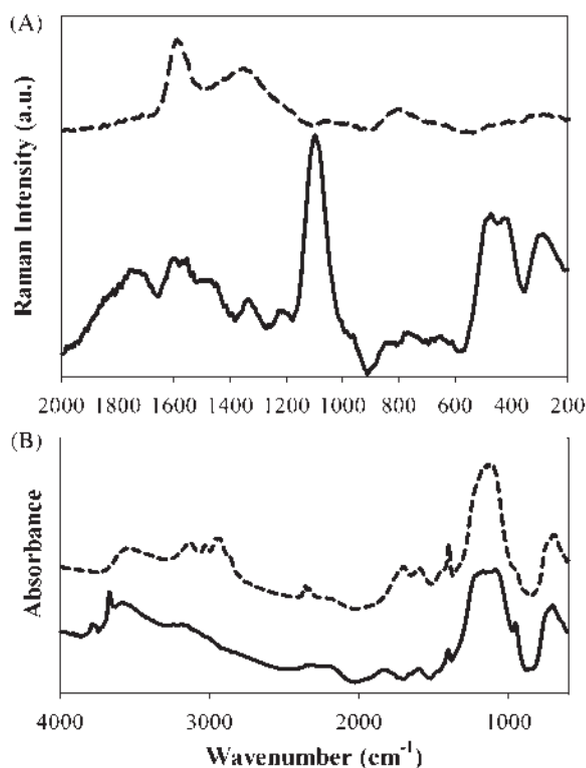


Figure 11. (A) Raman spectra and (B) DRIFT spectra of AlPO_4450 before (—) and after (---) 22 h on stream.

metal added, with a yield to acrolein oscillating between 49% (AlCuPO650) and 54% (AlCoPO650) at 3 h of time on stream, with 50 mg of catalyst and 280°C reaction temperature. The yield to the second main product, hydroxyacetone, oscillated between 28% (AlCrPO650) and 33% (AlCuPO650). Furthermore, a change in the formation of dehydration products with time on stream took place on all the phosphates. Thus, the yield to acrolein decreased and similarly the yield to hydroxyacetone increased, attaining values greater than 60% on phosphates modified with Fe and Co, at 22 h. This result could be explained by a change in the equilibrium between Lewis and Brønsted acid sites, favouring the formation of Brønsted sites, given the presence of water in the feed (36 wt% glycerol aqueous solution).

None of the phosphates suffered deactivation after 22 h of operation, whereas the zeolites underwent a strong loss of activity, caused by blockage of the micropores, which were predominant, by coke. The formation of coke at the beginning of the reaction, involving strong acid sites, preferably Brønsted acid, has also been verified. Unsaturated and aromatic compounds were the main components of coke.

We propose that on the phosphates studied, the acrolein formation from glycerol could take place preferably through an E_2 mechanism involving a pair of Lewis acid and base sites, whereas the hydroxyacetone formation could occur mainly through an E_1 mechanism with participation of the Brønsted acid sites. The participation of the Lewis acid (and base) sites associated with the presence of a transition metal, in addition to those related to aluminium, would explain the greater activity in the glycerol dehydration exhibited by the modified phosphates with respect to AlPO_4 , as well as their capacity for the propanone formation from 2-propanol.

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Supporting Information

Supporting information may be found in the online version of this article.

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