Rare Earth metal triflates $M(O_3SCF_3)_3$ (M = Sc, Yb, La) as Lewis acid catalysts of depolymerization of poly-(bisphenol A carbonate) via hydrolytic cleavage of carbonate moiety: catalytic activity of La(O_3SCF_3)_3

Eugenio Quaranta*

Dipartimento di Chimica, Università degli Studi "Aldo Moro" di Bari, Campus Universitario, Via E. Orabona, 4, 70126, Bari, Italy

DOI: 10/1016/j.apcatb.2017.01.007 ABSTRACT

Lewis acids such as $M(O_3SCF_3)_3$ (M = Sc, Yb, La) triflate salts have been used for the first time as catalysts of the hydrolysis reaction of poly-(bisphenol A carbonate) (PC), which is an attractive route to the chemical recycling and valorization of waste PC. In THF, the $M(O_3SCF_3)_3$ (M = Sc, Yb, La) salts effectively catalyzed the hydrolysis of PC to bisphenol A (2,2-bis(4-hydroxyphenyl)propane; BPA) and CO₂. La(O₃SCF₃)₃ promoted the depolymerization of the polymer with the highest BPA yield and selectively, as, unlike the other triflates investigated, it displayed also very poor tendency to promote, under the working conditions, the decomposition of the monomer (BPA) to phenol, 4-isopropenylphenol (4-IPP) and 4-isopropylphenol (4-IPPH). The influence of temperature, reaction time, catalyst load and H₂O/PC weight ratio on the catalytic activity of La(O₃SCF₃)₃ has been investigated. The La(O₃SCF₃)₃ catalyst can be recovered quantitatively at the end of the reaction and reused. The catalytic behavior of the M(O₃SCF₃)₃ salts has been compared with that of a strong proton donor such as CF₃SO₃H. The results obtained allow to exclude that protons arising from [L_nM(OH₂)]³⁺ (L = ligand (H₂O, THF, etc)) aquo-complexes might play a catalytically important role in promoting the depolymerization process.

Keywords: poly-(bisphenol A carbonate); plastics chemical recycling; hydrolysis; Lewis acid catalysis; lanthanum triflate

* Corresponding Author. Tel.: +39 080 5442093. *E-mail address:* <u>eugenio.quaranta@uniba.it</u> (E. Quaranta).

1. Introduction

In recent years, recycling of plastics has gained growing attention as a suitable strategy to reduce the environmental impact of this typology of wastes. This approach is noteworthy as it regards waste plastics as a potential resource rather than a mere refuse and responds to the current widespread demand of saving both carbon and energy and protecting fossil resources from depletion [1-4]. Mechanical, chemical and quaternary recycling (or energy recovery) are the main methods for recycling polymer wastes from the post-consumer stage. Chemical recycling, in which the polymer chains are converted to smaller molecules through chemical processes, is particularly intriguing, although, to date, industrial implementation of this technology has generally found economic restraints related to the plant and processing costs. A significant boost in overcoming these problems may come from the search for new technologies, simpler from the operational standpoint, based on selective chemical processes which avoid harsch reaction conditions and make use of selective catalytic systems susceptible to be reused.

Because of the excellent physical and chemical properties (transparency, thermal stability, etc) poly-(bisphenol A carbonate) (PC) is one of the most interesting thermoplastics and is widely used in several fields (optics; electronic components; data storage; automotive, aircraft, and security components; medical applications). PC consumption reached 3.4 Mton in 2010 and its market is continuously expanding year by year [5-7]. The rise in the utilization of PC calls for the developments of after-use treatments. Valorization of waste PC by chemical recycling can provide an attractive alternative to other options, such as landfill disposal, mechanical recycling, energy production (energy recovery) [1-3,6]. This approach aims at regenerating the starting monomer 2,2-bis(4-hydroxyphenyl)propane (bisphenol A; BPA), which can be reused to produce new virgin plastic, or also at obtaining chemicals with added value.

In principle, different strategies can be pursued for recycling waste PC by chemical methods. Pyrolysis has been proposed as a promising approach, but, usually, it suffers from low monomer selectivity because of formation of large amount of by-products [8]. Another viable option implies the fission of carbonate bond through a chemical reaction [4,9-35] such as, for instance, hydrolysis [21-34], aminolysis [19,20] or alcoholysis and glycolysis [9-18]. These methods are generally directed to regenerate BPA, but can also cogenerate high added value coproducts, for instance organic carbonates or ureas if the polymer is reacted with alcohols or amines, respectively. In this work the attention has been turned to the hydrolytic route (Eq. (1)). Reaction (1) has been investigated under a variety of conditions¹, often very severe because of poor solubility of PC in water. In order to promote the process under less severe conditions the use of a few reaction media able to dissolve the polymeric material has been also explored [24,33,34].

Insert equation 1

Nevertheless, the hydrolysis reaction (Eq. (1)) usually requires the assistance of a suitable catalyst. Most of studies have focused on the utilization of base [24-29,31,33,34] rather than acid catalysts.² Usually, these protocols make use of large loads of base, which cannot be reused, and imply other disadvantages such as equipment corrosion and environmental problems [32]. In the attempt to overcome the above shortcomings ionic liquids have been also employed both as reaction media and catalysts [32].

Compared with base catalysts, acid catalysts have been much less investigated [23,28,29]. Proton donors, such as HCl, H₂SO₄, HNO₃, CH₃CO₂H, were tested as catalysts in sub-critical water, but under the working conditions they showed no or poor activity [28]. More recently, the steam hydrolysis of PC has been investigated at $T \ge 572$ K in a fluidized bed reactor in the presence of quartz sand. SiO₂, by acting as a proton donor through its surface OH groups, mainly caused the fission of the BPA propylidene bridge, with generation of volatile fragments that were further hydrolyzed in the steam phase [29]. Recently, PC hydrolysis has been also studied in hot compressed water ($T \ge 523$ K) using Mn(O₂CCH₃)₂ as the catalyst [23]. A Broensted acid catalysis (Scheme 1) was operating under the working conditions and CH₃CO₂H, generated by hydrolysis of the Mn-salt, played the role of the catalytically active species. The selectivity to BPA was moderate in the above processes [23,29] due to the side-formation of phenol, 4-isopropenylphenol, and other species.

The behavior of Lewis acids (LAs) as catalysts of reaction (1) still remained unexplored. In principle, also Lewis acids, likewise the proton (Scheme 1), may promote the hydrolysis reaction by activating the carbonyl moieties along the polymer chain to nucleophilic attack by H₂O, provided that these systems are stable enough in the presence of water. Rare Earth triflates M(O₃SCF₃)₃ [37,38] are well known in the literature for their oxophilicity and stability in the presence of water and have been used as water-tolerant Lewis acid catalysts in a variety of reactions. Differently from conventional Lewis acids such as BF₃, AlCl₃, TiCl₄, etc., M(O₃SCF₃)₃ salts can be easily recovered and reused without any loss of catalytic activity. The above properties as well as the low toxicity

¹ For instance, using: high pressure and/or high temperature steam [22,25,26,29]; either supercritical or sub-critical water [21,23,28,30]; non conventional heating techniques such as microwave (MW) irradiation [31-33].

² The hydrolysis of strictly related compounds, such as diaryl- and aryl alkyl carbonates (or also $ArCO_2R$ (R = alkyl or aryl) esters), in the presence of a base (NaOH) proceeds faster than in the presence of a strong proton donor such as $HCIO_4$ [36].

and environmentally benign character, make Rare Earth triflates attractive reagents for many organic trasformations. Recently this group has shown that both $Sc(O_3SCF_3)_3$ and $La(O_3SCF_3)_3$ salts are effective Lewis acid catalysts for methoxycarbonylation of aliphatic and aromatic amines with carbonic acid diesters, such as dimethyl carbonate (DMC) or methyl phenyl carbonate, to afford carbamate methyl esters [39-43]. In these processes the metal center can act as a Lewis acid catalyst by coordinating the organic carbonate through the carbonyl oxygen atom and activating the carbonyl carbon to the nucleophilic attack by amine. The ability of the metal center to interact with the carbonyl group of the organic carbonate has been fully demonstrated by isolating, for the first time, a DMC–metal complex characterized as $[(\eta^1-O(C=O)-DMC)Sc(O_3SCF_3)_3]$ [44]. Herein, $Sc(O_3SCF_3)_3$ and $La(O_3SCF_3)_3$ have been studied as potential catalysts of the hydrolysis reaction of poly-(bisphenol A carbonate). The study has considered also $Yb(O_3SCF_3)_3$, which is known to show excellent catalytic activity in many LA-catalyzed organic reactions. The catalytic behavior of $La(O_3SCF_3)_3$ has been focused, as this salt can promote reaction (1) selectively with higher BPA yield than the other metal triflates $M(O_3SCF_3)_3$ (M = Sc, Yb) investigated.

Insert Scheme 1

2. Experimental

2.1. General methods and materials

M(O₃SCF₃)₃ (M = Sc, Yb, La) salts, as well as La(NO₃)₃6H₂O, LaCl₃ and La₂O₃, were from Aldrich. Commercial THF used as reaction medium was pretreated according to conventional methods (Na/benzophenone) [45] and stored under N₂. In this study pure PC pellets (3 mm length × 2 mm diameter) were used as a model of waste PC. Polycarbonate used in this work was from Aldrich (M_w ~ 64000). Scheme 2 shows the repeating unit (254.29 g/mol) of the polymer and highlights the nature of the additive, 4-(2-phenylpropan-2-yl)phenol (4-cumylphenol), used as a chain-terminator [46]. The spectroscopic characterization (¹H NMR, ¹³C NMR, ATR-FTIR) of the used polymeric material is reported in the Supplementary Material (Fig. S1-S3). GC analyses were performed with a HP 5890 Series II gas-chromatograph (capillary column: Heliflex AT-5, 30 m × 0.25 mm, 0.25 µm film thickness). GC-MS analyses were carried out with a Shimadzu GC-17A linked to a Shimadzu GC-MS QP5050 selective mass detector (capillary column: Supelco MDN-5S, 30 m × 0.25 mm, 0.25 µm film thickness). IR spectra were taken on a Shimadzu FTIR Prestige 21 spectrophotometer or a Perkin Elmer Frontier MIR/FIR spectrophotometer equipped with a Pike GladiATR (diamond crystal) accessory. NMR spectra were recorded with a Varian Inova 400 spectrometer or with an Agilent 500 instrument. Chemical shifts are in δ (ppm) vs TMS (¹H, ¹³C) or CFCl₃ (¹⁹F).

Insert Scheme 2

2.2. Hydrolysis of poly-(bisphenol A carbonate): general procedure

Into a glass tube, containing the M(O₃SCF₃)₃ catalyst, if used, and the polymer (0.5 g), the solvent (THF, 6 mL) was introduced. After solubilization of the polymer a measured volume of distilled H₂O was added.³ The glass reactor containing the reaction mixture was transferred into a steel autoclave previously purged under a N₂ stream. The reaction mixture was reacted at the working temperature for the established time under autogenous pressure (as for the details of the experimental conditions used in the catalytic runs, see Table 1, 3, 4 and 6, as well as Figure 2 and 3) and then cooled to room temperature. The liquid phase was analyzed by FTIR, GC/MS and/or GC, using biphenyl as the internal standard.⁴ BPA yield was calculated through equation (2), wherein n_{BPA} are the moles of BPA determined *via* GC in the reaction mixture or isolated (see Section 2.2.1) and n°_{BPA} are the moles of BPA units incorporated in the mass *w* of PC feed and calculated through BPA yield (%) = ($n_{\text{BPA}}/n^{\circ}_{\text{BPA}}$)×100 (2) equation (3),⁵ where MM_{UR} (254.28 g/mol) is the molar mass of the repeating unit,

⁵ Indicating with $n^{\circ}_{4-\text{cumylphenol}}$ the moles 4-cumylphenol incorporated in the mass *w* of PC feed, equation (3) can be derived by solving the following system of equations:

$n^{\circ}_{4-\text{cumylphenol}}/n^{\circ}_{\text{BPA}} = 0.035$	(determined by NMR, see Supplementary Material)	(3')
$w = X_{UR} \times MM_{UR} + Y \times MM_{bis(4-current)}$	$nylphenyl)$ carbonate = $n^{\circ}_{BPA} \times MM_{UR} + \frac{1}{2}n^{\circ}_{4-cumylphenol} \times MM_{bis(4-cumylphenyl)}$ carbonate	(3'')

where equation (3") is the mass balance stating that the mass w of PC feed consists formally of X moles of repeating units UR (X = n°_{BPA}) and Y mol of bis(4-cumylphenyl) carbonate (Y = $\frac{1}{2} n^{\circ}_{4-cumylphenol}$), see also Scheme 2. As $n^{\circ}_{4-cumylphenol} = 0.035 \times n^{\circ}_{BPA}$, then

³ The used triflate salts $M(O_3SCF_3)_3$ (M = Sc, La, Yb) were found to be soluble in the THF/H₂O solvent mixture. Although the polymer and H₂O are, separately, easily soluble in THF, the solubility of the polymer in a THF/H₂O solution is affected by the water content of the solution because of the poor solubility of PC in H₂O and decreases as the mixture becomes too much rich in water. For instance, on adding H₂O to a THF (3 mL) solution of PC (0.250 g) at room temperature (293 K) the system remained initially homogeneous, but it turned opalescent after addition of 0.24 mL of H₂O because of precipitation of the polymer. Continuing the addition of H₂O to the system caused the precipitation of greater and greater amounts of PC.

Raising temperature favours the solubilization of the polymer by increasing the solubility of PC in the solvent mixture. For instance, adding H₂O (0.5 mL) to a THF (6 mL) solution of PC (0.514 g) ($m_{H2O}/m_{PC} = 0.97$) caused, at ambient temperature, the partial precipitation of the dissolved polymer, which, however, easily re-dissolved upon heating the suspension to 373 K. Cooling the system again to room temperature caused the re-precipitation of part of PC. In another test, to a solution of PC (0.513 g) in THF (6 mL) a greater volume of H₂O (1 mL; $m_{H2O}/m_{PC} = \sim 2$) was added. In this case the amount of PC which precipitated at room temperature was markedly more copious and the system remained heterogeneous even when the resulting suspension was heated to temperatures ≤ 393 K. On the basis of the above tests it can be inferred that, at the working temperatures (see Results), reaction (1) can proceed homogeneously for values of m_{H2O}/m_{PC} weight ratio lower than or around ~ 1 . However, it is very likely that for significantly higher values of m_{H2O}/m_{PC} weight ratio the reacting system may remain heterogeneous at the working temperature because of the uncomplete solubilization of the polymer.

⁴ An analogous procedure was followed when $La(NO_3)_3$ $^{\circ}6H_2O$, $LaCl_3$ and La_2O_3 were used as catalysts (see Table 5 for the experimental conditions).

MM_{bis(4-cumylphenyl) carbonate} (450.58 g/mol) is the molar mass of bis(4-cumylphenyl) carbonate (see also Scheme 2) and 0.035 is the molar ratio 4-cumylphenol/BPA in the used polymer (Supplementary Material).

$n^{\circ}_{\text{BPA}} = w/(\text{MM}_{\text{UR}} + \frac{1}{2} \times 0.035 \times \text{MM}_{\text{bis}(4-\text{cumylphenyl})\text{carbonate}})$

The recovery of unreacted polymeric material can be accomplished by evaporating in vacuum the THF solution, treating the residue with methanol and filtering the resulting suspension. The solid on the filter was washed with a small volume of MeOH, dried in vacuum and weighted. By indicating with *w*' the mass of the solid on the filter, the PC conversion can be estimated through equation (6): PC conversion (%) = $[(w - w')/w] \times 100$ (6)

In a blank experiment carried out in the absence of any catalyst, to the solution of PC (502.5 mg) in THF (6 mL) 0.500 mL of H₂O were added (H₂O/PC: 0.995 (m_{H2O}/m_{PC}), 14.2 (mol_{H2O}/mol_{CO3})). The glass reactor containing the reaction mixture was transferred into a steel autoclave previously purged under a N₂ stream and stirred at 433 K for 3 h. The analysis of the reaction mixture, once cooled to room temperature, did not show any significant conversion of the polymer to BPA (traces).

2.2.1. Work-up of the reaction mixture: isolation of BPA

After complete conversion of the polymer, the reaction mixture, cooled to room temperature, was evaporated under vacuum at ambient temperature and the solvent was recovered by condensation in a cold bath. The residue was extracted with diethyl ether/H₂O and the two phases were separated. Pure BPA can be isolated (85-90% yield; Fig. S4-S7) from the ethereal phase (previously dried on MgSO₄) by chromatography on a silica gel column, using, as eluent, petroleum ether/ethyl acetate 6:1 (v/v) until elution of the minor amounts of 4-cumylphenol (Fig. S8 and S9) and, afterwords, petroleum ether/ethyl acetate 2:1 (v/v).

2.2.2. Recovery of the catalyst $(La(O_3SCF_3)_3)$

The reaction mixture, after removal of solvent, was extracted with diethyl ether/H₂O (see also 2.2.1). The white solid isolated from the aqueous phase upon evaporating the solvent (H₂O) under vacuum was characterized by FTIR, ¹³C NMR and ¹⁹F NMR (Fig. S10-S12) and identified as pure

from which equation (3) can be obtained.

 $mol_{CO3} = X + Y = n^{\circ}_{BPA} + 0.5 \times n^{\circ}_{4-cumylphenol} = n^{\circ}_{BPA} + 0.5 \times 0.035 \times n^{\circ}_{BPA}$

(3)

 $w = n^{\circ}_{\text{BPA}} \times \text{MM}_{\text{UR}} + \frac{1}{2} (0.035 \times n^{\circ}_{\text{BPA}}) \times \text{MM}_{\text{bis}(4\text{-cumylphenyl}) \text{ carbonate}}$

Knowing the value of n_{BPA}° (Eq. (3)), the moles of hydrolysable CO₃ groups (mol_{CO3}) contained in the mass *w* of PC feed can be calculated as follows (Eq.(5)):

 $La(O_3SCF_3)_3$ (yield: 90-95%) by comparison with the corresponding spectra of a commercial sample of $La(O_3SCF_3)_3$.

3. Results

3.1. Hydrolysis of poly-(bisphenol A carbonate) in the presence of $M(O_3SCF_3)_3$ (M = Sc, La, Yb) salts

The hydrolysis reaction (Eq. (1)) was carried out in THF, which can dissolve both the polymer and H₂O.

Table 1 shows that, at 433 K, $Sc(O_3SCF_3)_3$, $Yb(O_3SCF_3)_3$ and $La(O_3SCF_3)_3$ can promote reaction (1).⁶ After 6 h, the conversion of PC, monitored by FTIR, was found to be practically

Insert Table 1

quantitative in all the cases under study. Fig.1 (spectrum a) displays the FTIR spectrum of the reaction solution at the end of the catalytic run, when La(O₃SCF₃)₃ was used as the catalyst. The complete depolymerization of PC was confirmed by the absence of any significant absorption at 1778 cm⁻¹, where the stretching $v_{C=O}$ of PC can be located in THF solution (Fig. 1, spectrum b). The IR spectrum of the reaction solution also showed the typical bands of BPA at 1614 and 1593 cm⁻¹ (C=C skeleton vibrations) [47,48], partially masked by the band at 1647 cm⁻¹ due to the excess of H₂O present in the reaction medium, as well as the absorption at 2336 cm⁻¹ assigned to dissolved CO₂, the other product of the hydrolysis reaction (Eq. (1)).

Insert Figure 1

The generation of the monomer (BPA) was accompanied by that of 4-cumylphenol, which is not a side-product of reaction (1) (see also Supplementary Material), but the alkylphenol used as a chain terminator and molecular weight regulator in the synthesis of the used PC [46]. Phenol, 4-isopropenylphenol (4-IPP) and 4-isopropylphenol (4-IPPH) were found as side-products, as ascertained by GCMS analysis. These species formed in very low amounts when La(O_3SCF_3)₃ was used as the catalyst, but their formation became more significant switching to Yb(O_3SCF_3)₃ and, even more, to Sc(O_3SCF_3)₃.

The influence of reaction time on BPA productivity was probed for all the used triflate catalysts. Both scandium and ytterbium triflate were more active than the corresponding lanthanum salt. In

⁶ In the absence of any catalyst (see 2.2), PC conversion to BPA was found to occur in trace amounts. The depolymerization reaction can proceed catalytically also at a temperature as low as 373 K, but more slowly, with longer conversion times poorly appealing from an applicative point of view. As an example, at 373 K, in the presence of $Yb(O_3SCF_3)_3$ as the catalyst (THF, 6 mL; PC, 502 mg; H₂O, 0.5 mL; catalyst, 125 mg), PC conversion and BPA yield after 9.5 h were found to be equal to 60% and 53%, respectively.

fact, under working conditions analogous to those described in Table 1, polycarbonate conversion, monitored by FTIR, was almost complete within 2 h when using $Sc(O_3SCF_3)_3$ or $Yb(O_3SCF_3)_3$ as the catalysts, but it was quantitative after a longer time (6 h) in the presence of the lanthanum salt. However, the BPA yield measured at the above times (Sc: 28% after 2 h; Yb: 83% after 2 h; La: 97% after 6 h; see Fig. 2), indicated that selectivity decreases in the order La>Yb>Sc due to the growing incidence of the side-formation of PhOH, 4-IPP and 4-IPPH.⁷ In the presence of the Sc or Yb salt, BPA yield reached a maximum and then decreased with time once PC conversion was complete (~2 h), suggesting that BPA formed through the hydrolysis reaction may be involved in a subsequent decomposition process, probably promoted by the used M(O_3SCF_3)_3 catalyst and affording the observed side-products. This issue has been probed further.

Insert Figure 2

3.1.1. BPA cleavage promoted by $M(O_3SCF_3)_3$ (M = Sc, La, Yb) salts

4,4'-Dihydroxydiarylmethane derivatives have been recognized for a long time to undergo thermal and hydrolytic degradation (T > 523 K) [49]. More specifically, decomposition of BPA generates phenol, 4-isopropenylphenol, 4-isopropylphenol (Scheme 3) and resinous substances [49]. Phenol Insert Scheme 3

and 4-isopropenylphenol are obtained by cleavage of BPA propylidene bridge, while 4isopropylphenol forms by subsequent reaction of 4- isopropenylphenol initially produced [49,50]. Acid catalysts (H₂SO₄, 4-Me-C₆H₄-SO₃H, KHSO₄, ZnCl₂) can promote BPA cleavage at milder temperatures, also under anhydrous conditions [49]. Therefore, in order to account for the kinetic courses observed in Fig. 2, BPA cleavage was investigated in the presence of $M(O_3SCF_3)_3$ (M = Sc or Yb or La). Table 2 shows that the $M(O_3SCF_3)_3$ (M = Sc, Yb, La) salts can promote the decomposition of BPA to PhOH, 4-IPP and 4-IPPH catalytically even under anhydrous conditions, wherein the decomposition of the monomer is faster than in the presence of water (in Table 2, compare entries 3, 5, 7 with entries 2, 4, 6, respectively).⁸ Noteworthily, under conditions comparable with those used in reaction 1 (Table 2, entries 2, 4, 6), BPA decomposition proceeded slowly in the presence of the lanthanum salt (entry 6), while it was faster in the presence of

⁷ The reaction with $La(O_3SCF_3)_3$ was very selective. In this case (Figure 2 curve c), PhOH, 4-IPP and 4-IPPH formed in very low yield. The relative molar ratio 4-IPPH:4-IPP was found to be approximately equal to 0.1.

The formation of PhOH, 4-IPP and 4-IPPH was more important with $Yb(O_3SCF_3)_3$ and $Sc(O_3SCF_3)_3$. In the presence of $Yb(O_3SCF_3)_3$ (Figure 2, curve b), 4-IPPH always formed in lower amounts than 4-IPP (4-IPPH/4-IPP $\approx 0.2 \div 0.3$ mol/mol). However, in the presence of $Sc(O_3SCF_3)_3$ (Figure 2, curve a), the formation of 4-IPPH was much more important than with $Yb(O_3SCF_3)_3$ and the molar ratio 4-IPPH/4-IPP increased markedly with the reaction time to become close to 1.

⁸ In all the cases, the molar ratio 4-IPPH/4-IPP increased moving from $La(O_3SCF_3)_3$ to $Yb(O_3SCF_3)_3$ to $Sc(O_3SCF_3)_3$. However, whatever $M(O_3SCF_3)_3$ (M = Sc, Yb, La) catalyst was used, under anhydrous conditions a higher 4-IPPH/4-IPP molar ratio was found than in the presence of water. Remarkably, under anhydrous conditions and in the presence of $Sc(O_3SCF_3)_3$, BPA degraded to PhOH and 4-IPPH, while 4-IPP was detected in trace amounts.

 $Yb(O_3SCF_3)_3$ (entry 4) and, even more, with $Sc(O_3SCF_3)_3$ (entry 2). These results provide a rationale for the higher BPA productivity and selectivity displayed by $La(O_3SCF_3)_3$ and allow to conclude that $La(O_3SCF_3)_3$, albeit less active than $Sc(O_3SCF_3)_3$ and $Yb(O_3SCF_3)_3$, is by far the most selective towards BPA formation as, under the working conditions, it exhibits a very poor tendency to promote the cleavage of the monomer.

Insert Table 2

3.2. Hydrolysis of poly-(bisphenol A carbonate): catalytic activity of La(O₃SCF₃)₃

The above results emphasize the difficulty of controlling the selectivity of the hydrolysis reaction (Eq. (1)) relative to BPA cleavage when $Sc(O_3SCF_3)_3$ or $Yb(O_3SCF_3)_3$ were the used catalysts. In virtue of the higher BPA selectivity displayed by $La(O_3SCF_3)_3$ the catalytic behavior of the lanthanum salt was explored more thoroughly.

Fig. 3 shows the influence of H₂O/PC weight ratio on the productivity of the process (Eq. (1)). Various runs (Fig. 3) were carried out wherein a solution of PC (~0.5 g) in THF (6 mL) was catalytically hydrolyzed in the presence of higher and higher amounts of H₂O, using, therefore, a growing H₂O/PC weight ratio. Under the working conditions, BPA yield at first rose with increasing the availability of H₂O in the reaction medium (for kinetic reasons, being H₂O one of the reactants). However, on augmenting further the amount of feed H₂O (and therefore increasing the H₂O/PC ratio) the BPA yield decreased after reaching a maximum for a H₂O/PC ratio equal to 0.988 (m_{H2O}/m_{PC}). BPA yield was found to be practically null for a m_{H2O}/m_{PC} ratio as high as 5.65 (Fig. 3). The diminution of BPA productivity is not due to the increase of BPA degradation, but, at least in part (see also Section 4), may reflect the fact that, as previously noted,³ a too high concentration of H₂O in the reaction mixture diminishes the solubility of PC in the reaction medium and, therefore, for values of m_{H2O}/m_{PC} weight ratio significantly higher than 1 (for instance, m_{H2O}/m_{PC} ≥ 2 in Figure 3) the reacting system may remain heterogeneous at the working temperature because of the incomplete solubilization of the polymer.

Insert Figure 3

Therefore, the study was continued using a H_2O/PC weight ratio close to 0.988 (m_{H2O}/m_{PC}), corresponding to a molar ratio mol_{H2O}/mol_{CO3} close to 14.⁵

Table 3 illustrates the effect of catalyst loading. At 433 K, reaction (1) proceeded effectively with loads lower than 10 mol%. However, in the latter cases quantitative conversion of the polycarbonate required a longer reaction time. For instance, using a catalyst load as low as 1.0 mol% (Table 3, entries 4-6), PC conversion was quantitative within 21 h. Remarkably, under the latter conditions, prolonging the reaction time (24 h) did not cause any significant diminution of selectivity.

Insert Table 3

The depolymerization of the polycarbonate (Eq. (1)) has been investigated also at a temperature as high as 473 K. Table 4 shows the results obtained at this temperature when using markedly different catalyst loadings (1.0 and ~10 mol%), but keeping constant the other parameters. At 473 K, as expected, PC conversion was by far faster, but, whatever catalyst load was used (1 or 10 mol%), BPA selectivity was more difficult to control due to the higher incidence of the BPA cleavage reaction (Scheme 3). Nevertheless, the diminution of selectivity can be contained within modest limits by controlling the reaction time suitably. For instance, using the higher catalyst load (~10 mol%; see entries 1-3 in Table 4), PC conversion was quantitative within 1 h (vs 6 h, at 433 K), and the selectivity towards BPA was still as high as 91%. However, at this temperature a longer reaction time caused a more significant decrease of BPA yield because of the decomposition of the monomer. An analogous trend was noted when using a La(O₃SCF₃)₃ load equal to 1 mol% (Table 4, entries 4-6).

Insert Table 4

As a comparison, a few other La-based systems, such as $La(NO_3)_3$ $^{\circ}6H_2O$, La_2O_3 and $LaCl_3$, were also investigated as potential catalysts. Under the working conditions summarized in Table 5, both La_2O_3 and $LaCl_3$ showed negligible activity probably because of their poor solubility in the reaction mixture. $La(NO_3)$ $^{\circ}6H_2O$, which was soluble in the reaction medium (THF/H₂O), displayed only moderate catalytic activity. After 4 h at 433 K, the PC conversion was modest (~40%) and BPA was obtained in 23% yield (Table 5, entry 3) with poor selectivity: in addition to minor amounts of PhOH, 4-IPPH and 4-IPP, the formation of a new product was noted, which was identified as the nitric monoester of BPA, HOC₆H₄-C(Me)₂-C₆H₄O-NO₂, on the basis of GCMS analysis (Fig. S13). It is worth noting that, under comparable conditions (Fig. 2, curve c), $La(O_3SCF_3)_3$ promoted selectively the conversion of PC into BPA with 94% yield.

Insert Table 5

Reaction (1) was carried out also in the presence of trifluoromethanesulfonic acid (CF₃SO₃H, triflic acid), a very strong proton donor used in this context as a potential catalyst in place of $M(O_3SCF_3)_3$. Table 6 shows that also CF₃SO₃H can promote reaction (1). However, under comparable experimental conditions, CF₃SO₃H was not only by far less active than Sc(O₃SCF₃)₃ and Yb(O₃SCF₃)₃, but also a less effective and selective catalyst than La(O₃SCF₃)₃. In the presence of CF₃SO₃H the conversion of the polycarbonate was not yet complete after 6 h at 433 K and BPA was

obtained in 50% yield together with significant amounts of PhOH, 4-IPPH, 4- IPP and also 1,4butanediol.⁹

Insert Table 6

At the end of catalytic run the catalyst, $La(O_3SCF_3)_3$, can be recovered with high yield 90-95 %) from the reaction mixture by extraction with diethyl ether/H₂O, as described in detail in Section 2.2.2. The recovered catalyst displayed spectroscopic features (Fig. S10-S12) similar to those of commercial $La(O_3SCF_3)_3$ and its catalytic activity was comparable with that of the fresh catalyst (see 2.2.2).

Also isolation of BPA was quite straightforward. The purification procedure (Section 2.2.1) allowed to isolate as pure compounds, not only the monomer (BPA; Fig. S4-S7) but also the additive used as chain terminator in the synthesis of the feed PC (in the present case, 4-cumylphenol; Fig. S8 and S9).

4. Discussion

The oxophilic nature of Rare Earth triflate salts $M(O_3SCF_3)_3$ (M = Sc, Y, Ln) is firmly-established in literature [37,38], as well as the ability of Sc(OTf)₃, one of the most representative Rare Earth triflates, to coordinate an organic carbonate such as dimethyl carbonate (DMC) through the carbonyl oxygen [(η^1 -O(C=O) coordination], even in the presence of N- and/or O-donors (CH₃CN, amine, THF) [44]. A detailed spectroscopic (NMR, FTIR) study carried out on [(η^1 -O(C=O)– DMC)Sc(O₃SCF₃)₃], the first isolated DMC–metal complex, has also shown that η^1 -O(C=O) coordination of DMC to Sc³⁺ enhances the electropositive character of carbonyl carbon atom and activates it to the attack of nucleophiles (amines, [39-41, 44] for instance).

These remarks are of help in rationalizing a few aspects of the reactivity described in the present work. In fact, in the presence of $M(O_3SCF_3)_3$ (M = Sc, La, Yb) salts, the interaction of O-donors such as THF, H₂O and PC itself with the oxophilic Lewis acid cation M^{3+} (M = Sc, La, Yb) is very likely and may lead not only to the formation of aquo-solvato- complexes $[L_nM(OH_2)]^{3+}$ (L = ligand (H₂O, THF, etc)) but also to the coordination of the metal center to PC through the carbonyl oxygen of C=O groups along the polymeric chain. The latter interaction plays a key role in the catalytic process under study (Eq. (1)), as through it the carbonyl carbons of PC can be made more electrophilic and activated towards the nucleophilic attack by H₂O (Scheme 4).

⁹ Under the working conditions, 1,4-butanediol may form through CF_3SO_3H -assisted opening of THF ring by H₂O. The formation of the diol can be promoted also by $Sc(O_3SCF_3)_3$ (see Table 1, footnote (c) and Table 2, footnote (c)), but the diol was never observed to form in detectable amounts when $Yb(O_3SCF_3)_3$ or $La(O_3SCF_3)_3$ salts were used as catalysts of reaction (1) under the experimental conditions used in this work.

A Lewis acid catalysis agrees with the higher activity displayed by $Sc(O_3SCF_3)_3$ and $Yb(O_3SCF_3)_3$ relative to $La(O_3SCF_3)_3$, that reflects the higher Lewis acidity of $Sc(O_3SCF_3)_3$ and $Yb(O_3SCF_3)_3$ with respect to $La(O_3SCF_3)_3$ [48,49], and may provide an additional explanation for the descending branch of the curve in Fig. 3. In fact, an excessive concentration of a coordinating species such as H_2O in the reaction mixture may repress, to some extent, the interaction of the C=O groups of the polymeric chain with the M^{3+} cations (Scheme 4) and result, as observed, in a lower catalytic activity of the triflate salt.

Insert Scheme 4

An issue strictly related with the presence of $[L_nM(OH_2)]^{3+}$ (L = ligand (H₂O, THF, etc)) aquocomplexes in the reaction medium (THF/H₂O) concerns the potential catalytic role that these species might play in the process by acting as Broensted acid catalysts, as illustrated in Scheme 1 for a generic proton donor AH. In principle, this possibility cannot be excluded a priori (Eq. (4)) [51]. To shed light on this issue, reaction (1) was carried out also in the presence of a very strong

$$\left[L_{n}M(OH_{2})\right]^{3+} \quad \longleftarrow \quad \left[L_{n}M(OH)\right]^{2+} + H^{+} \tag{4}$$

proton donor such as CF_3SO_3H . The obtained results (Table 6) show that H^+ and M^{3+} exhibit quite a different catalytic behavior and bring to exclude that protons arising from $[L_nM(OH_2)]^{3+}$ aquocomplexes (weaker proton donors than CF_3SO_3H) may play a catalytically important role in promoting reaction (1).

The triflic salts can also activate the monomer (BPA) towards the cleavage of the propylidene bridge (Table 2) with formation of undesired side-products such as phenol, 4-IPP and 4-IPPH. The incidence of this side-reaction is particularly important with the stronger Lewis acids $Sc(O_3SCF_3)_3$ and $Yb(O_3SCF_3)_3$ rather than the weaker La $(O_3SCF_3)_3$ [52,53]. This fact, as well as the catalytic activity observed under anhydrous conditions (Entry 3, 5 and 7 in Table 2), suggest that the metal center can act as a Lewis acid catalyst also in this process, probably by coordinating a OH group of the substrate (BPA), as shown in Scheme 5. O-Coordination of BPA to the metal center may open Insert Scheme 5

the way to the cleavage of propylidene bridge (*a*) by weakening the bond C2-C1' (path a, Scheme 5), as formerly proposed by Schnell and Krimm for the analogous addition of proton to one of the OH groups of BPA [40], or (*b*) through path b (Scheme 5), which is reminiscent of the mechanistic pathway more recently proposed by Hunter and Sauvage [40]. The presence of water in the reaction mixture has an inhibitory effect on the bridge cleavage reaction (Entry 2, 4 and 6 in Table 2). This may be the result of a competition between water and BPA molecules for the coordination sites of the metal center (Eq. (7)). In principle, aquo-complexes[L_nM(OH₂)]³⁺ might act also as

Insert equation 7

proton donors towards the free substrate (Eq. (8)) with generation of protonated species B which might behave likewise A in Scheme 5. However, the inhibitory effect observed under non anhydrous conditions suggests that aquo-complexes $[L_nM(OH_2)]^{3+}$ cannot be so effective catalytically active species as L_nM -BPA adducts (Scheme 5).

Insert equation 8

5. Conclusions

Eco-friendly water tolerant Lewis acids such as the triflate salts $M(O_3SCF_3)_3$ (M = Sc, Yb, La) can promote effectively the depolymerization of poly-(bisphenol A carbonate) under hydrolytic conditions. The process is relevant to the chemical recycling and valorization of waste PC. The nature of the metal center affects the yield and selectivity of the process critically. In THF as cosolvent able to dissolve the polymer, La(O₃SCF₃)₃, a weaker Lewis acid than Sc(O₃SCF₃)₃ and Yb(O₃SCF₃)₃, promoted the depolymerization of the polymer with excellent BPA yield and selectively, as, unlike the other considered triflates, it displayed also very poor tendency to catalyze, under the working conditions, the cleavage of the propylidene bridge of the monomer (BPA).

The behavior of the system in the presence of the La-catalyst has been focused. The H₂O/PC weight ratio influences the BPA productivity significantly: as an excessive amount of H₂O in the reaction mixture diminishes the solubility of PC in the reaction medium and also restrains the activity of the Lanthanum salt as Lewis acid catalyst as, at higher concentrations, H₂O, by acting as a ligand, can more effectively compete with PC for the coordination sites on the metal center. The depolymerization can proceed under not severe temperature conditions (433 K) in the presence of moderate catalyst loads (1-10 mol%). Rising temperature increases the conversion rate of PC but also causes the diminution of BPA selectivity, which, however, can be kept in modest limits by suitably controlling the reaction time. For instance, at 433 K, using a H₂O/PC weight ratio close to unity and a catalyst load around 10 mol%, quantitative and selective (97%) depolymerization of the used polycarbonate was achieved within 6 h. At 473 K, under otherwise analogous reaction conditions, PC conversion was complete within 1 h, while diminution of BPA selectivity (91%) was substantially modest.

The process does not require complex equipments, is simple from the operational point of view, and does not make use of hazardous solvents as reaction medium [54]. THF used as the cosolvent

can be easily removed and recovered at the end of the process. Also the catalyst, $La(O_3SCF_3)_3$, can be recovered quantitatively at the end of the reaction and reused (Scheme 6). The recovered catalyst showed

Insert Scheme 6

spectroscopic and catalytic features similar to those of the fresh catalyst. The above features, as well as the high productivity/selectivity and environmentally benign nature of the catalyst [38], are important advantages of the described process.

Finally, the catalytic behavior of the used $M(O_3SCF_3)_3$ salts has been compared with that of a strong proton donor such as CF_3SO_3H . The results obtained allow to exclude that protons arising from $[L_nM(OH_2)]^{3+}$ (L = ligand (H₂O, THF, etc)) aquo-complexes may play a catalytically important role in promoting reaction (1).

Acknowledgements

This work was financially supported from Università degli Studi "A. Moro" di Bari (Fondi di Ateneo). Dr F. Iannone and Dr. M. Glaciale are acknowledged for their assistance in a few experiments.

References

- [1] F. La Mantia (Ed.), Handbook of Plastic Recycling, Rapra Technology, Shrewsbury, 2002.
- [2] J. Hopewell, R. Dvorak, E. Kosior, Phil. Trans. R. Soc. B 364 (2009) 2115-2126.
- [3] I.A. Ignatyev, W. Thielemans, B.V. Beke, ChemSusChem 7 (2014) 1579-1593.
- [4] E. Feghali, T. Cantat, ChemSusChem 8 (2015) 980-984.
- [5] HIS Markit, Newsroom, http://press.ihs.com/press-release/commodities-pricingcost/after-major-downturn-global-demand-polycarbonate-growing-agai, (August 2016).
- [6] E.V. Antonakou, D.S. Achilias, Waste Biomass Valor. 4 (2013) 9-21.
- [7] Mordor Intelligence, <u>http://www.mordorintelligence.com/industry-reports/pc-</u>polycarbonate-market?gclid=CKmv1NeW684CFasp0wod_r0F7w, (August 2016).
- [8] D.S. Achilias, E.V. Antonakou, E. Koutsokosta, A.A. Lappas, J. Appl. Polym. Sci. 114 (2009) 212-221.
- [9] L. Rosi, M. Bartoli, A. Undria, M. Frediani, P. Frediani, J. Mol. Catal. A: Chemical 408 (2015) 278-286.
- [10] A. Oku, S. Tanaka, S. Hata, Polymer 41 (2000) 6749-6753.

- [11] L.-C. Hu, A. Oku, E. Yamada, Polymer 39 (1998) 3841-3845.
- [12] H. Jie, H. Ke, Z. Qing, C. Lei, W. Yongqiang, Z. Zibin, Polym. Degrad. Stabil. 91 (2006) 2307-2314.
- [13] F. Liu, L. Li, S. Yu, Z. Li, X. Ge, J. Hazard. Mater. 189 (2011) 249-254.
- [14] C.-H. Lin, H.-Y. Lin, W.-Z. Liao, S.A. Dai, Green Chem. 9 (2007) 38-43.
- [15] R. Piñero, J. García, M. J.Cocero, Green Chem. 7 (2005) 380-387.
- [16] M.M.A. Nikje, Polimery 56 (2011) 381-384.
- [17] D. Kim, B. Kim, Y. Cho, M. Han, B.-S. Kim, Ind. Eng. Chem. Res. 48 (2009) 6591-6599.
- [18] D. Kim, B. Kim, Y. Cho, M. Han, B.-S. Kim, Ind. Eng. Chem. Res. 48 (2009) 685-691.
- [19] S. Hata, H. Goto, E. Yamada, A. Oku, Polymer 43 (2002) 2109-2116.
- [20] S. Hata, H. Goto, S. Tanaka, A. Oku, J. Appl. Polym. Sci. 90 (2003) 2959-2968.
- [21] Z. Pan, I.-M. Chou, R.C. Burrus, Green Chem. 11 (2009) 1105-1107.
- [22] M. Watanabe, Y. Matsuo, T. Matsushita, H. Inomata, T. Miyake, K. Hironaka, Polym. Degrad. Stabil. 94 (2009) 2157–2162.
- [23] Z. Pan, Z. Hu, Y. Shi, Y. Shen, J. Wang, I-M. Chou, RSC Adv. 4 (2014) 19992-19998.
- [24] F.-S. Liu, Z. Li, S.-T. Yu, X. Cui, C.-X. Xie, X.-P. Ge, J. Polym. Environ. 17 (2009) 208–211.
- [25] G. Grause, K. Sugawara, T. Mizoguchi, T. Yoshioka, Polym. Degrad. Stabil. 94 (2009) 1119–1124.
- [26] T. Yoshioka, K. Sugawara, T. Mizoguchi, A. Okuwaki, Chem. Lett. 34 (2005) 282-283.
- [27] K. Hatakeyama, T. Kojima, T. Funakuzuri, J. Mater. Cycles Waste Manag. 16 (2014) 124-130.
- [28] A. Ikeda, K. Katoh, H. Tagaya, J. Mat. Sci. 43 (2008) 2437-2441.
- [29] G. Grause, R. Karrbrant, T. Kameda, T. Yoshioka, Ind. Eng. Chem. Res. 53 (2014) 4215-4223.
- [30] M. Taguchi, Y. Ishikawa, S. Kataoka, T. Naka, T. Funazukuri, Catal. Commun. 84 (2016) 93-97.
- [31] G.P. Tsintzou, D.S. Achilias, Waste Biomass Valor. 4 (2013) 3-7.
- [32] X. Song, F. Liu, L. Li, X. Yang, S. Yu, X. Ge, J. Hazar. Mater. 244-245 (2013) 204-208.
- [33] N. Deirram, A.R. Rahmat, APCBEE Procedia 3 (2012) 172-176.
- [34] M.M.A. Nikje, M. Askarzadeh, Progr. Rubb. Plast. Recycl. Techn. 29 (2013) 169-176.
- [35] K. Troev, R. Tsevi, I. Gitsov, Polymer 42 (2001) 39-42.
- [36] G.D. Cooper, B. Williams, J. Org. Chem 27 (1962) 3717-3720.

- [37] S. Kobayashi, M. Sugiura, H. Kitagawa, W.W.-L. Lam, Chem. Rev. 102 (2002) 2227-2302.
- [38] V. Ladziata, ARKIVOC (2014(i)) 307-336.
- [39] M. Distaso, E. Quaranta, Tetrahedron 60 (2004) 1531-1539.
- [40] M. Distaso, E. Quaranta, Appl. Catal. B: Environ. 66 (2006) 72-80.
- [41] M. Distaso, E. Quaranta, J. Catal. 228 (2004) 36-42.
- [42] M. Carafa, E. Quaranta, Mini-Rev. Org. Chem. 6 (2009) 168-183.
- [43] E. Quaranta, M. Aresta, The Chemistry of N-CO₂ bonds: Synthesis of Carbamic Acids and their Derivatives, Isocyanates, Ureas, in: M. Aresta (Ed.), Carbon Dioxide as Chemical Feedstock, Wiley-VCH Verlag GmbH & Co.KGaA, Weinheim, 2010, pp. 121-167.
- [44] M. Distaso, E. Quaranta, J. Catal. 253 (2008), 278-288
- [45] D.D. Perrin, W.L.F. Armarego, D.R. Perrin, Purification of Laboratory Chemicals, Pergamon, Oxford, 1986.
- [46] D.J. Brunelle, P.M. Smigelski Jr., E.P. Boden, ACS Symposium Series Vol. 898 (2005) Chapter 2 (Advances in Polycarbonates) pp. 8-21.
- [47] R. Ullah, I. Ahmad, Y. Zheng, Journal of Spectroscopy, Volume 2016 Article ID 2073613, <u>http://dx.doi.org/10.1155/2016/2073613</u>.
- [48] R. Ullah, H. Li, YM. Zhu, J. Mol. Struct. 1059 (2014) 255-259.
- [49] H. Schnell, H.Krimm, Angew. Chem. Int. Edit. 2 (1963) 373-379.
- [50] S.E. Hunter, P.E. Sauvage, J. Org. Chem. 69 (2004) 4724-4731.
- [51] S. Kobayashi, SynLett (1994) 689-701.
- [52] H. Tsuruta, K. Yamaguchi, T. Imamoto, Chem. Commun. (1999) 1703-1704.
- [53] H. Tsuruta, K. Yamaguchi, T. Imamoto, Tetrahedron 59 (2003) 10419-10438.
- [54] D. Prat, A. Wells, J. Hayler, H. Sneddon, C.R McElroy, S.Abou-Shelada, P.J. Dunn, Green Chem. 18 (2016) 288-296.

LIST OF EQUATIONS

equation 1

equation 7

equation 8

LIST OF SCHEMES

Scheme 1. Hydrolysis of poly-(bisphenol A carbonate): Broensted acid catalysis.

Scheme 2. Schematic sketch of the polymeric chain: the repeating unit (UR) and the nature of the species, 4-cumylphenol, used as chain-terminal stopping agent in the synthesis of the used polymer have been highlighted. Accordingly, it may be assumed that a mass w(g) of feed PC consists formally of X mol of repeating units (UR) and Y mol of bis(4-cumylphenyl) carbonate (see later Eq. (3")).

Scheme 3. Decomposition of BPA.

Scheme 4. Hydrolysis of poly-(bisphenol A carbonate) promoted by M(O₃SCF₃)₃: Lewis acid catalysis.

Scheme 5. Possible reaction pathways for M(O₃SCF₃)₃ catalyzed cleavage of BPA.

Scheme 6. Hydrolysis of poly-(bisphenol A carbonate) promoted by M(O₃SCF₃)₃.

LIST OF FIGURES

Fig.1. Hydrolysis of poly-(bisphenol A carbonate) in the presence of $La(O_3SCF_3)_3$. FTIR spectrum (CaF₂ windows) of (a) the reaction solution at the end of the catalytic run; (b) the used polycarbonate in THF solution.

Fig. 2. Hydrolysis of poly-(bisphenol A carbonate) in the presence of $M(O_3SCF_3)_3$ (M = Sc (a), Yb (b), La (c)): influence of reaction time. THF, 6 mL; PC, ~0.5 g; H₂O/PC, ~0.98 m_{H2O}/m_{PC} (~14 mol_{H2O}/mol_{CO3});² catalyst load, ~10.2 mol% ((mol_{M(O3SCF3)3}/mol_{CO3})×100);² T, 433 K.

Fig. 3. Hydrolysis of poly-(bisphenol A carbonate) in the presence of La(O₃SCF₃)₃: influence of H₂O/PC weight ratio. THF, 6 mL; PC, ~0.5 g; catalyst load, ~10 mol% (($mol_{La(O3SCF3)3}/mol_{CO3}$) ×100);² T, 433 K; reaction time, 4 h.



equation 1



equation 7



equation 8



Fig.1. Hydrolysis of poly-(bisphenol A carbonate) in the presence of $La(O_3SCF_3)_3$. FTIR spectrum (CaF₂ windows) of (a) the reaction solution at the end of the catalytic run; (b) the used polycarbonate in THF solution.



Fig. 2. Hydrolysis of poly-(bisphenol A carbonate) in the presence of $M(O_3SCF_3)_3$ (M = Sc (a), Yb (b), La (c)): influence of reaction time. THF, 6 mL; PC, ~0.5 g; H₂O/PC, ~0.98 m_{H2O}/m_{PC} (~14 mol_{H2O}/mol_{CO3});² catalyst load, ~10.2 mol% ((mol_{M(O3SCF3)3}/mol_{CO3})×100);² T, 433 K.



Fig. 3. Hydrolysis of poly-(bisphenol A carbonate) in the presence of La(O_3SCF_3)₃: influence of H₂O/PC weight ratio. THF, 6 mL; PC, ~0.5 g; catalyst load, ~10 mol% ((mol_{La(O3SCF3)3}/mol_{CO3}) ×100);² T, 433 K; reaction time, 4 h.



Scheme 1. Hydrolysis of poly-(bisphenol A carbonate): Broensted acid catalysis.



Scheme 2. Schematic sketch of the polymeric chain: the repeating unit (UR) and the nature of the species, 4-cumylphenol, used as chain-terminal stopping agent in the synthesis of the used polymer have been highlighted. Accordingly, it may be assumed that a mass w(g) of feed PC consists formally of X mol of repeating units (UR) and Y mol of bis(4-cumylphenyl) carbonate (see later Eq. (3")).



Scheme 3. Decomposition of BPA.



Scheme 4. Hydrolysis of poly-(bisphenol A carbonate) promoted by M(O₃SCF₃)₃: Lewis acid catalysis.



Scheme 5. Possible reaction pathways for $M(O_3SCF_3)_3$ catalyzed cleavage of BPA.



Scheme 6. Hydrolysis of poly-(bisphenol A carbonate) promoted by M(O₃SCF₃)₃.

Table 1 Hydrolysis of poly-(bisphenol A carbonate) (PC) in the presence of a few Rare Earth triflate salts $M(O_3SCF_3)_3$ (M = Sc, Yb, La)^a

50, 10,	D u)						
entry	catalyst	H ₂ O/PC	Catalyst load	Т	reaction	BPA	PC
		$m_{\rm H2O}/m_{\rm PC};$	$(m_{catalyst}/m_{PC}) \times 100;$	(K)	time	GC-yield	conversion ^b
		mol _{H2O} /mol _{CO3}	$(mol_{catalyst}/mol_{CO3}) \times 100$		(h)	(%)	(%)
1 ^c	$Sc(O_3SCF_3)_3$	0.964;	19.7;	433	6	14	100
		13.8	10.3				
2	$La(O_3SCF_3)_3$	0.973;	23.3;	433	6	97	100
		13.9	10.3				
3	$Yb(O_3SCF_3)_3$	0.991;	24.5;	433	6	68	100
		14.2	10.2				

^a THF (solvent), 6 mL; PC, ~0.5 g. ^b As ascertained by FTIR of the reaction mixture (see Section 3.1). ^c The GC analysis of the reaction mixture displayed also the formation of minor amounts of 1,4-butanediol.

entry	catalyst	H ₂ O/BPA	Catalyst load	Т	reaction	BPA conversion (GC)
		mol_{H2O}/mol_{BPA}	$(mol_{catalyst}/mol_{BPA}) \times 100$	(K)	time	(%)
					(h)	
1	none	13.8	0	433	4	b
2^{c}	$Sc(O_3SCF_3)_3$	13.9	10.0	433	4	82
3	$Sc(O_3SCF_3)_3$	0	10.2	433	4	100
4	Yb(O ₃ SCF ₃) ₃	14.0	10.2	433	4	32
5	Yb(O ₃ SCF ₃) ₃	0	10.1	433	4	89
6	La(O ₃ SCF ₃) ₃	13.3	10.0	433	4	3
7	La(O ₃ SCF ₃) ₃	0	9.5	433	4	42

Table 2 Decomposition of BPA in the presence of $M(O_3SCF_3)_3$ salts (M = Sc, Yb, La).^a

^a THF (solvent), 6 mL; BPA, ~2.1 mmol.

^b The analysis of the reaction mixture did not show any evidence of BPA conversion into PhOH, 4-IPPH, 4-IPP or other products.

^c The GC analysis of the reaction mixture displayed also the formation of minor amounts of 1,4butanediol.

Table 3

Т entry H₂O/PC Catalyst load BPA PC Conversion t m_{H2O}/m_{PC} ; $(m_{catalyst}/m_{PC}) \times 100;$ (K) (h) GC Yield (%) mol_{H2O}/mol_{CO3} ($mol_{catalyst}/mol_{CO3}$)×100 (%) 100^b 1 23.3; 433 97 0.973; 6 13.9 10.3 с 2 79 0.980; 11.6; 433 8 5.1 14.0 100^{b} 3 0.978; 11.4; 433 12 96 5.0 14.0 с 4 0.971; 2.3; 433 18 85 14.0 1.0 100^{b} 5 0.987; 2.3; 433 21 97 14.1 1.0 100^{b} 6 0.975; 2.3; 433 24 97 13.9 1.0

Hydrolysis of poly-(bisphenol A carbonate) in the presence of La(O₃SCF₃)₃: influence of catalyst loading.^a

^a THF (solvent), 6 mL; PC, ~0.5 g.

^b As ascertained by the FTIR spectrum of the reaction solution.

^c PC conversion was not yet quantitative, as supported by the FTIR spectrum of the reaction solution showing the absorption of unreacted PC at 1778 cm⁻¹ (vC=O).

Hydrolysis of poly-(bisphenol A carbonate) in the presence of La(O ₃ SCF ₃) ₃ at 473 K. ^a									
entry	H ₂ O/PC	Catalyst load	Т	t (h)	BPA	PC Conversion			
	$m_{H2O}/m_{PC};$	$(m_{catalyst}/m_{PC}) \times 100;$	(K)		GC Yield	(%)			
	mol _{H2O} /mol _{CO3}	$(mol_{catalyst}/mol_{CO3}) \times 100$			(%)				
1	0.981;	23.1;	473	0.75	60	71			
	14.0	10.1							
2	0.980;	22.9;	473	1	91	100 ^b			
	14.0	10.1							
3	0.981;	23.1;	473	2	82	100 ^b			
	14.0	10.1							
4	0.984;	2.3;	473	3	74	b			
	14.1	1.0							
5	0.990;	2.3;	473	4	92	100 ^b			
	14.1	1.0							
6	0.979;	2.3;	473	6	76	100 ^b			
	14.0	1.0							

Table 4

^a THF (solvent), 6 mL; PC, ~0.5 g. ^b As ascertained by the FTIR spectrum of the reaction solution. ^c PC conversion was not yet quantitative, as supported by the FTIR spectrum of the reaction solution showing the absorption of unreacted PC at 1778 cm⁻¹ (vC=O).

entry	La catalyst	H ₂ O/PC	Catalyst load	Т	reaction time	BPA
		$m_{H2O}/m_{PC};$	$(m_{catalyst}/m_{PC}) \times 100;$	(K)	(h)	GC yield
		mol _{H2O} /mol _{CO3}	$(mol_{catalyst}/mol_{CO3}) \times 100$			(%)
1	LaCl ₃	0.967;	14.7;	433	4	<1
		13.8	15.4			
2	La_2O_3	0.969;	12.7;	433	4	~1
		13.8	10.0			
3	La(NO ₃) ₃ ⁻⁶ H ₂ O	0.982;	17.2;	433	4	23
		14.0	10.3			

Table 5
Hydrolysis of poly-(bisphenol A carbonate) in the presence of other La-based systems. ^a

^a THF (solvent), 6 mL; PC,~ 0.5 g.

Table of Hydrolysis of poly-(displicitor A carbonate). W(035CF3)3 vs CF35O311 catalytic activity								
Entry	Catalyst	H ₂ O/PC	Catalyst load	Т	t	BPA	PC	
		$m_{H2O}/m_{PC};$	$(m_{catalyst}/m_{PC}) \times 100;$	(K)	(h)	GC yield	conversion	
		mol _{H2O} /mol _{CO3}	$(mol_{catalyst}/mol_{CO3}) \times 100$			(%)	(%)	
1	CF ₃ SO ₃ H	0.986;	6.0;	433	6	50	90	
		14.1	10.0					
2	$Sc(O_3SCF_3)_3$	0.980;	19.6;	433	2	28	~100	
		14.0	10.2					
3	$La(O_3SCF_3)_3$	0.973;	23.3;	433	6	97	100	
		13.9	10.3					
4	$Yb(O_3SCF_3)_3$	0.973;	24.4;	433	2	83	~100	
		13.9	10.2					

Table 6 Hydrolysis of poly-(bisphenol A carbonate): M(O₃SCF₃)₃ vs CF₃SO₃H catalytic activity^a

^a THF (solvent), 6 mL; PC, ~0.5 g; T, 433 K.