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Catalysis Today xxx (2008) xxx-xxx

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### Ti-MCM-41 catalysts prepared by post-synthesis methods Limonene epoxidation with $H_2O_2$

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#### Abstract

Ti-MCM-41 catalysts were prepared incorporating Ti by post-synthesis methods: wetness and wet impregnation of MCM-41 mesoporous solid. The samples were characterized by XRD at low angle, UV-vis DRS, Raman spectroscopy, BET, ICP and colorimetry. The results were compared with those obtained for Ti-MCM-41 hydrotermically prepared. All catalysts were tested in the limonene epoxidation with  $H_2O_2$ . Three kinds of Ti sites were observed in post-synthesis catalysts which were different from that found in the hydrothermal Ti-MCM-41. These structural differences do not produce important changes in the catalytic activity, instead a slight decrease in the epoxide selectivity can be observed. The catalysts prepared by post-synthesis methods are promising in the limonene epoxidation with  $H_2O_2$  since these methods are simple and reproducible.  $\bigcirc$  2007 Elsevier B.V. All rights reserved.

Keywords: Ti-MCM-41; Limonene epoxidation; Post-synthesis methods

### 1. Introduction

The chemical industry is often associated with the damage of the environment and, at least in a number of cases, this cannot be denied. In order to improve chemical processes using clean technologies, the study of selective catalytic oxidation of organic compounds with innocuous oxidants for the environment requirements are in progress. In the particular case of the terpenes epoxidation in fine chemical industry, the present methods use peracids as oxidants. However, equivalent amounts of acid waste are produced with these reactants. Therefore, there is a strong necessity for a development of new epoxidation methods using safer oxidants and producing little waste. In this way, hydrogen peroxide seems to be an attractive option as oxidant since it is cheap, easily available and only produces H<sub>2</sub>O as by-product. In relation with the nature of the catalysts, the heterogeneous ones are more adequate since they are easily separated from the product mixture and they can be recycled.

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Among terpenes, the limonene epoxidation using  $H_2O_2$  as oxidant has been carried out with different transition metal complexes supported on different solids. However, these catalysts have a poor stability and product yields, which leads to leaching during reaction and thus limits the recyclability [1,2].

Titanium has been widely used as catalyst for epoxidation reactions. In the last years, new solid materials that could be used as Ti support have been developed. Thus, the synthesis and characterization of different types of mesoporous solids, such as MCM-41 and SBA-15, have been reported by different authors [3–5]. These materials present a regular, hexagonal arrangement of uniform pores with diameters varying between 1.5 and 10 nm. In this way, some catalysts have been synthesized by grafting titanium on SBA-15, from TiCl<sub>4</sub> in gas phase. However, they also showed a very important Ti leaching with H<sub>2</sub>O<sub>2</sub> [6].

The titanium incorporation in the framework of MCM-41 by 60 hydrothermal methods was attained in order to use the peculiar 61 textural properties of these solids, turning them in attractive 62 catalysts to be used in epoxidation reactions of bulky 63 molecules, with  $H_2O_2$  or *tert*-butyl-hydroperoxide as oxidant 64 agents [7–9]. Notwithstanding, the low reproducibility of the 65

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Please cite this article in press as: D. Marino, et al., Ti-MCM-41 catalysts prepared by post-synthesis methods, Catal. Today (2008), doi:10.1016/j.cattod.2007.12.111

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preparation methods of these solids and the Ti leaching from 66 67 the framework of the MCM-41 appears as a problem to solve according to that reported by different authors [10,11]. In a 68 previous work [12], we have found that under certain reaction 69 conditions, no Ti leaching occurs during the limonene 70 epoxidation with H<sub>2</sub>O<sub>2</sub>. Nevertheless, to find catalysts that 71 allow to carry out the oxidation process with a minimum Ti 72 leaching using H<sub>2</sub>O<sub>2</sub> is still an interesting challenge. In this 73 way, other preparation methods have been implemented. They 74 75 consist in post-synthesis treatments, such as incipient wetness 76 and wet impregnation of MCM-41 mesoporous solid previously synthesized [13-16]. 77

Other authors [17,18] have shown that alumina itself is already active for the limonene epoxidation with  $H_2O_2$ . In spite of  $Al_2O_3$  is the simplest catalyst compared with those above mentioned, to improve the selectivity to epoxides it is necessary to use a continuous azeotropic distillation in order to remove the water from the reaction mixture. The necessity to use anhydrous  $H_2O_2$  is a disadvantage of this system.

Bearing in mind the previous discussion, the scope of the 85 present work is to study the effect that the preparation method 86 has on the location of titanium and its influence on the catalytic 87 activity in the limonene epoxidation with H<sub>2</sub>O<sub>2</sub>. With this 88 purpose MCM-41 catalysts modified with Ti were prepared 89 using incipient wetness and wet impregnation methods which 90 are the less studied in the bibliography. The solids were 91 92 characterized and tested in limonene epoxidation reaction with H<sub>2</sub>O<sub>2</sub>. The characterization and activity results were compared 93 with those obtained with a Ti-MCM-41 catalyst prepared by us 94 using the more traditional method, hydrothermal synthesis. All 95 catalysts were tested in such reaction conditions in which it was 96 97 demonstrated that the hydrothermal solid does not show Ti "leaching". 98

### 2. Experimental

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To obtain Ti-MCM-41 catalysts by post-synthetic techni ques, MCM-41 synthesized according to the methodology
 proposed by Ryoo and Kim [4] was used as support. The
 mesoporous material was subjected to two different ways for
 titanium incorporation:

(a) Wetness impregnation: a volume of Ti-tertbutoxide (Ti (*t*-BuO)<sub>4</sub>) (Sigma–Aldrich 99%) in isopropyl alcohol solution (IPA) (J.T.B. absolute, analytical quality) of the desired concentration equal to the MCM-41 pore volume was added, in order to obtain solids with Ti nominal loadings of 1, 2 and 3%, w/w.

112(b) Wet impregnation: the support was suspended in a Ti(t-113)113BuO)<sub>4</sub> in IPA solution. It was stirred during 5 h, and114centrifuged during 15 min. The solid was separated and115then washed with clean portions of IPA, until the turbidity116appearance (TiO<sub>2</sub> formation) was not observed by adding a119drop of water to the washing liquid.

121 Both impregnation methods were carried out in a glove box, 122 in  $N_2$  controlled atmosphere to avoid  $Ti(t-BuO)_4$  hydrolysis. In both cases the impregnated samples were dried in nitrogen atmosphere and then calcined in air stream (150 cm<sup>3</sup>/min) according to the following thermal program: 3 h at 393 K; the temperature was increased to 823 K at 10 K/min, remaining at this value during 5 h.

The solids prepared by wetness impregnation were called WNI-1, WNI-2 and WNI-3 for nominal Ti contents of 1, 2 and 3% (w/w)<sub>5</sub> respectively<sub>A</sub> and the solid prepared by wet impregnation was called WI.

Ti-MCM-41 catalyst prepared by hydrothermal synthesis, with a Ti content of 1.8% (w/w), was synthesized from tetraethyl-ortosilicate, cetyl-trimethyl-ammonium bromide and Ti-*tert*-butoxide, following the methodology described in ref [9]; this solid was called HT.

The samples were characterized by X-ray diffraction (XRD) at low angles, using Cu K $\alpha$  radiation, diffuse reflectance spectroscopy (DRS) in the UV–vis range, specific surface area and pore radii distribution by N<sub>2</sub> adsorption (BET), inductively coupled plasma (ICP) and UV–vis spectrometry to determine the content of Ti (expressed like TiO<sub>2</sub>). Raman spectra were adquired under room conditions using a Jasco TRS-600-SZ-P single monochromator spectrophotometer equipped with a coupled charge detector (CCD) cooled to about 150 K with liquid N<sub>2</sub>. The excitation source was the line of 514.5 nm of a Spectra 9000 Photometrics Ar ion laser. The laser power, measured at the samples, was set at 80 mW and the exposition time was of 180 s.

The activity and selectivity measurements have been carried out in a glass batch reactor at 343 K equipped with refrigerant, thermometer and magnetically stirred. For a standard reaction, 50 mg of catalyst were added to a reaction mixture of 4.32 mmol of limonene (L), 2.7 g of acetonitrile (CH<sub>3</sub>CN), used as solvent, and 1.17 mmol of H<sub>2</sub>O<sub>2</sub> 35%, w/w. The reactions were carried out in limonene excess since in a previous work we have found that in these conditions there is not titanium leaching [12]. Samples at different reaction times were obtained through a lateral tube of the reactor and were analyzed by GC using a capillary column (crosslinked methylsilicone gum, 30 m × 0.53 mm × 2.65 µm film thickness) and quantified with FID detector. Reaction products were identified by GC–MS. The H<sub>2</sub>O<sub>2</sub> conversion was measured by iodometric titration.

### 3. Results and discussion

#### 3.1. Solids characterization

XRD patterns of all samples (Fig. 1), are characteristics of materials with hexagonal arrangement [4,9]. The MCM-41 and the samples prepared by post-synthesis methods display an intense peak in  $2\theta = 2.1^{\circ}$ , two broad and less intense peaks in  $2\theta = 3.7^{\circ}$  and  $2\theta = 4.3^{\circ}$  and a very weak signal in  $2\theta = 5.7^{\circ}$  only in the samples obtained by wetness impregnation. The solid prepared by hydrothermal synthesis displays the main peak broadened, whereas two peaks less intense are not resolved. This result indicates that, in HT, a smaller structural regularity exists. In addition, the first peak appears shifted

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Fig. 1. XRD patterns of the catalysts.

towards high angles suggesting a different pore diameter or 177 wall thickness. 178

Table 1 shows Ti content (determined by colorimetry) and BET results for all solids. The values of specific surface area are characteristics of mesoporosous MCM-41. A decrease in the specific surface area with the increase of the Ti content is observed for the post-synthesis samples, compared with the support. However, important changes in the pore diameter and wall thickness values were not observed. This would indicate that the Ti species would partially occlude the channels. Sample HT displays channels of equal diameter but with small wall thickness. This result, jointly with the lower structural regularity determined by XRD, would indicate a smaller structural stability of this solid.

Fig. 2 shows DRS spectra of the solids. In HT a strong transition around 205 nm and a shoulder at 258 nm, can be observed. In a previous work [12] the band at 205 nm was assigned to charge transfer transition (CT) from an ion  $O^{2-}$  to a Ti<sup>4+</sup> ion located in the MCM-41 structure (closed sites) and the other band was assigned to open sites generated by the rupture of bridges Si-O-Ti of the closed sites, with the appearance of two OH groups, implying the insertion of a fifth oxygen atom to the first coordination sphere of titanium. The spectra of the samples obtained by wetness and wet impregnation methods are very different compared with HT. Considering that anatase displays a characteristic band centred at 330 nm [19] it would

Table 1

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Textural	properties	of	the	solids
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%TiO <sub>2</sub> (w/w)	$S_{\rm g}({\rm m}^2/{\rm g})$	$D_{\rm p}~({\rm nm})$	WT (nm)
0	1167	2.9	1.9
0.8	969	2.8	2.0
1.8	926	2.9	1.9
2.9	930	2.9	1.9
7.3	816	2.6	2.2
1.8	1104	2.8	1.6
	%TiO <sub>2</sub> (w/w) 0 0.8 1.8 2.9 7.3 1.8	%TiO <sub>2</sub> (w/w) $S_g(m^2/g)$ 0         1167           0.8         969           1.8         926           2.9         930           7.3         816           1.8         1104	%TiO <sub>2</sub> (w/w) $S_g(m^2/g)$ $D_p$ (nm)           0         1167         2.9           0.8         969         2.8           1.8         926         2.9           2.9         930         2.9           7.3         816         2.6           1.8         1104         2.8

 $S_{g}$ : specific surface area;  $D_{p}$ : pore diameter; WT: wall thickness.



Fig. 2. DRS spectra of the catalysts.

be useful to determine if this species is or not present in order to 203 carry out a more accurate assignement of the titanium species 204 present. With this purpose Raman spectroscopy was used since 205 it is well known that this technique is extremely sensitive to the 206 formation of crystalline  $TiO_2$  anatase particles [20]. The most 207 anatase intense peak  $(155 \text{ cm}^{-1})$  was not detected in any post-208 synthetic sample (Fig. 3), therefore, the crystalline  $TiO_2$ 209 (anatase) presence can be discarded. This result is in agreement 210 with that obtained by Gao et al. [20] for TiO<sub>2</sub> supported on 211 amorphous SiO<sub>2</sub>. They found trace amount of crystalline 212 anatase when the TiO<sub>2</sub> loading reaches 15%. The jointly use of 213 DRS and Raman techniques allow us to conclude that in post-214 synthetic samples, Ti<sup>4+</sup> ions are not located neither in 215 isomorphic substitutional positions of the MCM-41 framework 216 nor in anatase crystallites. In order to assign the possible Ti<sup>4+</sup> 217 ions locations, we fitted the DRS spectra with three Gaussian 218 curves whose maxima can be observed in Table 2. 219



Fig. 3. Raman spectra of the catalysts. The insert shows the pure anatase raman spectrum and the catalysts spectra with the same Raman intensity scale for illustrative purposes.

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Table 2

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Wavelength maxima and relative areas of the DRS UV-vis spectra fitting

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Sample	$\lambda_1(nm)$	(%)	$\lambda_2(nm)$	(%)	$\lambda_3(nm)$	(%)
WNI-1	240	25	269	29	297	46
WNI-2	241	21	263	21	300	58
WNI-3	241	23	265	19	304	58
WI	248	20	278	17	283	63

The fittings were carried out with Gaussian lines.

Gao et al. [20], showed that Ti species in isolated 220 221 tetrahedrons adsorbed on the pores surface of the SiO<sub>2</sub> support, show a band at 210 nm; dimeric or unidirectional chains of 222 tetrahedrons of  $TiO_4$  generate a band in 246 nm;  $TiO_5$ 223 bidimensional clusters display an absorption at 256 nm and 224 finally TiO<sub>6</sub> clusters absorb at 294 nm. According to this 225 description, the band corresponding to isolated TiO<sub>4</sub> was not 226 observed in our catalysts prepared by wetness or wet 227 impregnation, instead the other three bands could be detected. 228 Comparing the four samples, it can be seen that the Ti 229 coordination increases when the Ti loading increases. Besides, 230 it would be possible to conclude that in all samples the Ti ions 231 are mainly inside small clusters of  $(TiO_6)_n$  species, without 232 forming anatase crystals. It is known that the presence of this 233 compound in the catalyst is highly undesirable since it causes 234 the H<sub>2</sub>O<sub>2</sub> decomposition, diminishing the efficiency of this one 235 for the oxidation reaction. 236

### 3.2. Activity and selectivity measurements

Previous to the activity and selectivity measurements of the catalysts, a blank experiment was carried out mixing the reactants in absence of catalyst. A limonene conversion ( $\chi_L$ ) of 1% was obtained. This value was subtracted in all experiments in order to evaluate only the catalytic behavior. Table 3 shows the values of  $\chi_L$ , turnover frequency to epoxides (TOF), H<sub>2</sub>O<sub>2</sub> conversion ( $\chi_{H2O2}$ ), H<sub>2</sub>O<sub>2</sub> efficiency and reaction products distribution for all catalysts after 7 h of reaction. This time was selected since as it can be seen in Fig. 4, between 5 and 7 h of reaction time, all catalysts have reached a value of limonene conversion approximately constant. In each case it was corroborated by mass spectrometry, that the only obtained



Fig. 4. Limonene conversion (as a percentage of the maximun possible conversion) vs. reaction time for all catalysts. Reaction conditions: T = 343 K, 4.32 mmol of limonene (L), 2.7 g of acetonitrile, 1.17 mmoles of H<sub>2</sub>O<sub>2</sub> 35%<sub>7</sub> w/w and 50 mg of catalyst.

products are those listed in Table 3. On the other hand, the mass balance was verified making a balance of grams of carbon between the consumed limonene and the obtained products.

Considering that the limonene is in excess respect to  $H_2O_2$ ,  $\chi_L$  was calculated as a percentage of the maximum possible conversion, assuming complete conversion of  $H_2O_2$  to oxygenated products. TOF to epoxides, is defined as the number of epoxides moles obtained by mol of Ti per hour (h<sup>-1</sup>). It was assumed that all Ti atoms are exposed on the catalysts surface and all of them have the same activity. The  $H_2O_2$  efficiency represents the percentage of this reagent that leads to oxygenated products of limonene. Finally, considering that the only products detected by GC–MS are those shown in Table 3 (no by-products were detected), the selectivity to an "*i*" product (*S<sub>i</sub>*), is expressed as:

 $\frac{S_i = \text{``i'' product (mol)}}{\sum_{i=1}^n \text{``i'' product (mol)} \times 100.}$ 

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Table 3

Activity and selectivity	results of limonene	epoxidation with H <sub>2</sub> O <sub>2</sub>	after 7 h of reaction
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Catalyst	χ <sub>L</sub> (% max.)	TOF (h <sup>-1</sup> )	Х н2О2	Efficiency H <sub>2</sub> O <sub>2</sub> <sup>a</sup>	Selectivity (mol.%) <sup>b</sup>			
					Epoxides <sup>c</sup>	Carvone + Carveol	Diepoxide	Glycols
WNI-1	48	6.9	68	71	43	32	14	11
WNI-2	50	3.4	74	68	46	34	9	11
WNI-3	51	2.3	79	64	48	33	7	12
WI	48	0.9	81	59	50	31	5	14
HT	52	4.5	80	65	58	22	12	8

Reaction conditions: T = 343 K, 4.32 mmol of limonene (L), 2.7 g of acetonitrile, 1.17 mmol of H<sub>2</sub>O<sub>2</sub> 35%; w/w and 50 mg of catalyst.  $\chi_L$  (% max.): limonene conversion as a percentage of the maximum possible conversion. TOF (h<sup>-1</sup>): turnover frecuency to epoxides.  $\chi_{H2O2}$ : conversion of the oxidant determined by iodometric titration.

 $^{a}$  H<sub>2</sub>O<sub>2</sub> efficiency, percentage of H<sub>2</sub>O<sub>2</sub> consumed to produce oxygenated products.

<sup>b</sup> Expressed as a percentage of the total formed products.

<sup>c</sup> 1,2 and 8,9 epoxylimonene.

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The catalysts prepared by post-synthesis treatments show 267 very similar values of limonene conversion. However, the H<sub>2</sub>O<sub>2</sub> 268 conversion increases as the content of Ti is increased up to 81% 269 for WI catalyst, whereas the H2O2 efficiency follows an inverse 270 relation. Finally, the TOF to epoxides shows an important 271 decrease, about one order of magnitude from WNI-1 to WI. 272 These results can be explained if it is assumed that Ti located in 273  $(TiO_6)_n$  sites is inactive or less active for the selective oxidation 274 of limonene and besides these sites show tendency to 275 decompose the  $H_2O_2$ , since the relative population of these 276 277 sites is the only one that increases from WNI-1 to WI (Table 2). The solid prepared by hydrothermal synthesis, shows values of 278  $\chi_{\rm L}$ , TOF to epoxides,  $\chi_{\rm H2O2}$  and  $\rm H_2O_2$  efficiency similar to 279 WNI-2 catalyst. Both catalysts have the same Ti content 280 (1.8% w/w). However, HT solid has Ti sites different from the 281 ones of the post-synthetic samples. Therefore, the presence of 282 Ti<sup>4+</sup> sites replacing Si<sup>4+</sup> ions in the MCM-41 framework, would 283 not be necessary to produce the catalytic results mentioned 284 285 above.

When Ti loading increases for post-synthetic samples, the 286 selectivity results show that: the epoxides increase, the 287 diepoxide diminishes and glycols and carvone + carveol 288 remain approximately constant as it can be seen in Table 3. 289 It can be considered that the diepoxide synthesis can follow 290 two ways: a consecutive reaction where, first a double bond is 291 epoxidized and then the other one, or a simultaneous 292 epoxidation of both double bonds. The two mechanisms 293 could occur on Ti<sup>4+</sup> tetrahedral sites of dimeric or 294 unidirectional chains and/or on pentacoordinated Ti (TiO<sub>5</sub>). 295 Besides, the relative population of tetrahedral sites remains 296 nearly constant, whereas the pentacoordinated sites diminish 297 from WNI-1 to WI. Taking into account the above 298 considerations, if diepoxide production would occur on 299 TiO<sub>5</sub> sites following Scheme 1, the epoxide percentage might 300 diminish. However, epoxide percentage increases, therefore, 301 the diepoxidation through Scheme 2 would occur on TiO<sub>5</sub> 302 sites. Consequently, when TiO<sub>5</sub> population decreases, diep-303 oxide production also diminishes. 304



Scheme 1. Proposed consecutive diepoxidation mechanism.

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Scheme 2. Proposed simultaneous diepoxidation mechanism.

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The greater selectivity to products of allylic oxidation (carveol and carvone) showed by these catalysts, can be 306 explained by a radicalary mechanism [12], competitive with the 307 formation of an active intermediate, peroxo-metal complex, 308 showed in Schemes 1 and 2. The free-radicals (OH) production 309 310 would be favoured by Ti species with greater coordination number,  $TiO_5$  and  $(TiO_6)_n$  which are absent in HT. 311

On the other hand, HT catalyst shows the highest selectivity 312 to epoxides and the lowest production of allylic oxidation 313 (carvone + carveol) among all solids tested in this work. This 314 result demonstrates that the isolated tetrahedral sites of Ti<sup>4+</sup> in 315 framework replacing positions of the MCM-41 are the most 316 317 selective ones for the limonene epoxidation reaction when  $H_2O_2$  is used as oxidant. 318

Finally, as it was mentioned above, Ti-MCM-41 can 319 experiment Ti "leaching" when H2O2 is used as oxidant 320 leading to a little activity in homogenous catalysis [21]. Bearing 321 in mind that, the Ti<sup>4+</sup> ions in HT are hardly strained in 322 substitutional positions inside the SiO<sub>2</sub> lattice, the breaking of 323 324 Ti bond and its extraction by  $H_2O_2$  may be easy. However, in a previous work [12] we found the operative conditions to avoid 325 Ti leaching in HT. In the present work we have used the same 326 reaction conditions for post-synthesis catalysts and Ti leaching 327 was not observed, considering the detection limits of the 328 technique used (ICP). The assays were carried out in the liquid 329 phase after 7 h of reaction for all catalysts. Taking into account 330 that substitutional Ti<sup>4+</sup> was not detected in post-synthesis 331 catalysts, the Ti leaching absence is an expected result, since in 332 these solids the Ti<sup>4+</sup> ions are less strained. 333

### 4. Conclusions

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In the present work different Ti-MCM-41 catalysts were 335 prepared via post-synthesis (wetness and wet impregnation) 336 methods. The solids showed a better long order hexagonal 337 arrangement in comparison with a solid obtained by hydro-338

thermal method. Using DRS and Raman spectroscopy, it was demonstrated that these solids present three types of Ti sites, which are different from that found in HT catalyst. These structural differences do not lead to significant changes in  $\chi_{L}$ values, in limonene epoxidation with H<sub>2</sub>O<sub>2</sub>. However, the selectivity to epoxides appears diminished with an increase in the production of allylic oxygenated (carvone + carveol). These results indicate that the structural characteristics of the Ti<sup>4+</sup> sites do not influence on the catalyst activity but they are important in order to define the solid selectivity.

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Besides, the greater structural quality of post-synthesis mesoporous solids compared with HT, does not produce beneficial effects on the activity and/or selectivity in the limonene epoxidation with  $H_2O_2$ .

On the other hand, in the post-synthesis solids, Ti loadings higher than about 2% do not produce a  $\chi_L$  increase and H<sub>2</sub>O<sub>2</sub> efficiency is noticeable lower. Therefore, it is not recommended to surpass this loading. Finally, we think that the epoxide selectivity differences between HT and post-synthesis solids are compensated by the simplicity and reproducibility of the last preparation methodology.

### Acknowledgements

The authors acknowledge support of this work by Consejo Nacional de Investigaciones Científicas y Técnicas, Comisión de Investigaciones Científicas de la Provincia de Buenos Aires, Universidad Nacional de La Plata, Universidad Tecnológica Nacional, Facultad Regional Córdoba and Agencia Nacional de Promoción Científica y Tecnológica (PICT No. 14-08456).

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Please cite this article in press as: D. Marino, et al., Ti-MCM-41 catalysts prepared by post-synthesis methods, Catal. Today (2008), doi:10.1016/j.cattod.2007.12.111

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