INGENIERÍA OUÍMICA

Preparación y caracterización del óxido de estaño y su uso como catalizador en la reacción de epoxidación de esteres grasos insaturados

CHEMICAL ENGINEERING

Tin oxide preparation and characterization and use as a catalyst in the reaction of unsaturated fatty esters epoxidation

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Resumen

La síntesis y la caracterización del óxido de estaño se estudiaron en este trabajo para evaluar su actividad catalítica en la reacción de epoxidación del metilricinoleato proveniente del aceite de ricino. La obtención del óxido se realiza por el método de precipitación controlada, encontrándose que se obtienen ventajas significativas al emplear soluciones acuosas de SnCl₂.2H₂O como precursor. Su caracterización se efectuó por medio de las técnicas de DTA, TGA, DRX, encontrándose que para obtener un alto rendimiento hacia el sistema casiterita el valor de pH óptimo fue de 6.25, mientras que un pH de 8.0 conlleva hacia el sistema romarchita. En el caso de las técnicas de MEB y área superficial, se determinó que el tamaño de partícula encontrado estaba por debajo de los 50 nm con un área superficial de 19 m²/g. El ester ricinoleico se obtuvo a partir del aceite empleando metilato de sodio como catalizador y seguidamente se hace reaccionar con peróxido de hidrógeno como agente oxidante y óxido de estaño para producir un epóxido. Finalmente, los productos de reacción se identifican por cromatografía de gases, encontrando que los óxidos de estaño son activos para la obtención de epóxidos con un mejor desempeño del sistema cristalino romarchita (SnO) principalmente hacia el metil-12-hidroxi-9,10-epoxioctadecanoato (MHEOD).

Palabras clave: Casiterita, epoxioctadecanoato, epoxidación, óxido de estaño, romarchita.

Abstract

Tin oxide synthesis and characterization were studied in the present work in order to evaluate its catalytic activity in the reaction of methyl ricinoleate from castor oil. Oxide is obtained through the controlled precipitation method, and significant advantages are obtained when using water solutions of SnCl₂.2H₂O as precursor. Its characterization was done using the techniques DTA, TGA, XRD, finding that in order to have high performance for the cassiterite system the optimum pH value was 6,25 whereas a pH of 8.0 was leading to the romarchite system. When using SEM techniques and a surface area, the size of the particle found was below 50 nm with a surface area of 19 m²/g. The ricinoleate ester was obtained from the oil using sodium methylate as catalyst and then making it react with hydrogen peroxide as oxidant agent and tin oxide to produce an epoxide. Finally, reaction products are identified through gas chromatography finding that tin oxides are active for obtaining epoxides with better performance of the romarchite crystal system (SnO) especially towards the methyl-12-hydroxi-9, 10-epoxioctadecanote (MHEOD).

Key words: Cassiterite, epoxi-octadecanoate, epoxidation, tin oxide, romarchite.

1. Introduction

Epoxidation of oleochemical products has been the object of academic and industrial research (L.A. Rios *et al*, 2005, R. E. Harry-O'kuru *et al*, 2005), and derived products of vegetal oil have a commercial importance due to their application in plasticizers and polymer industries (U. Biermann *et al*, 2000). Vegetal oil can be directly epoxidized but not much work has been done in this regard. One work done is the one reported by V.V. Goud *et al* (2006a), obtaining in this case mono, di- and tri-epoxides. It is clear that the results depend on the reactive used and in this sense A. Corma *et al* (2007) have reported the use of formic acid in the presence of a heavy metal as complexing agent.

Epoxidation of fatty acids or methyl esters (FAME) coming from vegetal oils has been more studied, and in this case the double bond is attacked (V. V. Goud *et al*, 2009; J. Sepulveda *et al*, 2007; A. Campanella *et al*, 2008; E. Poli *et al*, 2009; and K. Wadumesthrige *et al*, 2009). This process requires an acid catalyst and generally mineral acids have been used because of their low cost, although there are works giving details about the catalyst attack on the oxiran ring during the reaction (L.H. Gan *et al*, 1992, F.A. Zaher *et al*, 1989, and A. Campanella and M.A. Baltanas, 2006).

It is worth mentioning the use of the aluminium oxide as catalyst in FAME's epoxidation, focusing on two alumina, one produced by the sol-gel process, and recognizing that both systems present very good catalytic activity in this reaction, though the alumina prepared by the sol-gel process is more efficient (J. Sepulveda, *et al* 2007). These authors state that after 24 hours the attainted conversion is of the order of 95%, for a selectivity of 97%, though after its reutilization is less active.

On the other hand, an oxidizing species such as the molecular oxygen has also been tested in the epoxidation of heavy olefins by J-R. Monnier (2001). What is shown in the study is that epoxidation of 1,3-butadiene is carried out with an Ag catalyst with the formation of an oxidizing

species in situ (hydrogen peroxide from molecular O2). It is worth noticing that epoxidation of non-allylic species or sterically or kinetically-hindered ones require promoters such as Cs or Rb or some thallium salts to help in desorption of the epoxidized olefin. Other systems used successfully in the epoxidation reaction are the Cu/V by L. Yang *et al* (2010), Ag by J-R Monnier (2001) a polimetacrilate peroxotugstene or heteropolioxometaltes by G. Gelbard *et al* (2000) and Z. Xi *et al* (2001), or even complex Mn salt (III) and Mo (IV) by T. Luts *et al* (2007). The use of a tungsten-based solid has been reported, the phosphotungstate, but this time evaluating the absence of solvent over the selectivity in epoxidation (E. Poli *et al*, 2009).

Researchers conclude that the catalyst produces a selective transformation of cis-methyl oleate in cisepoxioleate under much more cleaner conditions, due to the use of a stoichiometric amount of of hydrogen peroxide and the suppression of solvent in the medium of reaction. Such a result is probably due to a synergic effect between the hydrogen peroxide and the air oxygen at a low temperature 313K, though they also state the possibility of a low dimerization of the methyloleate due to catalyst saturation because of the air oxygen.

As far as the tin oxide is concerned, this is a system with good affinity with the carbonyl and carboxyl functions, it is confirmed that it enhances its capacity when a second element is incorporated (P.J. Smith, 1998). Among the incorporated elements for such purpose there is phosphorus, antimony, vanadium, chromium, molybdenum, manganese, and copper. These systems have been used to promote the oxidation of carbon monoxide, sulfur dioxide and saturated and unsaturated hydrocarbons, as well as other organic compounds. Palladium can also be used in methane oxidation (K. Skizawa et al, 2000) and for the oil transesterification processes and for dimethyloxalate, where tin oxide is mixed with other solids, such as SiO2, ZnO, and TiO2 (J. Urresta et al, 2000, S.Wang et al, 2004 and S. Verdier et al, 2003). Finally, in the present work a tin oxide type romarchite and casserite was synthesized, through the method of controlled precipitation. The solid obtained was used in the methyl esters epoxidation reaction resulting from the castor oil plant with hydrogen peroxide as oxidizing agent.

2. Experimental

2.1 Materials and Reagents

Reactives used in the present work such as KOH, SnCl₂*2H₂O, and Al₂O₃ were obtained from Mallinckrot, the acetic acid from Aldrich, the castoroil acid in a composition of 90% in ricinoleic acid form *Genquímicos Ltda*, and chloroform, ethanol, sodium anhydrous sulfate, sodium methoxide, tert-butanol, hydrogen peroxide, ammonium hydroxide, and 1-fenilethanol from Merck. Tin oxide was obtained by the controlled precipitation method (CPM), using SnCl₂ as starting material and via the above mentioned process by Avila and Rodriguez-Paez (2009).

2. 2 Characterization of solids

To obtain the crystalline phases a Phillips PW1830/40 diffractometer with Co anode (k = 1,790A) and an iron filter, operating at 40kV and 30 mA, at a pace of 0.02° and a fixed time of 0.3 second for a range of 20 to 80° was used. Microstructural characterization of solids was determined via a SEM electronic microscope (JEOL 1200 EX) operating at 80 keV. Powders were previously dried at a furnace at 80° for 24 hours and finally macerated in an agate mortar. Solids were also characterized with the differential thermic and gravimetric analysis (Shimadzu TG / Dt - 50) in order to determine the endo- and exothermic processes that could be performed during thermic treatment. Analysis was carried out in the presence of dry air (7L/min flux) at a heating speed of 10°C/min with Al₂O₃ as reference material.

2.3 Preparation of methyl ester from castor-oil

To remove free fatty acids in the oil a treatment is performed with a concentrated ethanol solution of KOH at a temperature of 65°C, during 2 hours.

The mixture obtained is warmed in backflow during 4 hours and while it is hot it is diverted to a separation funnel. After an HCl diluted solution of 0.1 N is added, for the separation of the aqueous and organic phases, and the respective phase is extracted. To this phase 5mL of distilled water were added and an emulsion was then formed and treated with a similar quantity of chloroform/ 50/50 and a cooling-down period was ethanol allowed for the separation of the phases. To the organic phase selected solvent is evaporated, and then a small quantity of sodium anhydrous sulfate is added to absorb the remaining water. For the preparation of methyl esters of the castor-oil plant, to the previous product was added methanol in a relationship of 1/6 with sodium methoxide in methanol as catalyst. The mixture was put in backflow during 4 hours at 65°C and a washing process was performed as it was described before.

2.4 Epoxidation Reaction

In a three-neck flask with 50 ml capacity and equipped with a condenser, approximately 3 g (~10 mmol) of methyl ester were placed. Then, 1.2 ml of tert-butanol was added as solvent and immediately after 2 ml of a homogenized solution at 6% of hydrogen peroxide (H₂O₂) were added in 1-fenilethanol, the latter being added to stabilize the hydrogen peroxide and lastly 0.1 g of catalyst is added. The solution was heated to reflux at 10°C with constant agitation during 7 hours. The following conditions were also tested: reaction time (2, 4, 6, 7, 8, and 10 hours), reaction temperature (110 and 130°C), and heat treatment in muffle to SnO catalyst at 2% by mass (100,200,300, 450, and 600°C) and the quantity of oxidizing solution (1, 2, and 3 ml of 6% solution oh hydrogen peroxide in 1-fenilethanol). After completing the reaction time, catalyst was stirred for centrifugation.

2.5 Chromatographic Analysis

All compounds have been analyzed using a CG Hewlett Packard 6890 with a FID detector and equipped with a HP-% column (30mx0.32mmx0.25µm). Injector and detector

temperatures were 260° and 280°C respectively. Carrier gas was helium and the furnace temperature was 50°C for 3 min and then it increased to 10°C/min C, and keeping this temperature for 10 min. Dodecane was used as internal standard.

3. Results and discussion

3.1 Synthesis and characterization of tin oxide, through the determination of solid crystalline phases

Figure 1 shows pH change when doing titration with ammonium hydroxide of SnCl₂*2H₂O dissolutions using acetic acid and ethanol as solvent. Through addition of SnCl₂ in acetic acid, it is expected that the mains species in solution would be anionic

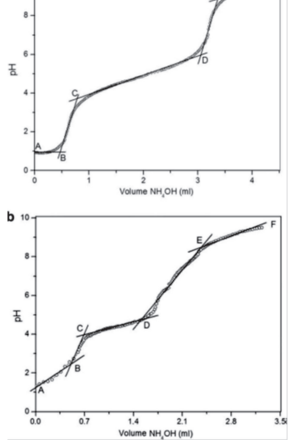


Figure 1. Potentiometric titration curves of 0.1 M solution SnCl₂·2H₂O in: (a) Acetic acid and (b) Ethanol as solvent.

compounds, because of the complexing nature of the ethyl anion; the most important species in the solution would be the tin tricarboxylate ones (II) [Sn(RCO₂)₃].

When analyzing the behavior of the SnCl₂-CH₃COOH system through the addition of NH₄OH in figure 1, we can observe five regions: two with significant pH changes in (BC and DE) and three plateaus (AB, CD, EF). Region AB corresponds to the H⁺ ion neutralization produced by the reaction (1), forming less stable compounds and the tricarboxylates-stannates already mentioned.

The third CD region represents high consumption of OH ions due to dissolvent neutralization (Reaction (1)) and the hydrolysis of the compounds formed in previous phases.

$$CH_3COOH + NH_4OH \rightarrow CH_3COONH_4 + H_2O$$

Reaction (1)

In this region there are also polynuclear species $[Sn_2(OH)_4O]_2^-$, where the oxygen bridge makes it possible for the pyramidal structure to get stable Sn(II). Polynuclear species such as $[Sn_2(OH)_2]_2^+$, $[Sn_3(OH)_4]_2^+$ and $[SnO(OH)_2]_2^-$ can be formed as products of the reactions of hydrolization reaction $SnCl_2$ (E. Ararat, 2007). The union of these polynuclear species would allow the embryos' growth which, upon reaching the critical size, would develop the nuclei of the solid phase.

The development of the nuclei in the DE region is the most important part of the process since the first ever precipitate appears at pH 6.2, right at the start of this region. In the last EF region, pH stabilization corresponds to the system saturation. Here, the tinoxy type complexes must prevail, benefiting the SnO formation, as indicated on the DRX and FTIR analysis. Moreover, when ethanol is employed as a solvent (Fig 1(b)), five similar regions are observed, as shown in figure 1 (a). Through this system, the following reactions may occur:

$$SnCl2+2H2O \xrightarrow{C_2H_5OH} Sn(OH)_2+2H^++2Cl^-$$
Reaction (2)

In the AB region an almost linear increase in pH is shown due to the neutralization of H+ in the system, representing the dissolving of tin chloride (II). The second BC area represents the formation of soluble complexes originated in hydrolysis reactions (Reaction 2). In the third CD region, the vast consumption of OH- is due to the condensation reaction that lead to the polynuclear species such as $[Sn_2(OH)_2]^{2+}$, $[Sn_3(OH)_4]^{2+}$ y $[SnO(OH)_2]^{2-}$. The last EF region shows the saturation dissolving due to the excess of ammonium hydroxide in the system of ethanol chloride-Sn (II).

Figure 2(a) shows the X-ray diffractogram of the solid obtained from the SnCl₂–CH₃CO₂H at pH 8.0, where a crystalline romarchite phase is mainly found according to the JCPDS 06-0395 files. A small formation of cassiterite SnO₂ (JCPDS 21-1250) is observed in the unwashed system. These results allow concluding that it is possible to obtain SnO by employing acetic acid as a solvent with a small formation of SnO₂. When the cleansings

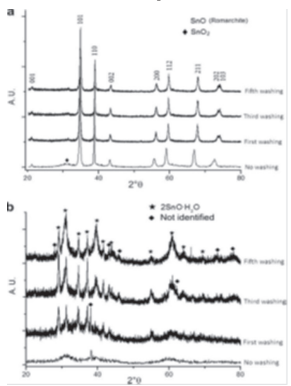


Figure 2. XRD patterns of calcined solids at 600 °C and obtained by means of: (a) SnCl₂–CH₃CO₂H, and (b) SnCl₂–CH₃CH₂OH, both with and without washing with diethylamine solution 0.05 M.

are done with an aqueous dissolution of 0.05 M in diethylamine, the presence of the crystalline romarchite phase is exclusively observed. Figure 2(b) shows the diffractograms of prepared solids in ethanol at pH 6.5, being observed that in the unwashed sample an amorphous solid with the presence of an incipient formation of cassiterite (SnO₂) is found. Furthermore, when the elimination of the chlorines with the diethylamine solution is carried out, the crystallization of a romarchite hydrate is favored SnO*2H₂O (JCPDS 07-0194). The presence of other peaks is also observed, but they are not related to any reference compound in particular. Of the previous results, it can be concluded that when using acetic acid as a solvent and taking the pH to 8.0, a crystalline romarchite phase is obtained, whilst upon employing ethanol as a solvent and afterward taking the pH to 6.5 the crystalline phase obtained in this simple after thermal treatment to 400 °C was cassiterite.

3.2 Morphology and Size of the Particles

Figure 3 shows particles of an irregular shape

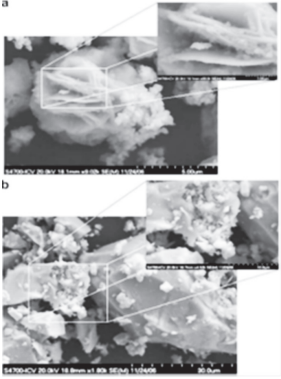
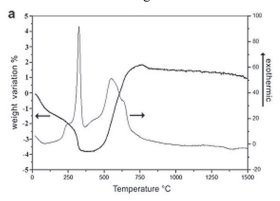


Figure 3. SEM images obtained from the solid SnO treated at 600 °C for 2 h: (a) SnCl₂-CH₃CO₂H, y (b) SnCl₂-CH₃CH₂OH.

with the presence of agglomerates of 200 μm . The primary particles have a smaller size of 2 μm . Specifically in Figure 3(a) a micrometric cluster flower type SnO is observed, obtained by the Sn (II) salt in acetic acid. The formation of polycrystalline leaves made of spherical nanoparticles. In Fig. 3(b) a cluster with a dense structure and of a micrometric size is shown, obtained by the SnCl₂-etanol system.

3.3 Thermic Analysis of Solids

Figure 4(a) shows the DTA/TGA curve of the precipitate solids at pH 8.0 with acetic acid as a solvent. In the TGA curve, the loss of a small weight below the 100°C is due to the loss of adsorbed water. The weight loss between 200-250°C, as is with the exothermic peak in the DTA curve, might be associated with water and ammonia, chemically and structurally absorbed. At 300°C there is weight loss associated with the exothermic peak in the DTA curve which indicates organic matter combustion



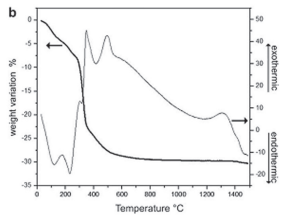


Figure 4. DTA curves and corresponding to the solids obtained from the solution of $SnCl_2*2H_2O$ in TGA: (a) acetic acid at pH = 8.0 and (b) ethanol at pH = 6.5.

and the oxidation of the hydroxyl groups (F. Lui *et al*, 2005). The TGA curve at higher temperatures between 400 and 700°C may be associated with the SnO oxidation or with the SnO disproportion expressed in the following equations:

$$2SnO + O_2 \rightarrow 2SnO_2$$

Reaction (3)

Nevertheless, it must be known that tin may be present during the thermic treatment, due to stoichiometric reactions of disproportionation as shown by H. Giefersa *et al.* (2005).

$$2SnO \rightarrow SnO_2 + Sn$$
Reaction (4)
$$3SnO \rightarrow Sn_2O_3 + Sn$$
Reaction (5)
$$2Sn_2O_3 \rightarrow 3SnO_2 + Sn$$
Reaction (6)

The DTA and TGA curves in Fig. 4(b) correspond to the solid obtained from SnCl₂–C₂H₅OH a pH 6.5 system. 9% of the weight loss between 110 and 250°C is associated with two endothermic

Table 1. Specific values for castor oil and its composition.

Acid value (mg/1g sample)	0.808
Saponification value (mg/1g sample)	312.66
Iodine value (g/100g sample)	13.06
Peroxide value (meq/1000g)	2.74
Physical state	Yellow clear viscous liquid
Solubility	Ethanol 98%, chloroform 99%
Melting point	5.5 °C

Methyl ester	Percentage	
C16:0	0.5	
C18:0	1.3	
C18:1	2.3	
C18:2	6.8	
C18:1-OH	89.0	

Table 2. Epoxidation of methyl esters of castor oil

Catalyst	Conversion (%)	Selectivity to epoxide				
		MEOD	MEODECEN	MDEOD	MHEOD	
Al2O3	9.5		3.0	9.9	87.1	
SnO	39.0	2.4	4.0	8.0	85.7	
SnO2	5.1		3.5	6.5	90.5	

Reaction temperature $T_r = 110$ °C, reaction time $t_r = 7$ h, tert-butanol as solvent, 2 ml of a homogenized solution 6% hydrogen peroxide (H_2O_2) in 1-phenylethanol, catalyst mass $m_{cata} = 0.1$ g of catalyst. Methyl-12-hydroxy-9,10-epoxyoctadecanoate (MHEOD), Methyl-9,10- epoxyoctadecanoate (MEOD), Methyl -9,10- epoxyoctadecanoate (MEODECEN), Methyl -9,10,12,13-diepoxyoctadecanoate (MDEOD).

peaks of the DTA curve. This corresponds to chemically connected water. The second weight loss (11%) at 250-400° would be associated with two narrow exothermic peaks which can correspond to organic matter decomposition (particularly ethoxy groups). Above the 400°C a constant weight loss is obtained and the DTA peaks may be associated to crystallization material.

3.4 Determining Composition and Characteristics of Castor Oil and Methyl Ester

Table 1 shows the values of the different determined rates for the commercial castor oil.

The physical state shows us that the liquid oil possesses a light yellow coloring and is soluble in ethanol and chloroform, due to the OH group present in the ricinoleic acid. The value of the acidity rate (0.808) indicates that the oil has a low quantity of free fatty acid, so it can be assumed that the oil, in a great percentage, is found in a triglyceride state and this is demonstrated with

the value of the saponification or esterification rate (312.66). The seemingly low iodine rate (13.06) for the composition of the castor oil might be due to sterical impediment. The peroxide rate determines the quality of the oil in terms of its state of rancidity and for this case it is 2.74. This low value shows that the oil is in good state and, therefore, the double bonds are not altered.

The ester stemming from the castor oil presents 5 products (see table 2), considering the majority product the ester of the ricinoleic acid.

3.5 Study of the Reaction of the Epoxidation of the Methyl Ester

Firstly the epoxidation reaction of the methyl esters is carried on with (table 3), employing 3g (\sim 10 mmol) of methyl ester, 1.2 ml of tert-butanol as a solvent, 2 ml of a 6% hydrogen peroxide ($\rm H_2O_2$) homogeneous mixture in 1-phenylethanol and 0.1 g of catalyst, with a reaction temperature of 110°C in constant mixing for 7 hours. For this reaction, the solids obtained

Table 3. Effect of SnO calcination temperature on the reaction epoxidation.

Calcination temperature °C	Conversion (%)	Selectivity to epoxide		
		MEOD	MEODECEN + MDEOD	MHEOD
100	5.6			100
200	21.8		10.0	90.0
300	22.9			100
450	39.0	2.4	11.9	85.7
600	76.8		9.5	90.5

Reaction temperature $T_r=110$ °C, reaction time $t_r=7$ h, tert-butanol as solvent, 2 ml of a homogenized solution 6% hydrogen peroxide (H_2O_2) in 1-phenylethanol, catalyst mass $m_{cata}=0.1$ g of catalyst.

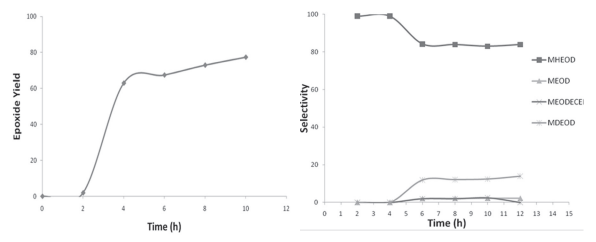


Figure 5. Epoxidation reaction with SnO calcined at 600 ° C. (a) yield (b) Selectivity. Reaction conditions: Temperature $T_{Rx} = 130$ ° C, tert-butanol as solvent, 2 ml of hydrogen peroxide (H_2O_2) 6% 1-phenylethanol, mass of catalyst $m_{cata} = 0.1$ g.

through the method of controlling precipitation, (Romarchite (SnO), Cassiterite (SnO $_2$)), and the Al_2O_3 as a reference catalyst.

Comparing the different catalysts, it can be observed that the Romarchite (SnO) presents the biggest conversion. Regarding Aluminium oxide (Al₂O₃), it has a low conversion percentage, but it also generates products of epoxidation. According to the obtained results, it were obtained methyl-12-hydroxy-9, 10-epoxyoctadecanoate (MHEOD), methyl-9,10-epoxyoctadecanoate (MEOD), methyl-9, 10-epoxyoctadecanoate (MEODECEN) y methyl-9, 10, 12, 13-diepoxyoctadecanoate (MDEOD). The most efficient solid for the epoxidation reaction is

the romarchite and it is thus chosen as a catalyst to continue with the study.

3.6 The effect of the SnO calcination temperature on the epoxidation reaction

In table 4 the results of the epoxidation results for the SnO at different calcination temperatures are shown. In the figure 5 it can observe that the efficiency increases in the measure that the temperature of the thermic treatment does so as well with a slight exception at 300°C, being the thermic treatment at 600°C the one that presents the highest conversion percentage in the epoxidation reaction. The selectivity remains practically

Table 4. Effect of the amount of oxidizing solution on SnO activity against the epoxidation reaction of the
methyl ester from castor oil.

Oxidizing solution in mL	The reaction temperature in °C	Conversion (%)		Selectivity t	o epoxide	
			MEOD	MEODECEN	MDEOD	MHEOD
1	110	25.6	0	0	9.2	90.8
	130	52.9	0	0	14.3	85.7
2	110	76.8	0	0	9.5	90.5
	130	66.0	1.7	2.0	10.9	85.4
3	110	75.6	1.5	2.9	8.1	87.5
	130	96.2	1.5	2.4	10.5	84.7*

^{*}Other products possibly forming epoxide decomposition products found. Reaction conditions: Temperature $(T_{Rx}) = 110$ and $130 \,^{\circ}$ C , tert -butanol as the solvent , 1, 2 , 3 mL of hydrogen peroxide (H_2O_2) 6% 1-phenylethanol , catalyst mass $(m_{coor}) = 0.1$ g .

constant for the 5 tested thermic treatments, being the majority product the methyl-12-hydroxy-9, 10-epoxyoctadecanoate (MHEOD).

When evaluating the behavior of the solid processed at 600°C regarding time and to a reaction temperature of 130°C, an increase is observed in the efficiency percentage in the measure that the reaction time increases with an almost constant selectivity. It can also be observed that the reaction at 130°C tends to degrade the products after the 12 hours of reaction, given that a coloring in the reaction products is seen.

3.7 The effect of oxidizing mixture on the efficiency of the epoxidation reaction

Under the established conditions different quantities of oxidizing mixture were tested for two of the worked on temperatures, 110 and 130°C. The table 4 shows the principal results obtained...

The results for 110°C indicate that the enzyme substrate/oxidizing agent relationship is directly proportional to the conversion factor, and when working with a temperature of 130°C, a more important conversion is observed, but similar selectivity than found at 110°C. This last behavior seems logical because in the measure that the enzyme substrate/oxidizing agent relationship increases there will be greater conversion percentage, though the efficiency of the catalyst, which may be affected by the concentration of the oxidizing agent mixture, is not to be dismissed. Thus, the reaction needs a quantity of 3mL of the oxidizing agent.

4. Conclusions

Based on the obtained results, it can be concluded that for the synthesis of the tin oxide the type of solvent has an influence on the synthesis process and it affects the morphology and the size of the particle. Indeed, when employing acetic acid, the presence of polynuclear species [Sn₂(OH)₄O]²⁻ is favored, in which the oxygen bridge would allow the stability of the pyramidal structure Sn(II) and

polynuclear species like [Sn₂(OH)²]²⁺, [Sn₃(OH)₄]²⁺ and [SnO(OH)₂]²⁻ could be formed too. The union of these polynuclear species would allow the growth of the embryos which, upon reaching critical size, would develop tin-oxy type complexes, benefitting the formation of SnO.

Regarding the epoxidation reaction, the efficiency of the romarchite was determined, with the solid treated thermically at 600°C being the most active one. The reaction temperature of 130°C and of an oxidizing mixture of hydrogen peroxide is necessary to epoxide the ricinoleic ester effectively towards the methyl-12-hydroxy-9, 10-epoxy octadecanoate (MHEOD) (85% selectivity), with a 96% conversion. Shown was also the need to add 3 mL of oxidizing mixture of hydrogen peroxide (H₂O₂) at 6% in 1-phenylethanol to obtain better results.

5. Acknowledgements

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6. References

Ararat C., Mosquera A., Parra R., Castro M.S., & Rodríguez-Páez J.E., (2007). Synthesis of SnO₂ nanoparticles through the controlled precipitation route. *Materials Chemistry and Physics* 101, 433-440

Ávila H.A., & Rodríguez-Páez J.E., (2009). Solvent effects in the synthesis process of tin oxide, *Journal of Non-Crystalline Solids* 355, 885–890.

Biermann U., Friedt W., Lang S., Lühs W., Machmüller G., Metzger J.O., Rüsch gen Klaas M., Schäfer H.J., & Schneider M.P., (2000) . New Syntheses with Oils and Fats as Renewable Raw Materials for the Chemical Industry. *Angewandte Chemie International Edition* 39, 2206–2224

Campanella A., Fontanini C., & Baltanas M-A., (2008). High yield epoxidation of fatty acid

methyl esters with performic acid generated in situ. *Chemical Engineering Journal* 144, 466–475.

Campanella A., & Baltanas M.A., (2006). Degradation of the oxirane ring of epoxidized vegetable oils in liquid–liquid heterogeneous reaction systems. *Chemical Engineering Journal* 118, 141–152.

Corma A., Iborra S., & Velty A., (2007). Chemical Routes for the Transformation of Biomass into Chemicals. *Chemical Reviews* 107, 2411-2502

Gan L.H., Goh S.H., & Ooi K.S., (1992). Kinetic studies of epoxidation and oxirane cleavage of palm olein methyl esters. Journal of the American Oil Chemists' Society 69, 347–351.

Gelbard G., Breton F., Quenard M., & Sherrington D-C., (2000). Epoxidation of cyclohexene with polymethacrylate-based peroxotungstic catalysts. *Journal of Molecular Catalysis A: Chemical* 153, 7–18.

Giefersa H., Porschb F., & Wortmann G., (2005). Kinetics of the disproportionation of SnO. *Solid State Ionics* 176, 199–207.

Goud V. V., Patwardhan A.V., & Pradhan N.C., (2006a). Studies on the epoxidation of mahua oil (Madhumica indica) by hydrogen peroxide. *Bioresource Technology* 97, 1365–1371.

Goud V.V., Pradhan N.C., Patwardhan A.N., (2006b). Epoxidation of karanja (Pongamia glabra) oil by H2O2. *Journal of the American Oil Chemists' Society* 83, 635-640

Harry-O'kuru R.E., Gordon S.H., Biswas A., (2005). A facile synthesis of aminohydroxy triglycerides from new crop oils. *Journal of the American Oil Chemists' Society* 82, 207-212.

Lui F., Quan B., Lui Z., & Chen L., (2005). Surface characterization study on SnO2 powder modified by thiourea. *Materials Chemistry and Physics* 93, 301-304.

Luts T., Frank R., Suprun W., Fritzsche S., Hey-Hawkins E., & Papp H., (2007). Epoxidation of olefins catalyzed by novel Mn(III) and Mo(IV)-Salen complexes immobilized on mesoporous silica gel Part II: Study of the catalytic epoxidation of olefins. *Journal of Molecular Catalysis A: Chemical* 273, 250–258.

Monnier J-R., (2001). The direct epoxidation of higher olefins using molecular oxygen. *Applied Catalysis A: General* 221, 73–91.

Poli E., Clacens J-M., Barrault J., & Pouilloux Y., (2009), "Solvent-free selective epoxidation of fatty esters over a tungsten-based catalyst", *Catalysis Today* 140, 19–22.

Rios L.A., Weckes P., Schuster H., & Hoelderich W.F., (2005). Mesoporous and amorphous Tisilicas on the epoxidation of vegetable oils. *Journal of Catalysis* 232, 19-26.

Sekizawa K., Widjaja H., Maeda S., Ozawa Y., & Eguchi K., (2000). Low Temperature of Methane over Pd/SnO2 Catalyst. *Applied Catalysis A: General* 200,211-217.

Sepulveda J., Teixeira S., & Schuchardt U., (2007). Alumina-catalyzed epoxidation of unsaturated fatty esters with hydrogen peroxide. *Applied Catalysis A: General* 318, 213–217.

Smith P.J., (1998). Chemistry of Tin, Ed. Blackie Academic & Professional – Chapman & Hall, Great Britain, 2nd ed. pp. 35 – 36, 443 – 475.

Urresta J., Ramírez A., Martínez C., & Vargas L., (2000). Transesterificación del aceite de Palma empleando Catalizadores Soportados a base de Sn y Ni. Palmas, 21 No. Especial, 381-385

Verdier S., Didillon B., Morin S., & Jumas J.C., (2003). Pd-Sn/Al₂O₃ Catalyst from Colloidal Oxide Synthesis. *Journal of Catalysis* 218, 280-287.

Wadumesthrige K., Salley S.O., & Simon Ng K.Y., (2009). Effects of partial hydrogenation,

epoxidation, and hydroxylation on the fuel, properties of fatty acid methyl esters. Fuel Processing Technology 90, 1292–1299.

Wang S., Ma X., & Yang X., (2004). Transesterification of Dimethyl Oxalate with Phenol under SnO2 / SiO2 Catalysts. *Industrial & Engineering Chemistry Research* 43, 4027-4030.

Xi Z., Wang H., Sun Y., Zhou N., Cao G., & Li M., (2001). Direct epoxidation of olefins catalyzed by heteropolyoxometalates with molecular oxygen

and recyclable reductant. *Journal of Molecular Catalysis A: Chemical* 168, 299–301.

Yang L., He J., Zhang Q., & Wang Y., (2010). Copper-catalyzed propylene epoxidation by oxygen: Significant promoting effect of vanadium on unsupported copper catalyst. *Journal of Catalysis* 276, 76-84.

Zaher F.A., El-Mallah M.H., & El-Hefnawy M.M., (1989). Kinetics of oxirane cleavage in epoxidized soybean oil. *Journal of the American Oil Chemists' Society* 66, 698–700.