Acid functionalized carbons as catalyst for glycerol etherification with benzyl alcohol

María E. Chiosso1 · Ileana D. Lick1 · Mónica L. Casella1 · Andrea B. Merlo1

Received: 14 December 2018 / Revised: 7 August 2019 / Accepted: 12 September 2019 / Published online: 7 January 2020
© Associação Brasileira de Engenharia Química 2020

Abstract

In this paper, the catalytic behaviour of two different carbonaceous systems, commercial carbon (Cc) and synthetic carbon (Cs) functionalized with concentrated sulfuric acid (–SO3H) and with reduced aryl diazonium salt (–PhSO3H), was studied in the etherification of glycerol (Gly) with benzyl alcohol (BA). The catalytic activity and selectivity were studied varying the catalyst percentage (5, 10 and 15 wt%) and the initial reactant mass ratio. Taking into account the results obtained, the experimental conditions selected to continue with the catalytic studies were: temperature 393 K, Gly:BA molar ratio 3:1 and a catalyst loading of 10 wt%. Mono- and diethers were the main products. For the Cc-based catalysts, higher conversion (66%) and selectivity (90%) were obtained with the most acidic system, Cc–PhSO3H. Changing the carbon functionalization method (–PhSO3H for –SO3H groups) produced a considerable increase in glycerol conversion (from 23 to 66%) and in selectivity to ME + DE (from 62 to 90%). On the other hand, for both Cs-based systems a great increase in performance was obtained, about 95% conversion after 360 min of reaction. Monoether was the major product of the reaction with a selectivity of 79% for Cs–SO3H and 87% for Cs–PhSO3H, which was maintained with reuse.

Graphic abstract

Etherification of glycerol (Gly) with benzyl alcohol (BA) with two carbonaceous systems synthesized in the laboratory and functionalized with –SO3H and –PhSO3H.

Keywords Glycerol · Etherification · Acid catalysts · Sulfonated carbon · Benzyl alcohol

Introduction

Current problems of the energetic crisis due to the depletion of fossil fuels, high oil prices and the effects on the environment caused by the emission of hazardous and chemicals products into the atmosphere have forced us to look for alternatives to replace these fuels. Biodiesel is a renewable and biodegradable alternative to fossil fuels, and because of this, it is expected that its use will continue growing in the future (Anitha et al. 2016; Bozkurt 2015).
Biodiesel produced from vegetable oil, animal tallow, oils and fats is a promising biofuel to be mixed with diesel or used directly. However, one of the main obstacles to the commercialization of biodiesel is the by-product glycerol, since each ton of biodiesel originates 100 kg of glycerine that saturates the market, causing a dramatic reduction in prices and, consequently, negatively effects biodiesel economy (Anitha et al. 2016; Janaun and Ellis 2010).

The direct combustion of glycerol is not feasible in engines because of its high viscosity, low heating value, high auto-ignition temperature and corrosive salt content (for crude glycerol). Moreover, its incomplete combustion can be carcinogenic due to the production of acroleine (Bohon et al. 2011).

Therefore, in recent years several processes to produce value-added chemicals from glycerol have been reported, such as hydrogenolysis, etherification, esterification, dehydration, oxidation and others (Len and Luque 2014; Faroppa et al. 2016; Jaworski et al. 2015; Pagliaro et al. 2007; Alonso et al. 2012).

A wide range of chemical products can be derived from glycerol, making sure that in the future glycerol will become an important renewable raw material for biorefineries. Although researchers have developed various methods for using glycerol as fuel additives, these methods are still at the laboratory scale and their actual cost for production is unknown.

The etherification process of glycerol is an interesting application that could be industrially important to obtain compounds of lower viscosity and higher volatility than glycerol, which could be used as oxygenated additives for liquid fuels, intermediates for pharmaceutical products, agrochemicals, nonionic surfactants, etc. (Gonçalves et al. 2013, 2016). As a function of the alcohol used, selective glycerol etherification may also lead to obtaining a wide range of products having biological applications, such as anti-inflammatory, antibacterial, antifungal and antitumor products, among others (Gu et al. 2008).

Glycerol etherification with benzyl alcohol (BA) is an acid-catalyzed reaction, resulting in a mixture of three products. Besides, a by-product of BA self-condensation (dibenzyl ether, BE), can also be formed (Scheme 1) (Jaworski et al. 2015).

This reaction can be carried out using homogeneous or heterogeneous acid catalysts.

There are several reports of acid catalysts used in the etherification of glycerol with benzyl alcohol. Gu et al. (2008) obtained 73% and 96% yield with the catalyst SiO$_2$–SO$_3$H (under conditions Gly:BA = 1:1 and 4.2:1, respectively) and 100% of ME. In that publication the results with a C–SO$_3$H system were also shown, but only 12% yield was reported (Gu et al. 2008). Da Silva et al. (2009) obtained the best performance with the systems Amberlyst-35 (37% ME, 20% DE, 3% TE and 30% BE) and Zeolite β (55% ME, 3% DE, 1% TE and 38% BE), for a 100% conversion after 120 min of reaction. Conversion of 70% of BA was reported (for 360 min of reaction) using 2.5% of sulfated zirconia (2S/ZrO$_2$) as catalyst and the distribution of products was the following: 40% S ME, 30% S DE and 30% S BE (Jaworski et al. 2015). On the other hand, Zeolite Z15c as catalyst for the etherification of Gly with BA (ratio 1:1 and 393 K) produced ME (60%) and DE (38%), with little BE formation (2%), for 80% of BA conversion after 8 h reaction (Gonzalez-Arellano et al. 2015).

**Scheme 1** Reaction network for glycerol etherification with benzyl alcohol
Carbonaceous materials are a good alternative. Scientific production with these materials during the last 20 years has been increasing considerably (González-García 2018).

The surface chemistry of carbonaceous materials determines their physicochemical and acid–base properties, and hydrophobic characteristics. This is defined by the presence of different atoms of carbon and the formation of functional groups. Oxygenated groups are generated in large quantities as a result of the chemical reaction between the active centres of the carbon with the oxygen coming from the precursor material or any other oxidizing agent present in the reaction medium (Martínez Fierro 2012). Of the functional groups that can be found on the surface of activated carbon, carboxylic acids, lactones, anhydrides or phenols confer an acidic character. Nevertheless the presence of groups such as quinones, pyrones and chromenes give it a basic character. In general, the surface of active carbon is hydrophobic; however, the presence of acidic oxygenated groups gives the surface a certain hydrophilic character (Boehm 1994).

Among the supports and catalysts used in the etherification of glycerol is also carbon. The most direct way to obtain a carbonaceous material is by pyrolysis of carbohydrates, such as glucose or sucrose, at relatively mild temperature (573–673 K) in an inert atmosphere. Biomass resources can also be used (Gonçalves et al. 2013, 2014, 2016; Rodríguez-Reinoso 1998; Han et al. 2003). Among these materials, those functionalized with sulfonic acid groups have been investigated as potential catalysts (Toda et al. 2005; Janaun and Ellis 2010; Sánchez et al. 2011; Liu et al. 2008, 2013).

In this work, carbon acid catalysts were prepared from dextrose and functionalized with sulfonic acid groups to be used in the etherification of glycerol with benzyl alcohol. The effect of several operational variables (percentage of catalyst, Gly:BA molar ratio) as well as the reusability of the catalysts, have been evaluated. There are many reports of glycerol etherification using acidic carbons, but there are very few papers where benzyl alcohol is used and the results are not good (Gu et al. 2008). Because of this, our research addresses the challenge of finding an acid carbonaceous system that will be an efficient catalyst in this reaction. Therefore, the main objective of this work was to obtain (with good performance and low cost) a functionalized carbonaceous system, comparable to a commercial carbon in efficiency in the etherification reaction.

Materials and methods

**Reagent for C synthesis:** dextrosa (anhydrous, Aendra Por-analysis), sodium silicate solution (J.T. Baker 98% SiO₂), hydrochloric acid (Aendra analytic reagent), ethyl alcohol (Baker analy-sed), ethyl alcohol (Cicarelli 99.5% absolute).

**Reagent for functionalization of carbon:** sulfuric acid (J.T. Baker 98%), 4-aminobenzenesulfonic acid (Biopack Pro-analysis), hypophosphorous acid (Biopack 50% Pro-analysis).

**Etherification Reagent:** glycerol (Biopack 99.5%), benzyl alcohol (Aendra analytic reagent), 1-butanol (Baker analysed), ethyl alcohol (Cicarelli 99.5% absolute).

**Catalyst preparation**

NORIT C (GAC 1240 PLUS), which was ground to a particle size of 60 to 100 mesh, was used as commercial carbon catalyst (Cc). The synthesized carbon catalyst (Cs) was obtained from dextrose, following similar procedure that published by Han et al. (2003). This method comprises a template-assisted carbonization process for the synthesis of the carbonaceous material. The anhydrous dextrose was used as carbon precursor (15 g) and the commercial sodium silicate solution (20 mL) as source of SiO₂ to develop the template. The key step in the procedure is the co-polymeriza-tion of dextrose with the sodium silicate. After complete dissolution, hydrochloric acid was added, producing a brown solution. The reaction mixture was kept for 24 h at 343 K to achieve vaporization of residual water and polymerization. Carbonization of the material was performed under a nitrogen atmosphere in two steps: the first one at 473 K and the second one at 723 K, both for a period of 5 h. The resulting material was stirred in 3 M NaOH for 5 h to remove the silica “template”. Then, the carbon was recovered by filtration and dried in an oven at 378 K.

**Carbon functionalization**

Two methods for the sulfonation of the carbon materials were used. In the first, the solid was contacted with concentrated H₂SO₄, using a ratio of 15 mL of acid per gram of solid, at 423 K and under a N₂ current. Then, the sulfonated material was washed with distilled water, filtered and dried for 2 h at 378 K (C–SO₃H). The second method consists of the reaction between 4-benzene-diazonium sulfonate and the carbon material, using hypophosphorous acid (H₃PO₂) as the reducing agent (Liu et al. 2010). The synthesis of 4-benzene-diazonium sulfonate was carried out by diazotization of 4-aminobenzenesulfonic acid. The C–PhSO₃H obtained was filtered and washed with distilled water (Ph: phenyl group).

**Catalyst characterization**

The total acid site density was determined by back titration. In a typical test, 50 mL NaOH were added to 0.1 g of dried carbon and kept under stirring for 30 min at room temperature. Subsequently, the excess NaOH was titrated with HCl.
solution until the endpoint. This analysis was performed on carbon samples both with and without functionalization.

The surface properties and porosity of the carbonaceous materials were analyzed by isothermal adsorption/desorption of N$_2$ at 77 K in a Micromeritics Accusorb 2100E equipment, Scanning Electron Microscopy (SEM) was performed using a Philips SEM 505. The carbon samples were placed inside the microscope chamber and evacuated to high vacuum. Energy dispersive X-ray spectroscopic (EDX) analysis was performed to detect the presence of sulphur in the functionalized systems.

Fourier transform infrared spectroscopy (FTIR) was recorded in a Jasco 4200 spectrometer in transmittance mode, on samples embedded in KBr pellets. The spectra were obtained between 400 and 4000 cm$^{-1}$.

Thermogravimetric analysis (TGA) was performed using a Shimadzu TGA-50 thermal analyzer. Sample mass was recorded during heating at a constant temperature rate of 10 K min$^{-1}$, from 298 to 973 K, under a N$_2$ flow of 100 mL min$^{-1}$.

**Catalytic tests**

Etherification of Gly with BA was conducted under a N$_2$ atmosphere, at 393 K and atmospheric pressure, with constant stirring in a glass reactor. The reaction time was 360 min. Cc–SO$_3$H was selected to optimize the Gly:BA molar ratio (1:1, 3:1 or 5:1) and the mass of catalyst in relation to the mass of glycerol employed (5, 10 or 15 wt%). The sample was taken at the end of the reaction, filtered and analysed by gas chromatography. The analyses were carried out using a Varian CP WAX 52 CB (30 m, 0.53 mm i.d.) column.

The response factors of Gly and BA were determined using butyl alcohol as internal standard for the calibration curve. Commercial (+) 3-benzylol-1,2-propanediol and 1,3-dibenzylol-2-propanol were employed to obtain the corresponding response factors. EB was calibrated as well.

The reaction products were identified using a GC/MS HP 5890 gas chromatograph equipped with a capillary column CP CP 3800 gas chromatograph equipped with a capillary column CP WAX 52 CB (30 m, 0.53 mm i.d.) and a FID detector.

The response factors of Gly and BA were determined using butyl alcohol as internal standard for the calibration curve. Commercial (+) 3-benzylol-1,2-propanediol and 1,3-dibenzylol-2-propanol were employed to obtain the corresponding response factors. EB was calibrated as well.

The reaction products were identified using a GC/MS HP 5890 gas chromatograph equipped with a capillary column CP WAX 52 CB (30 m, 0.53 mm i.d.) column.

For reuse tests, the catalysts, in a typical experiment after finishing the etherification reaction, were filtered, washed with water, dried in an oven at 378 K for 24 h, and reused under these conditions: 393 K, 10% of catalyst and Gly:BA molar ratio 3:1.

**Results and discussion**

**Catalyst characterization**

Table 1 shows the results obtained for surface analysis and acid site density of the materials used. The amount of acid sites is very important in the glycerol etherification reaction. In particular, sulfonic groups play a crucial role as active sites for such reaction. In this sense, Table 1 shows that the acidic sites of both carbons increase significantly after functionalization, reaching the highest values for synthesized carbon (Cs–SO$_3$H and Cs–PhSO$_3$H). The high density of sulfonic groups can be explained by the relatively soft carbonization temperature used (723 K), which produces an amorphous but more active surface towards the sulfonation (Sánchez et al. 2011). It can also be seen that, when comparing Cs and Cc carbons, a higher acid density is obtained with the first one (0.4 vs. 0.05 mmol g$^{-1}$), a fact that can be attributed to the additional contribution of –COOH and –OH groups generated as products of the incomplete carbonization of the starting sugar. Strong sulfonation can also oxidize aliphatic CH$_3$/CH$_2$ groups to carboxylic acid groups, which could contribute to explain the increase in acid site density (Mo et al. 2008).

The N$_2$ adsorption/desorption results shown in Table 1 are in agreement with those published by other authors. In the case of Cs, the use of Na$_2$SiO$_3$ plays a very important role as a “director” of the pore structure. This Table also shows that, after functionalization treatment, the surface area and V$_p$ decreased for both carbon systems (Cc and Cs). These results show that sulfonic groups were introduced successfully on the surface (Sánchez et al. 2011).

Figure 1 shows the SEM micrographs and the EDX measurements confirm the presence of sulfur on both catalysts. For both functionalized Cs systems, the micrographs depict rather small carbon aggregates with flat surfaces, according to the low surface area values measured. The EDX measurements confirm the presence of sulfur on both catalysts. For the Cc–SO$_3$H and Cc–PhSO$_3$H systems, although not shown, similar diagrams were obtained (Goncalves et al. 2015; Tao et al. 2015).

FTIR spectra were recorded in order to confirm that the carbonization and sulfonization processes were successful.

| Catalyst      | Acid density (mmol H$^+$/g$^{-1}$) | $S_{BET}$ (m$^2$/g$^{-1}$) | V$_p$ (cm$^3$/g$^{-1}$) |
|---------------|----------------------------------|---------------------------|------------------------|
| Cc            | 0.05                             | 1011                      | 0.533                  |
| Cc–SO$_3$H    | 0.4                              | 975                       | 0.513                  |
| Cc–PhSO$_3$H  | 0.6                              | 793                       | 0.408                  |
| Cs            | 0.4                              | 523                       | 0.315                  |
| Cs–SO$_3$H    | 3.7                              | 34                        | 0.028                  |
| Cs–PhSO$_3$H  | 2.2                              | 10                        | 0.011                  |

Cc commercial carbon NORIT (GAC 1240 PLUS), Cs carbon synthesized from dextrose

$^a$BET surface area

$^b$Pore volume
and to corroborate the presence of the characteristic bands that allow the identification of sulfonic, carboxylic and polycyclic aromatic carbons. Figure 2 shows the FTIR spectra for Cs, Cs–SO₃H and Cs–PhSO₃H catalysts. All of them show a band around 1590 cm⁻¹, which can be attributed to the stretching of the C=C bonds formed as a consequence of the carbonization process (Zong et al. 2007; Toufiq Reza et al. 2016). Besides, a broad band located at 3440 cm⁻¹ is found, which is associated with the –OH stretching mode. This band, which is more intense in the spectra of samples containing –SO₃H groups, could originate from carboxylic and sulfonic groups. All the studied samples also present a band located at ca. 710 cm⁻¹, associated with the stretching mode of the C=O bond of the carboxylic group.

It should be noted that the band at about 3350 cm⁻¹, assigned to the N–H stretching absorption, does not appear in the Cs–PhSO₃H spectrum. The absence of this band is indicative of a successful diazotation step during the preparation of the functionalized carbon catalysts. On the other hand, a strong band at 1175 cm⁻¹, that is assigned to the
antisymmetric stretching mode of the S=O group of SO₂, can be observed in the Cs–SO₃H and Cs–PhSO₃H spectra. This band may be accompanied by a symmetrical stretching band at 1060 cm⁻¹. The spectra also show absorption bands located at 1035 cm⁻¹ and 1004 cm⁻¹; these bands are associated with stretching modes of the –SO₃ group. In the spectrum of the Cs–PhSO₃H sample, a band at 834 cm⁻¹, which is attributed to S–O bonds, can be observed. In addition, a band at 575 cm⁻¹ appears, assigned to the C–S stretching mode. These bands confirm that the –HSO₃ groups were successfully introduced onto the carbonaceous materials (Gonçalves et al. 2016; Figueiredo et al. 1999; Nersasian and Johson 1965).

Figure 3 shows a thermogravimetric analysis of Cs, Cs–SO₃H and Cs–PhSO₃H catalysts. In this figure, the mass loss (wt%) as a function of temperature (TGA curve) and the first derivative of the TGA curve (DrTGA curve) for each sample is shown. The DrTGA curves indicate at which temperature a mass loss process begins and ends.

The three catalysts present a first mass loss process, representing less than 15%, which starts at low temperature (<323 K) and ends at approximately 403 K. This process can be attributed to the loss of water. The base carbon, Cs (profile a), also exhibits a continuous mass loss between 474 and 1023 K, attributed to the loss of functional groups in the form of gaseous molecules such as CO, CO₂. These compounds could be generated from the decomposition of several types of oxygen functional groups such as: carboxylic groups, anhydrides, phenols, carbonyl, quinones, lactones, polycyclic aromatic compounds, etc. (Figueiredo et al. 1999; Figueiredo and Pereira 2010). The total mass loss associated with these processes is approximately 40%.

The Cs–SO₃H catalyst (profile b) shows three mass loss processes in addition to the water loss: one in the 473–533 K range, another from 533 to 700 K and the last one from 700 to 1023 K. The total mass loss for these processes is 56%. When comparing Cs and Cs–SO₃H catalysts, it is worth noting the higher mass loss observed at temperatures lower than 673 K for the latter. This result could be associated with the loss of functional groups that are not present in the Cs. It has been reported that part of the sulfonic groups decompose in this temperature range (Gonçalves et al. 2014; Zhao et al. 2010a).

The Cs–PhSO₃H catalyst (profile c) shows two mass loss processes, in addition to the water loss: the first one from 546 to 716 K and the other one from 716 to 1023 K. The total mass loss associated with these processes is 44%. The higher thermal stability of this catalyst with respect to the Cs–SO₃H catalyst is remarkable, a fact that could be associated with the preparation procedure. Nevertheless, it can be stated that all the catalysts are thermally stable in the temperature range in which they are used in the reaction.

Catalytic results

The catalysts (Cc and Cs) were studied in glycerol etherification using benzyl alcohol, at 393 K reaction temperature. The results obtained (conversions below 3%) demonstrated that the base carbons do not have sufficient acidity and therefore are inactive for this reaction. This behavior can be explained by taking into account the fact that the etherification reactions require the presence of an acidic active site to take place.

Influence of the Gly:BA molar ratio and of the catalyst percentage

The effect of the initial concentration of the reactants was evaluated at 393 K using three Gly:BA molar ratios (1:1, 3:1 and 5:1) and Cc–SO₃H as catalyst. As can be seen in Table 2, the conversion of BA does not significantly change between the 1:1 and 3:1 molar ratios; however, a marked effect on the product distribution is observed. The selectivity for the desired products (ME and DE) was 62% when working under conditions of excess glycerol (3:1), while for the 1:1 molar ratio, the main product obtained was the undesired BE. Therefore, by increasing the Gly:BA initial mass ratio, the BE selectivity decreased (Jaworski et al. 2015).

In Table 2, it can also be seen that, if the Gly concentration continues to increase (5:1 molar ratio), the selectivity to ME + DE is even more favored. However the conversion obtained reaches only an 8% value after 360 min of reaction.
The catalyst deactivation could be a result of strong adsorption of glycerol on its surface. On the other hand, the results presented in Table 3 show that the increase in the percentage of catalyst, based on the mass of glycerol, reflects a higher BA conversion (5, 23, and 30% conversion for 5, 10, and 15 wt% of catalyst, respectively). Nevertheless, the selectivity for the desired products decreases for the highest catalyst concentration.

Previous to this work, a lower dosage of catalyst was tested (2.5 wt%), but we did not obtain significant benzyl alcohol conversion. Taking into account the results obtained in these exploratory tests, the experimental conditions selected to continue with catalytic studies are: temperature 393 K, Gly:BA molar ratio 3:1 and a catalyst loading of 10 wt%.

Comparison of the different carbonaceous systems

Figure 4a depicts the conversion and selectivity values obtained for etherification of Gly with BA, using a 3:1 molar ratio and 10% of Cc–sulfonated catalysts. The results show that changes in the carbon functionalization method, Cc–SO$_3$H vs. Cc–PhSO$_3$H, produce a considerable increase in the conversion (23 vs. 66%, respectively) after 360 min of reaction. Although the difference in the acidity of both catalytic systems (0.4 and 0.6 mmol H$^+$ g$^{-1}$, respectively) is not quite significant, it is evident that the acid sites of the Cc–PhSO$_3$H catalyst are more efficient than those generated from the treatment with concentrated sulfuric acid (Cc–SO$_3$H). That is, the H$^+$ of –PhSO$_3$H group are more available for the protonation of BA, which will then lose a water molecule to become a carbocation, which in turn will be attacked by either Gly or another BA molecule.

The distribution of products also shows differences between the catalysts. The selectivity for ME + DE increased from 62 to 90% and the selectivity for BE diminished from 37 to 10% for Cc–SO$_3$H and Cc–PhSO$_3$H, respectively.

The formation of ME and DE agrees with the results published in the literature, where it has been reported that etherification of glycerol is preferentially in the primary hydroxyl groups of glycerol. ME formation is favored during the etherification, due to the reaction equilibrium reached in the presence of the water produced. The presence of water decreases the amounts of DE and TE formed, through their own hydrolysis (Gonçalves et al. 2014; Cannilla 2014). Therefore, the high monoether selectivity observed, along with the appreciable amount of BE found in the present study, could be explained taking into account that the water produced was not removed from the reaction medium.

The catalytic performance of Cc–functionalized carbon was compared to carbon synthetized in our laboratory.
(Cs) in the glycerol etherification with BA. The results for the Cs–SO$_3$H and Cs–PhSO$_3$H catalysts are presented in Fig. 4b. As can be seen, an almost complete conversion of BA (97 and 95% for Cs–SO$_3$H and Cs–PhSO$_3$H, respectively) is achieved after 360 min of reaction. The remarkable improvement in the catalytic performance can be attributed to successful incorporation of sulfonic groups, as well as other oxygenated surface groups, in particular carboxylic acids, which cause an increase of hydrophilicity of the solid surface, favoring the adsorption of Gly and BA. Zhao et al. (2010b) suggest that sulphonic acid groups are responsible for the improvement in the conversion of glycerol to its ethers. The presence of these groups does not limit the accessibility of reagents and products inside and outside the active sites; thus, high catalytic performance is reached despite the small surface area of the systems.

Both Cs–SO$_3$H and Cs–PhSO$_3$H catalysts were selective towards ME + DE, giving 87% and 93% selectivity, respectively (at the final time of reaction). For the Cs–SO$_3$H system, 11% selectivity was also observed for the self-condensation product (BE), whereas for Cs–PhSO$_3$H catalyst, BE only reached 5% selectivity. Thus, it can be concluded that the –PhSO$_3$H group functionalized system resulted in the most active and selective of the prepared catalysts.

Further, the results obtained in this work using Cs–SO$_3$H and Cs–PhSO$_3$H are similar and/or better than those reported in the literature. Gu et al. (2008) obtained with the C–SO$_3$H system only 12% yield. Gonzalez-Arellano et al. (2015) concluded that Zeolite ZSM-5 could be used as a good catalyst to produce ME (60%) and DE (38%), with little BE formation (2%), for 80% of BA conversion after 8 h reaction. Da Silva et al. (2009) obtained 100% of conversion after 120 min of reaction with the systems Amberlyst-35 and Zeolite β, but the product distribution was not entirely satisfactory.

Reusability of the catalysts

The reuse reactions were carried out at 393 K using the 3:1 Gly:BA molar ratio and the functionalized carbons Cs–SO$_3$H and Cs–PhSO$_3$H. It is important to note that no reactivation treatment was performed on the catalyst after each reaction cycle. The catalyst was subject only to a few washing and drying stages.

For both systems, Cs–SO$_3$H and Cs–PhSO$_3$H, a similar behavior is observed (Fig. 5) after each reuse. The final conversion decreased between 20 and 30% for each reaction cycle. According to the results of our studies, the vulnerability of active sites can be assigned to different deactivation factors, such as leaching or blocking of acidic sites because the adsorbed substrates cannot be removed by washing the catalyst. These results are in agreement with other publications (Gonçalves et al. 2016; Fraile et al. 2012). These catalysts were employed in different experiments published in the literature and the results confirm that leaching is one of motive of loss catalytic activity (Suwannakarn et al. 2008). Nevertheless the selectivity to ME + DE was kept between the second and third cycle for the best catalyst (Cs–PhSO$_3$H).

Conclusions

The catalytic behavior of Cs–SO$_3$H was studied in the etherification of Gly with BA. The results obtained allowed us to optimize the reaction parameters. A molar ratio Gly:BA of 3:1 and a catalyst percentage of 10% were chosen. Changing the carbon functionalization method (–PhSO$_3$H for –SO$_3$H groups) produced a considerable increase in glycerol conversion (from 23 to 66%) and in selectivity for ME + DE (from 62 to 90%). The Cs–SO$_3$H and Cs–PhSO$_3$H catalysts were obtained from synthesized carbon. These catalytic systems were highly active (97 and 95%, respectively) and selective. The main reaction product was ME (79 and 87%, respectively). The possibility of Cs–SO$_3$H and Cs–PhSO$_3$H catalyst reuse was investigated in three successive reactions. It was observed that the final conversion decreased between 20 and 30% for each cycle, but the selectivity for ME + DE was kept between the second and third cycle for Cs–PhSO$_3$H.

Acknowledgement

This work was supported by the Universidad Nacional de La Plata (Projects X633 and X700), CONICET (PIP Nº
0276) and ANPCyT (PICT 2014-0679) from Argentina. The authors are grateful to Pablo Fettis, CINDECA technician, for BET analysis.

**References**

Alonso DM, Wettstainea SG, Dumesic JA (2012) Bimetallic catalysts for upgrading of biomass to fuels and chemicals. Chem Soc Rev 41:8075–8098

Anitha M, Kamarudin SK, Kofli NT (2016) The potential of glycerol as a value-added commodity. Chem Eng J 295:119–130

Boehm HP (1994) Some aspects of the surface chemistry of carbon blacks and other carbons. Carbon 32(5):759–769

Bohon MD, Metzger BA, Linak WP, King CJ, Roberts WL (2011) Glycerol etherification with TBA: high yield to valuable products: an overview. Sustain Chem Process 2:1

Brunault J, Gu Y, Azzouzi A, Pouilloux Y, Jérôme F, Barrault J (2008) Heterogeneously catalyzed etherification of glycerol: new pathways for transformation of glycerol to more valuable chemicals. Green Chem 10:164–167

Han S, Kim M, Hyeron T (2003) Direct fabrication of mesoporous carbons using in situ polymerized silica gel networks as a template. Carbon 41:1525–1532

Janaun J, Ellis N (2010) Glycerol etherification by tert-butanol catalyzed by sulfonated carbon catalyst. J Appl Sci 10(21):2633–2637

Jaworski MA, Vega S, Síri GI, Casella ML, Romero Salvador A, Santos López A (2015) Glycerol etherification with benzyl alcohol over sulfated zirconia catalysts. Appl Catal: Gen 505:36–43

Len C, Luque R (2014) Continuous flow transformations of glycerol to valuable products: an overview. Sustain Chem Process 2:1

Liu R, Xiqing W, Zhao X, Fen P (2008) Sulfonated ordered mesoporous carbon for catalytic pre-treatment of biodiesel. Carbon 46:1664–1669

Liu X-Y, Huang M, Ma H-L, Zhang Z-Q, Gao J-M, Zhu Y-L, Han X-J, Guo X-Y (2010) Preparation of a carbon-based solid acid catalyst by sulfonating activated carbon in a chemical reduction process. Molecules 15:7188–7196

Liu T, Li Z, Li W, Shi C, Wang Y (2013) Preparation and characterization of biomass carbon-based solid acid catalyst for the esterification of oleic acid with methanol. Bioresour Technol 133:618–621

Martínez Fierro MC (2012) Preparación y caracterización de carbón activo a partir de lignina para su aplicación en procesos de descontaminación de aguas. Universidad Autónoma de Madrid, Madrid

Mo X, López DE, Suwannakarn K, Liu Y, Lotero E, Goodwin JG Jr, Lu C (2008) Activation and deactivation characteristics of sulfonated carbon catalysts. J Catal 254:332–338

Nersasan A, Johnson PR (1965) Infrared spectra of alkanesulfonic acids, chlorosulfonated polyethylene and their derivatives. J Appl Polym Sci 9:1653–1668

Pagliaro M, Ciriminnna R, Kimura H, Rossi M, Della Pina C (2007) From glycerol to value-added products. Angew Chem Int Ed 46:4343–4440

Rodriguez-Reinoso F (1998) The role of carbon materials in heterogeneous catalysis. Carbon 36:159–175

Sánchez JA, Hernández DL, Moreno JA, Mondragón F, Fernández JJ (2011) Alternative carbon based acid catalyst for selective esterification of glycerol to acetylglycerols. Appl Catal A 405:55–60

Suwannakarn K, Lotero E, Goodwin JG Jr, Lu Ch (2008) Stability of sulfated zirconia and the nature of the catalytically active species in the transesterification of triglycerides. J Catal 255:279–286

Tao M-L, Guan H-Y, Wang X-H, Liu Y-Ch, Louh R-F (2015) Fabrication of sulfonated carbon catalyst from biomass waste and its use for glycerol esterification. Fuel Process Technol 138:355–360

Toda M, Takagaki A, Okamura M, Kondo JN, Hayashi S, Domen K, Hara M (2005) Biodiesel made with sugar catalyst. Nature 438:178–179

Toufiq Reza M, Nover J, Wirth B, Coronella CJ (2016) Hydrothermal carbonization of glucose in saline solution: sequestration of nutrients on carbonaceous materials. AIMS Energy 4(1):173–189

Z hang W, Yang B, Yi C, Lei Z, Xu J (2010a) Etherification of glycerol with isobutylene to produce oxygenate additive using sulfonated peanut shell catalyst. Ind Eng Chem Res 49:12399–12404

Zhang Y, Wang H, Zhao Y, Shen J (2010b) Preparation of a novel sulfonated carbon catalyst for the etherification of isopentene with methanol to produce tertamyl methyl ether. Catal Commun 11:824–828

Zong M, Duan Z, Lou W, Smith TJ, Wu H (2007) Preparation of a sugar catalyst and its use for highly efficient production of biodiesel. Green Chem 9:434–437

**Publisher’s Note** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.