

# **Diols Production From Glycerol Over Pt-Based Catalysts: On the Role Played by the Acid Sites of the Support**

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Abstract A series of 1 wt% AuPt (6:4) catalysts were prepared by sol immobilization using acidic [TiO2, H-Mordenite, SiO<sub>2</sub>, MCM-41, sulphated ZrO<sub>2</sub> (S-ZrO<sub>2</sub>)] and one basic (MgO) oxide supports. A careful characterization of the catalyst was performed by HRTEM and FTIR spectroscopy. EDX analysis showed that on all catalysts only alloyed AuPt nanoparticles are present, but the final size of AuPt particles is significantly affected by the support. Indeed, on TiO<sub>2</sub> the mean AuPt nanoparticles diameter is 3.7 nm, whereas for all the remaining supports, larger AuPt nanoparticles with diameter of 6-7.5 nm were obtained. AuPt catalysts result very active in catalyzing the liquid phase hydrogenolysis of glycerol to 1,2-propandiol (68% conversion after 16 h) and 90% selectivity with ethylene glycol, 1-propanol and 2-propanol as main by-products. The role of the support has been highlighted in terms of acidic properties. The medium strength of Lewis acid sites of TiO<sub>2</sub> leads to the best performance in terms of activity, selectivity and stability of the catalytic system.

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# **Graphical Abstract**



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# **1** Introduction

The use of biomass for the production of renewable raw materials and their conversion to high value chemicals and materials show a significant potential [1-3]. The utilization of vegetable oil for the production of biodiesel has led to an increase in glycerol production, as it constitutes the main coproduct (about 20 wt%). Glycerol is a highly functionalized

molecule, which is recognized as a promising chemical building block for the synthesis of fine chemicals [1–6]. One attractive route involves the catalytic hydrogenolysis of glycerol to diols and alcohols (as reported in Scheme 1) [7–9].

1,2 Propandiol (1,2-PD), is commercially produced from propylene oxide and it is a valuable chemical for the production of polyesters, resins, and polyurethanes [10]. 1,3-PD is an important monomer utilized for the production of polyester fibers and films [11]. Ethylene glycol (EG) is produced via hydration of ethylene oxide and it is used as antifreezing agent [12].

In the past decade, glycerol hydrogenolysis has been extensively studies using different metal based catalysts, including Pt, Ru, Ni and Cu [7–9]. Among all noble metal based catalysts, Pt has been recognized as one of the most effective in the glycerol hydrogenolysis [7–9, 13–15]. For example, commercial Pt/C showed high selectivity to 1,2-PD (82.7%) but only at low conversion (35%) [16]. Similar results were obtained using PtSn/SiO<sub>2</sub> with a good selectivity to 1,2-PD (84%) but even at lower conversion (16%) [17]. Nevertheless, the drawback of most of the monometallic Pt systems seems to be the low resistance to deactivation and the low capacity to maintain high selectivity at high conversion. We recently reported that the addition of Au to Ru improves the stability of the catalysts, maintaining a good selectivity also at high conversion [18]. Moreover, the beneficial effect of Au addition to Pt catalysts in terms of activity, selectivity and stability was already verified in the case of oxidation of glycerol [19, 20]. Recent papers illustrated that the nature of the support can significantly affect the catalytic performance [21, 22]. In particular, the acid base properties have been reported to modify the activity and, primarily, the selectivity in the glycerol oxidation [20]. Therefore, this study encompasses the investigation of the effect of the acid base properties of the support in AuPt catalyzed glycerol hydrogenolysis to 1,2 PD.

Many mechanisms corresponding to glycerol hydrogenolysis to 1,2 PD, in which the acidic/basic properties seem to have a key role in determining the final pathway, have been proposed. Three main pathways (Scheme 2) are generally accepted: dehydration-hydrogenation, dehydrogenation-dehydration-hydrogenation and direct-hydrogenolysis [9].

In the first path, glycerol is initially dehydrated through an acid-catalyzed reaction to form an intermediate, which is subsequently hydrogenated to generate propanediol [23]. Following the second possible path, which usually occurs in neutral and alkaline conditions, glycerol is dehydrogenated to glyceraldehyde with a consecutive dehydration-hydrogenation to 1,2-PD [24]. The last possibility is the directhydrogenolysis that was initially reported for Ir catalysts [25]: glycerol is adsorbed on the metal surface to form 1,3 dihydroxyisopropoxide that is successively transformed to 1,2-PD. According to this mechanism, the low stability of the intermediate dihydroxyisopropoxide can result in the formation of over-hydrogenolysis products (1-propanol and 2-propanol). In all the possible pathways, the role of the support properties appears fundamental and yet not fully understood.



Scheme 1 Products obtained by glycerol hydrogenolysis



Scheme 2 Proposed reaction pathways of glycerol hydrogenolysis to 1,2-PD

Therefore, to clarify the possible synergy between the acid base properties of the support and AuPt nanoparticles, different catalysts have been prepared supporting preformed AuPt NPs on acidic (TiO<sub>2</sub>, H-Mordenite, SiO<sub>2</sub>, MCM-41, S-ZrO<sub>2</sub>) and basic (MgO) oxides. High-resolution transmission electron microscopy (HRTEM) analysis of the AuPt catalysts was performed to investigate the morphology as well as the composition of the nanoparticles. The nature (Lewis and/or Brønsted) and acid site density were determined by a quantitative analysis of the FTIR bands of adsorbed 2,6 dimethyl-pyridine (2,6-DMP). The catalysts have been then tested in glycerol hydrogenolysis.

# 2 Experimental

#### 2.1 Materials

NaAuCl<sub>4</sub>·2H<sub>2</sub>O, and K<sub>2</sub>PtCl<sub>4</sub> were from Aldrich (99.99% purity), TiO<sub>2</sub> P25 (SA = 48 m<sup>2</sup>/g) was from Evonik and H-Mordenite (SA = 450 m<sup>2</sup>/g) was from Degussa. Sulphated zirconia (SA = 78 m<sup>2</sup>/g), SiO<sub>2</sub> (SA = 148 m<sup>2</sup>/g) and MgO (SA = 38 m<sup>2</sup>/g) were from Alfa Aesar. MCM-41 (SA = 980 m<sup>2</sup>/g) have been prepared following the procedure reported in Mori et al., [26]. NaBH<sub>4</sub> of purity >96% from Fluka, polyvinylalcohol (PVA) (Mw = 13,000–23,000, 87–89% hydrolyzed), from Aldrich were used. A 1 wt% PVA solution in water was prepared. Gaseous oxygen from SIAD was 99.99% pure.

### 2.2 Catalyst Preparation

#### 2.2.1 Monometallic Pt Catalyst

Solid  $K_2$ PtCl<sub>4</sub> (0.051 mmol) and PVA (1 wt%) solution (Pt/ PVA 1/0.5 wt%) were added to 100 mL of H<sub>2</sub>O. After 5 min, H<sub>2</sub> was bubbled (50 mL/min) under atmospheric pressure and room temperature for 2 h. The colloid was immobilized by adding the support under vigorous stirring. The amount of support was calculated as having a total final metal loading of 1 wt%. After 2 h the slurry was filtered, the catalyst washed thoroughly with distilled water (neutral mother liquors) and dried at 80 °C for 4 h.

## 2.2.2 Au-Pt Bimetallic Catalysts

For the sake of clarity, the main steps of the preparation of the bimetallic catalysts are summarized in Scheme 3. The bimetallic AuPt catalysts were prepared by sol immobilization through a two steps procedure, which ensures the formation of alloyed bimetallic particle [19, 20].

In detail, NaAuCl<sub>4</sub>·2H<sub>2</sub>O (Au: 0.031 mmol) was dissolved in 60 mL of H<sub>2</sub>O, and PVA (1%, wt%) was added (Au/PVA = 1:0.5 wt/wt). The yellow solution was stirred for 3 min, after which 0.1 M NaBH<sub>4</sub> (Au/NaBH<sub>4</sub> = 1:4 mol/mol) was added under vigorous magnetic stirring. The ruby-red Au(0) sol was formed immediately. Within a few minutes of sol generation, the gold sol was immobilized by adding the support (acidified to pH 2 by sulphuric acid) under vigorous stirring. The amount of support was calculated as having a gold loading of 0.60 wt%. After 2 h, the slurry was filtered and the catalyst washed thoroughly with distilled water (neutral mother liquors). The Au/support was dispersed in 40 mL of water, therefore  $K_2PtCl_4$  (Pt: 0.021 mmol) and PVA solution (Pt/PVA = 1:0.5 wt/wt) were added. H<sub>2</sub> was bubbled (50 mL/min) under atmospheric pressure and room temperature for 2 h. After an additional 18 h, the slurry was filtered and the catalyst washed thoroughly with distilled water. ICP analyses were performed on the filtrate using a Jobin Yvon JV24 instrument to verify the metal loading on the support. The total metal loading was 1 wt%. AuPt nanoparticles were supported on five acidic oxides, i.e. TiO<sub>2</sub>, H-Mordenite, SiO<sub>2</sub>, MCM-41, S-ZrO<sub>2</sub>, and a basic support (MgO). The obtained catalysts are listed in Table 1.

#### 2.3 Catalytic Tests

Glycerol hydrogenolysis was performed at 150 °C, using a stainless steel Parr reactor (50 mL capacity), equipped with heater, mechanical stirrer, gas supply system and thermometer. The glycerol solution (30 mL; 0.3 M) was added into the reactor and the desired amount of catalyst (glycerol/metal ratio = 1000, mol/mol) was suspended in the solution. The



Scheme 3 Synthesis of the bimetallic catalysts

**Table 1**Catalytic evaluation inthe glycerol hydrogenolysis

Catalyst	$\frac{\text{Activity}^{a}}{\binom{(h^{-1})}{\text{mol} \times h}}$	$\frac{Activity \ A^b}{\frac{mol}{N \times h}}$	Selectivity (%) <sup>c</sup>					AuPt size
			1,2-PD	1-PrOH	2-PrOH	EG	MeOH	(nm)
AuPt/MgO	34	176	93	_	_	3	2	6.9
AuPt/TiO <sub>2</sub>	56	165	90	-	-	4	2	3.7
AuPt/MCM-41	22	111	52	-	-	22	16	6.7
AuPt/SiO <sub>2</sub>	21	117	35	-	-	28	22	7.5
AuPt/H-Mordenite	14	65	30	8	12	20	16	6.2
AuPt/S-ZrO2	10	52	31	10	11	18	15	6.9
Pt/TiO <sub>2</sub>	41	119	86	-	-	3	4	3.6
Au/TiO <sub>2</sub>	1	2	-	-	-	-	-	3.5

Reaction conditions: glycerol = 0.3M, pH<sub>2</sub> = 7 bar, T = 150 °C, metal/glycerol = 1/1000 mol/mol. *EG* ethylene glycol, *1,2-PD* 1,2 propandiol, *1-PrOH* 1-propanol. The carbon balance around 90–96% observed in all the reactions was ascribed to the formation of CH<sub>4</sub> and CO<sub>2</sub> not detectable in liquid phase

<sup>a</sup>Mol of glycerol converted per hour per mol of metal, calculated after 0.5 h of reaction

<sup>b</sup>Mol of glycerol converted per hour per fraction of atoms lying at the surface, A  $[A = (Ns/NT) \times 100)$  after 0.5 h of reaction

<sup>c</sup>Selectivity at 60% of conversion

autoclave was purged three times with nitrogen before charging 10 bar of  $H_2$ . The mixture was heated to the reaction temperature, 150 °C, and mechanically stirred (1250 rpm).

Products analysis: the reaction mixture, after separation from the catalysts by filtration, was analysed using high performance liquid chromatography (HPLC). Samples were removed periodically (0.5 mL) under stirring and analysed by HPLC using a column (Alltech OA-10308 300 mm  $\times$  7.8 mm) with UV and refractive index (RI) detection in order to analyse the product mixtures. H<sub>3</sub>PO<sub>4</sub> 0.1 wt% solution was used as the eluent. The identification of the possible products was done by comparison with the original samples. Activity was calculated on the total mol of metal and, alternatively, on the total amount of exposed surface atoms. Calculations of the number of exposed surface atoms were performed by assuming that all the nanoparticles had cuboctahedral morphology with cubic close-packed structure in this size range, the model of full-shell nanoparticles was adopted [27]. The total number ( $N_t$ ) of M atoms in the cluster for a given cluster size can be calculated using the following Eq. (1):

$$d_{\rm sph} = 1.105 \, d_{\rm at} N_{\rm T}^{-1/3} \tag{1}$$

where  $d_{sph}$  is the mean diameter of the Au, Pt or AuPt particles obtained from transmission electron microscopy (TEM) analysis and  $d_{at}$  is the atom diameter of Au (0.288 nm), Pt (0.256 nm) and AuPt (0.264 nm), respectively. The number of surface atoms (N<sub>s</sub>) and n can be calculated from Eqs. (2) and (3), based on the values of N<sub>T</sub>:

$$N_{\rm T} = \left(10n^3 - 15n^2 + 11n - 3\right) / 3 \tag{2}$$

$$N_s = 10n^2 - 20n + 12 \tag{3}$$

The activity based on the surface atoms can then be calculated as follows:

fraction of atoms lying at the surface:

 $A = (N_s/N_T) \times 100$ 

Activity based on  $N_s$  = activity (calculated for total metal mols)/A.

Recycling tests were carried out under the same experimental conditions. The catalyst was recycled in the subsequent run after filtration without any further treatment.

## 2.4 Characterisation

- a. The metal content was checked by Atomic Absorption Spectroscopy (AAS) analysis of the filtrate, on a Perkin Elmer 3100 instrument.
- b. Samples for TEM characterization were prepared by depositing an ethanol suspension of the catalyst onto lacey carbon coated 300 mesh copper grids. The particle morphology of the supported nanoparticles, specifically the nanoparticle size, was first investigated by TEM using a Philips LaB<sub>6</sub> electron microscope, operating at 200 kV and equipped with a Gatan CCD camera. Detailed high resolution High Angle Annular Dark Field Scanning TEM (HAADF-STEM) imaging and energydispersive X-ray spectroscopy (XEDS) analyses were carried out using FEI Titan3 microscope operated at 200 kV accelerating voltage for a deeper investigation of the AuPt alloy structure of AuPt/TiO<sub>2</sub> catalyst. This microscope is equipped with double aberration correctors, providing ultrahigh-resolution HAADF-STEM images, and an Oxford Inca energy dispersive X-ray (EDX) spectrometer equipped with a 30 mm<sup>2</sup> ultrathin window Si/Li X-ray detector. XEDS data were collected either as spectrum images, in which a focused electron probe was scanned across a region of interest during data collection, or in stationary spot mode, where an emitted X-ray spectrum from 0 to 20 kV is acquired from a specific point on a particle using a probe size <0.5 nm. Spectra were acquired with a probe current of approximately 0.5 nA and dwell times between 200 and 400 ms/pixel, in the case of maps, and 20-30 s per analysis in spot mode. STEM digital images were

acquired using FEI TIA software and Oxford INCA microanalysis software was used for XEDS acquisition and analysis. The atomic fractions of gold and platinum were quantified by the Cliff–Lorimer method on relative intensities of the Pt-L $\alpha$  and Au-L $\alpha$  peaks using k-factors provided by the XEDS system manufacturer. The AuPt particle size distribution and the total metal dispersion were determined by counting 250 particles in HAADF-STEM images using GAUSS software.

FTIR spectra were taken on a Perkin-Elmer 2000 specc. trometer (equipped with a MCT detector) with the samples in self supporting pellets introduced in cells allowing thermal treatments in controlled atmospheres and spectrum scanning at room temperature (r.t.) in vacuum or in the presence of probe gases. The 2,6-dimethylpyridine (2,6-DMP) adsorption/desorption experiments were performed on the samples after activation in vacuum at 393 K for 1 h. The following procedure was adopted: (1) inlet of an excess dose of 2,6-DMP vapor (~2 Torr), and equilibration at r.t. for 10 min; (2) evacuation at r.t. for 30 min. From each spectrum, the spectrum of the sample before the inlet of the probe was subtracted. The spectra have been normalised with respect to the weight of the pellets and to the surface areas of the different supports.

The processing of the FTIR spectra of adsorbed 2,6-DMP was performed by using Fityk 0.9.8, an open-source curvefitting and data analysis software, that allows to obtain accurate curvefits (either Gaussian or Lorentzian curves were used). As output files, the software permits the realisation of one figure each time. Each obtained figure contains the files related to one specific sample in the optimised spectral range, therefore it was not possible to plot all files in the same figure.

# **3** Results and Discussion

The catalysts were evaluated in glycerol hydrogenolysis (0.3 M glycerol, glycerol/metal = 1000 mol/mol, 10 atm. H<sub>2</sub>, T = 150 °C) and both activity and selectivity, at isoconversion (60%) are reported for each system in Table 1.

The activity was calculated firstly basing on the moles of glycerol converted per hour per total mol of metal. By using this approach, AuPt/TiO<sub>2</sub> showed a high activity (56 converted mol of glycerol (AuPt mol)/h), superior than AuPt supported on MgO (34 converted mol of glycerol (AuPt mol)/h) and the other acidic oxides (<22 mol of glycerol (AuPt mol)/h for AuPt/MCM41, AuPt/SiO<sub>2</sub>, AuPt/H-Mordenite, AuPt/S-ZrO<sub>2</sub>, accordingly). Moreover, monometallic Pt/TiO<sub>2</sub> and Au/TiO<sub>2</sub> were also tested as reference catalysts: it was found that monometallic Pt showed intermediate



Fig. 1 Reaction profile of AuPt catalysts



Fig. 2 Stability test using AuPt/TiO<sub>2</sub>

activity (41 converted mol of glycerol (Pt mol)/h), whereas monometallic Au was inactive (1 converted mol of glycerol (Au mol)/h). These findings indicate that Au has a synergist effect on the activity of Pt, maximized in the case of  $\text{TiO}_2$  support.

In addition, the comparison of the reaction profile of the two most active catalysts put in evidence that  $AuPt/TiO_2$  showed a better resistance to deactivation than AuPt/MgO (Fig. 1). Interestingly, AAS analysis revealed the presence of Au, Pt and Mg in the reaction media as for AuPt/MgO, whilst AuPt/TiO<sub>2</sub> did not show any leaching. Moreover, the performance of bimetallic AuPt compared to monometallic Pt on TiO<sub>2</sub> (Table 1) showed not only the best activity of the bimetallic sample but also its best resistance to deactivation (Fig. 1). To confirm the stability of AuPt/TiO<sub>2</sub>, recycling tests have been performed (Fig. 2). Recycling experiments

were carried out by filtering and using the catalyst in the next run without any further purification. The catalyst was stable for six run. Therefore, these results demonstrate that when  $TiO_2$  is used as support the addition of Au to Pt is beneficial in terms either of activity or of stability in glycerol hydrogenolysis, similarly as previously observed for glycerol oxidation [19, 20]. Similar findings were achieved by some of us when Au was added to Ru supported on active carbon (AC), with AuRu/AC displaying an higher activity than Ru/AC in glycerol hydrogenolysis to 1,2-PD [18].

As for the obtained selectivity, also reported in Table 1, it was found that AuPt supported on MgO and TiO<sub>2</sub> were extremely selective toward the formation of 1,2-PD (93 and 90%, respectively. All the other catalysts showed lower selectivity to 1,2-PD, 30–50%). In particular, these catalysts produced high amounts of EG and methanol deriving from the C–C cleavage (see Scheme 1; Table 1). Moreover, 1-propanol and 2-propanol, deriving from the over-hydrogenolysis of 1,2-PD (Scheme 1, Table 1) were produced in the presence of the AuPt/H-Mordenite and AuPt/S-ZrO<sub>2</sub> catalysts.

A combined HRTEM and FTIR characterization was considered out to unravel on one hand, the influence of both AuPt morphology and size and, on the other hand, to focus on the effect of the nature of the support on the catalytic activity and selectivity. For the latter reason, due to the low stability of MgO based catalyst during the reaction, the attention will be entirely focused on the acid catalysts which all presented a pretty good stability, as shown for TiO<sub>2</sub>.

A detailed HRTEM characterization of the bimetallic AuPt catalysts has been previously reported by Chan-Thaw et al. [28] for AuPt/TiO<sub>2</sub> and by Villa et al. [20] for all the other catalysts. In all cases, TEM images revealed a good AuPt dispersion and EDS analysis showed the exclusive presence of AuPt alloyed nanoparticles [20, 28]. HAADF images of AuPt/TiO<sub>2</sub> catalyst are shown in Fig. 3 for the sake of clarity. A representative STEM-HAADF image of AuPt/ TiO<sub>2</sub> and the corresponding XEDS elemental analysis are also reported in Fig. 4a, b, respectively. It is clearly shown that both Pt and Au are present in the same nanoparticle. Moreover, the distribution profiles of the metals shown in Fig. 4c confirmed the coexistence of Pt and Au within the same nanoparticle, suggesting the formation of a random gold-platinum alloy structure.

However, despite the same alloy nanostructure, the AuPt bimetallic nanoparticles differ in term of size, as reported in Table 1. Indeed, AuPt/TiO<sub>2</sub> average diameter is 3.7 nm, whereas for all the remaining supports larger AuPt nanoparticles, with mean diameter of about 6–7.5 nm, were observed. The two metals have been deposited on the different supports by following the same synthetic procedure summarized in Scheme 3: it is evident that the main difference in the particle size was observed when Pt and Au were supported on TiO<sub>2</sub>. Therefore, in the case of AuPt/



Fig. 3 HAADF images of AuPt/TiO<sub>2</sub>. Images a, b, show the presence of nanoparticles ca. 3.7 nm in size



**Fig. 4** a STEM-HAADF image and **b** the corresponding XEDS spectra taken from the AuPt nanoparticle. **c** The corresponding Au L $\alpha$  and Pt L $\alpha$  linescan profile taken across the diameter of the same AuPt nanoparticle (*dashed orange arrow* in **a**)

TiO<sub>2</sub>, the support played an active role during the metal insertion. Indeed, a combined scanning tunnelling microscopy and density functional theory study demonstrated that bridging oxygen vacancies are the active nucleation sites

for Au clusters on the rutile  $\text{TiO}_2$  (110) surface [29]. In particular, by monitoring the temperature dependence of the cluster size distribution and the oxygen vacancy density, it was found that a single Au atom-vacancy complex is stable up to room temperature and a single oxygen vacancy can bind three Au atoms on average. For larger clusters, the Au-substrate interface contains a high density of oxygen vacancies, which enhances the binding of Au particles to the substrate. A synthesis model involving the gold nucleation and growth on the oxygen vacancies of titania during the preparation (Scheme 3) can be proposed. These preformed nanoparticles will act themselves as nucleation sites for the Pt precursor, according to the EDS findings showing exclusively the presence of AuPt alloyed nanoparticles within the whole catalysts [20, 28].

It is well known that smaller nanoparticles are more active than larger ones in liquid phase reactions, probably due to the higher amount of exposed atoms [1-3]. Keeping this observation in mind, in order to rule out the effect of the particle size and to focus on the support effect, the activity was calculated by normalizing with respect to the fraction of exposed atoms (Table 1). In that case, the most active catalyst was AuPt/MgO, in agreement with what reported in the literature, i.e. that metal nanoparticles on basic supports are more active than on acid ones for this reaction [7-9, 30]. The AuPt/MgO catalyst resulted also very selective toward the formation of 1,2-PD (93%) with a small amount of EG and methanol produced (Table 1). Under basic conditions, it was reported that the formation of 1.2-PD proceeded via dehydrogenation of glycerol to glyceraldehyde as firs step (Scheme 2) [7, 8]. The limited formation of EG can be due to the difficulty for the retro-aldol reaction of glyceraldehyde to take place under alkaline conditions [7].

To be noted that the value obtained for AuPt/TiO<sub>2</sub> is very closed to the one of AuPt/MgO (Table 1).

The normalized activity seems to decrease by increasing the support activity. Therefore, to elucidate the effect of the nature of the support on the catalytic performance, a detailed characterization of the surface properties of the catalysts has been performed, in particular by focusing on the acid supports.

The normalised FTIR spectra collected on the different catalysts exposed to 2 mbar 2,6-DMP at r.t. and after 30 min outgassing at the same temperature are shown in the wavenumber range 1500–1700 cm<sup>-1</sup>, (dotted curves in Figs. 5, 6, 7, 8, 9). It is worth noting that the use of 2,6-DMP allowed to characterise both Lewis and Brønsted acidic centers present at the surface and offered the advantage to distinguish among Lewis and/or Brønsted sites of different acidic strength [31, 32].

The former species, together with H-bonded species, gave rise to three bands in the  $1620-1580 \text{ cm}^{-1}$  range, where the 8a and 8b ring stretching modes can be observed [33]. The presence of the latter species, i.e. Brønsted sites, was revealed by the formation of the 2,6-dimethylpyridinium species (2,6-DMPH<sup>+</sup>), that presented the 8a-8b bands of the ring stretching modes, at  $1640-1655 \text{ cm}^{-1}$  and about



Fig. 5 FTIR difference spectrum reported in the spectral region of 8a–8b ring modes of 2,6-DMP collected exposed to 2,6-DMP (2 mbar) at r.t. on AuPt/MCM-41 (*blue dotted curve*), curvefit of the spectrum (*orange curve*) and deconvoluted bands (*grey curves*)



**Fig. 6** FTIR difference spectrum reported in the spectral region of 8a–8b ring modes of 2,6-DMP collected after the inlet of 2,6-DMP (2 mbar) at r.t. on AuPt/SiO<sub>2</sub> (*blue dotted curve*), curvefit of the spectrum (*orange curve*) and deconvoluted bands (*grey curves*)

1630 cm<sup>-1</sup>, respectively. A careful deconvolution of the normalised spectra collected upon interaction with 2,6-DMP and after outgassing at r.t. for 30 min was carried out (see Figs. 5, 6, 7, 8, 9) and the calculated integrated areas of the produced absorption bands (normalized to the weight of the pellets and to the SSA) are reported in Table 2. This allowed us to compare the obtained quantitative information about the different acidic sites present on the catalysts and to afford structure-activity correlations with the catalytic results reported in Table 1.

Complex bands related to liquid like and H-bonded 2,6-DMP were produced upon 2,6-DMP adsorption at r.t. (see Figs. 5, 6, 7, 8, 9). Therefore, the 2,6-DMP interaction was given mainly via H bonding with the OH groups exposed at the surface of the samples, suggesting that OH groups with weak acid strength play a role in the C–C cleavage to produce EG and methanol (see Tables 1, 2). This is not true Fig. 7 FTIR difference spectrum reported in the spectral region of 8a–8b ring modes of 2,6-DMP collected after the inlet of 2,6-DMP (2 mbar) at r.t. and *inset* after 30 min outgassing, on AuPt/S-ZrO<sub>2</sub> (*blue dotted curves*), curvefits of the spectra (*orange curves*) and deconvoluted bands (*grey curves*)





Fig. 8 FTIR difference spectrum reported in the spectral region of 8a–8b ring modes of 2,6-DMP collected after the inlet of 2,6-DMP (2 mbar) at r.t. and *inset* after 30 min outgassing, on AuPt/H-Mordenite (*blue dotted curves*), curvefits of the spectra (*orange curves*) and deconvoluted bands (*grey curves*)

in the case of AuPt/TiO<sub>2</sub>, which exposed almost exclusively Lewis sites, according to the residual intensities of the bands after outgassing (given by the integrated areas in Table 2).

No Lewis and Brønsted acidity was observed in the case of AuPt/MCM-41 and AuPt/SiO<sub>2</sub> (the total normalised area of the bands related to H-bonded and physisorbed species was 12.783 and 18.91, respectively). The higher stability to the outgassing shown by the bands related to AuPt/MCM-41, with respect to those detected on AuPt/SiO<sub>2</sub>, was due to the presence of mesopores, where the H-bonded 2,6-DMP can be removed only by prolonging the outgassing time (data not shown for the sake of brevity). On the contrary, both Lewis and Brønsted acid sites were observed on AuPt/S-ZrO<sub>2</sub> (total normalised areas 1.831 and 8.368, respectively) and AuPt/TiO<sub>2</sub> (total normalised areas 4.401 and 8.472, respectively), whereas AuPt/H-Mordenite showed almost exclusively Brønsted acidity (2.38). Interestingly, as revealed by the deconvolution of the spectra collected after outgassing for 30 min at r.t., the amount of Lewis sites present on AuPt/TiO<sub>2</sub> was higher than that obtained for AuPt/S-ZrO<sub>2</sub>, where the sulphate groups are occupying these sites. Moreover, the apparent absence of Lewis sites on the AuPt/H-Mordenite catalyst was ascribed to the presence of residual water Fig. 9 FTIR difference spectrum reported in the spectral region of 8a–8b ring modes of 2,6-DMP collected after the inlet of 2,6-DMP (2 mbar) at r.t. and *inset* after 30 min outgassing, on AuPt/TiO<sub>2</sub> (*blue dotted curves*), curvefits of the spectra (*orange curves*) and deconvoluted bands (*grey curves*)



Table 2 Position and integrated areas of the bands observed upon 2,6-DMP dosage at r.t

	Band integrated areas of adsorbed 2,6-DMP									
	Brønsted acid sites			H-bonded	Physisorbed	H-bonded and physisorbed				
Position (cm <sup>-1</sup> )	1648	1640	1632	1603	1595	1581				
AuPt/MCM-41				2.404	5.256	5.123				
AuPt/SiO <sub>2</sub>				5.169	2.022	11.719				
	Brønsted acid sites			Lewis acid sites and Physisorbed H-bonded		H-bonded and physisorbed				
AuPt/S-ZrO <sub>2</sub> <sup>a</sup>	3.480	3.186	1.702	2.384	2.318	6.609				
	6.444		5.832	1.046		0.785				
AuPt/TiO <sub>2</sub> <sup>a</sup>	3.621	3.808	1.043	5.318	7.073	14.738				
	0.377	2.111	0.429	2.443	0.632	1.326				
AuPt/H-Mordenite <sup>a</sup>	0.365	0.107	1.908	0.9286	0.200	1.616				
	1.334		1.688							

<sup>a</sup>The deconvolution was performed also on the spectra collected after 30 min outgassing at r.t. (refer to the second line for each catalyst)

molecules adsorbed on H-Mordenite activated by simply outgassing at 120 °C for 1 h.

On the basis of the FTIR results, the low selectivity to 1,2-PD displayed by both AuPt/S-ZrO<sub>2</sub> and AuPt/H-Mordenite it has been related to the presence of Brønsted sites, that are able to give over-hydrogenolysis of 1,2-PD, forming 1-propanol and 2-propanol. An high 1,3-PD selectivity (48.6%) in the presence of a 2 wt% Pt/H-Mordenite catalyst at 94.9% glycerol conversion was also reported [21]. The selectivity to 1,3-PD was influenced by the Pt dispersion

(around 4 nm) and by the Brønsted acidity of the support. In our case, for the AuPt/H-Mordenite catalyst, an effect on the selectivity to 1,2-PD due to the additional presence of gold, which acts on the electronic properties of Pt, can be inferred.

Interestingly, Brønsted sites are also present at the surface of AuPt/TiO<sub>2</sub>, i.e. the most active catalyst (according to Table 1). Our results gave precise indication that despite the amount of these sites is comparable to that detected on AuPt/S-ZrO<sub>2</sub>, and higher than that observed for AuPt/H-Mordenite, the behaviour towards the outgassing at r.t. is

completely different. Indeed, on both AuPt/S-ZrO<sub>2</sub> (Fig. 7) and AuPt/H-Mordenite (Fig. 8) the amount of Brønsted sites is enhanced after outgassing (the integrated area values move from 8.368 up to 12.276 and from 2.38 up to 3.022, respectively) indicating that at low coverages a rearrangement of the 2,6-DMP molecules on the most acidic sites has occurred. Differently, a decrease in intensity of the bands related to the Brønsted sites is observed in the case of AuPt/TiO<sub>2</sub> (from 8.472 down to 2.917, see Fig. 9), giving evidence of the presence of less strong Brønsted sites on this catalyst.

On the contrary, the FTIR spectroscopic characterisation indicated that the superior activity as well as the high selectivity displayed by AuPt/TiO<sub>2</sub> can be related to the presence of Lewis sites with moderate strength. More acidic supports present a lower activity. Similar results were reported in the literature by Guo et al. [34]. Indeed, they reported that Cu immobilized on alumina with moderate acid sites is more active than more acidic zeolite supports.

## **4** Conclusion

Bimetallic AuPt catalysts were tested in the glycerol hydrogenolysis, using supports with different acid–base properties. It was found that the smallest AuPt nanoparticles (with average diameter equal to 3.7 nm) showed a better activity than largest ones (6–7 nm).

However, by normalizing the activity (conversion) on the number of exposed atoms, it appears clearly that the activity and the selectivity were significantly influenced by the acid base properties of the support. As expected, AuPt on MgO appears the most active catalyst, despite it deactivates quite rapidly. The activity in the case of acidic supports follows the order  $(AuPt/TiO_2) > (AuPt/MCM41 \ge AuPt/$  $SiO_2$  > (AuPt/H-Mordenite > AuPt/S-ZrO<sub>2</sub>). The observed trend can be directly related to the acid character of the support: the higher the strength of the acidic sites, the lower the activity. In particular, the presence of Lewis acid sites is required for achieving good catalytic performances. Moreover, it was shown that the nature of the acidic sites plays also an important role in tailoring the selectivity, by lowering the production of 1,2-PD and by directing the glycerol hydrogenolysis reaction to EG and methanol by C-C cleavage and/ or 1-propanol and 2-propanol by over-hydrogenolysis of 1,2-PD. Finally, it has been shown that the intermediate acidic character of TiO<sub>2</sub> leads to an improved selectivity towards 1,2-PD.

## References

- 1. Gallezot P (2012) Chem Soc Rev 41:1538-1558
- 2. Huber GW, Iborra S, Corma A (2006) Chem Rev 106:4044-4098

- 3. Corma A, Iborra S, Velty A (2007) Chem Rev 107:2411-2502
- Katryniok B, Kimura H, Skrzyńska E, Girardon J-S, Fongarland P, Capron M, Ducoulombier R, Mimura N, Paul S, Dumeignil F (2011) Green Chem 13:1960–1979
- 5. Davis SE, Ide MS, Davis RJ (2013) Green Chem 15:17-45
- Villa A, Dimitratos N, Chan-Thaw CE, Hammond C, Prati L, Hutchings GJ (2015) Acc Chem Res 48(5):1403–1412
- 7. Nakagawa Y, Tomishige K (2011) Catal Sci Technol 1:179–190
- Nakagawa Y, Tamura M, Tomishige K (2014) J Mater Chem A 2:6688–6702
- 9. Wang Y, Zhou J, Guo X (2015) RSC Adv 5:74611-74628
- Martin A, Armbruster U, Gandarias I, Arias PL (2013) Eur J Lipid Sci Technol 115:9–27
- Zhou C-H, Beltramini JN, Fan Y-X, Lu GQ (2008) Chem Soc Rev 37:527–549
- 12. Yue H, Zhao Y, Ma X, Gong J (2012) Chem Soc Rev 41:4218-4244
- Zhao X, Wang J, Yang M, Lei N, Li L, Hou B, Miao S, Pan X, Wang A, Zhang T (2017) ChemSusChem 10:819–824
- Bhanuchander P, Shanthi Priya S, Pavan Kumar V, Hussain S, Pethan Rajan N,, Bhargava SK, Chary KVR (2017) Catal Lett 147:845–855
- Fan Y, Cheng S, Wang H, Ye D, Xie S, Pei Y, Hu H, Hua W, Hua Li Z,a Qiao M, Zong B (2017) Green Chem 19:2174–2183
- Dasari MA, Kiatsimkul P-P, Sutterlin WR, Suppes GJ (2005) Appl Catal A 281:223–231
- 17. Barbelli ML, Santori GF, Nichio NN (2012) Bioresour Technol 111:500–503
- Villa A, Chan-Thaw CE, Campisi S, Bianchi CL, Wang D, Kotula PG, Kuebel C, Prati L (2015) Phys Chem Chem Phys 17:28171–28176
- Villa A, Veith GM, Prati L (2010) Angew Chem Int Ed 49:4499–4502
- Villa A, Campisi S, Mohammed KMH, Dimitratos N, Vindigni F, Manzoli M, Jones W, Bowker M, Hutchings GJ, Prati L (2015) Catal Sci Technol 5:1126–1132
- Shanthi Priya S, Bjanuchander P, Pavan Kumar V, Dumbre DK, Periasamy SR, Bhargava SK, Lakshmi Kantam M (2016) ACS Sustain Chem Eng 4:1212–1222
- Miyazawa T, Kusunoki Y, Kunimori K, Tomishige K (2006) J Catal 240:213–221
- 23. Montassier C, Menezo JC, Moukol J, Naja J (1991) J Mol Catal 70:65–84
- 24. Nakagawa Y, Shinmi Y, Koso S, Tomishige K (2010) J Catal 272:191–194
- 25. Khalil KMS (2007) J Colloid Interface Sci 315:562-569
- 26. Mori K, Hara T, Mizugaki T, Ebitani K, Kaneda K (2004) J Am Chem Soc 126:10657–10666
- Dimitratos N, Villa A, Prati L, Hammond C, Chan-Thaw CE, Cookson J, Bishop PT (2016) Appl Catal A 514:267–275
- Chan-Thaw CE, Chinchilla LE, Campisi S, Botton GA, Prati L, Dimitratos N, Villa A (2015) ChemSusChem 8:4189–4194
- Wahlström E, Lopez N, Schaub R, Thostrup P, Rønnau A, Africh C, Lægsgaard E, Nørskov JK, Besenbacher F (2004) Phys Rev Lett 303:026101-1–026101-4
- 30. ten Dam J, Hanefeld U (2011) ChemSusChem 4:1017-1034
- Morterra C, Meligrana G, Cerrato G, Solinas V, Rombi E, Sini MF (2003) Langmuir 19:5344–5356
- Toledo-Antonio JA, Cortés-Jácome MA, Navarrete J, Angeles-Chavez C, López-Salinas E, Rendon-Rivera A (2010) Catal Today 155:247–254
- 33. Morterra C, Cerrato G, Meligrana G (2001) Langmuir 17:7053-7060
- Guo LY, Zhuo JX, MaO JB, Guo XW, Zhang SG (2009) Appl Catal A 367:93–98