



Article

Study of Ni/Al-Fe Catalyst Stability in the Aqueous Phase Hydrogenolysis of Glycerol

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Abstract: The present work studied the stability and reusability of Ni/Al-Fe catalyst in the aqueous phase hydrogenolysis of glycerol without external hydrogen addition. The catalyst based on 28 molar % of Ni with 3/1 molar ratio of Al/Fe was prepared through co-precipitation. This catalyst presented the best performance in our last study which compares several Ni/Al-Fe catalysts with different molar ratios of Al/Fe. To see the influence of the pressurized water on the physicochemical characteristics of Ni/Al-Fe catalyst, a test of up to 9 h has been carried out. Fresh and used catalysts were characterized by various techniques: X-ray Diffraction (XRD), N2-physisorption, field emission scanning electron microscopy (FESEM) and STEM. Glycerol conversion and carbon yield to gases and liquids did not vary significantly when compared at 3 h and 9 h. Furthermore, the morphology of the catalyst remains stable after continuous recycling under severe hydrothermal conditions. The nickel rich phase of the catalyst, which was determined by XRD and scanning transmission electron microscopy (STEM) techniques, showed a stable size after 9 h under reaction.

Keywords: glycerol; in situ hydrogen; fixed bed; stability

1. Introduction

Nowadays, the production of biodiesel has been given considerable attention due to the growing concerns regarding environmental contamination and the depletion of existing fossil fuel reserves. Biodiesel is mainly produced by transesterification, where several triglycerides (vegetable oils, animal fats or waste oils, among others) react with an alcohol such as methanol or ethanol in the presence of a catalyst. From the biodiesel production, glycerol is obtained as a subproduct; about 0.1 ton is generated for every 1 ton of biodiesel. The surplus glycerol negatively affects the economics of the biodiesel industry [1–3]. Therefore, the conversion of glycerol into value-added products, such as H₂ and 1,2-propanediol, among others, is a method to improve the economics of biodiesel as it has been the case during the last decade. Many works have been published on aqueous phase hydrogenolysis (APH) of glycerol using different catalysts such as: noble metals (Pt, Ru, Pd or Rh), transition metals (Ni, Cu or Zr) and bi-metallics (Ni-Cu or Pt-Fe) supported on activated carbon, several oxides, or mixed oxides [4–18].

In recent years, Ni-based catalysts have attracted considerable attention due to their low price and high activity [19,20]. However, the major disadvantage of Ni-based catalysts is the low stability because of coke deposition and Ni sintering which cause catalyst deactivation and block the catalytic active sites and catalyst pores [15,19]. Hence, work continues on the development of multi-functional catalysts with coking inhibition, anti-sintering, significant activity, and stability [19]. Bastan et al. [21] found that mixed-oxide catalysts (Ni/Al $_x$ Mg $_y$) and the Ni/Mg catalyst were relatively more stable for 25 h of reaction time than the Ni/Al catalyst. Moreover, the mixed-oxide catalysts demonstrated best glycerol

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conversion and selectivity towards H₂ production in comparison to the Ni/Al and Ni/Mg catalysts in the following order: $Ni/Al_2Mg_1 > Ni/Al_1Mg_1 > Ni/Al_1Mg_2 > Ni/Al > Ni/Mg$. Therefore, the results revealed that the mixed oxide supports had a good stability and activity in converting biomass into hydrogen through aqueous phase reforming of glycerol [21]. In addition, Fan et al. [22] found that Mg-Al oxide supported Ni catalyst (Ni/Mg₂Al(O)), indicated higher activity at low reaction temperature and had a much better thermal stability (for 8 h of reaction duration at 700 °C) than the Ni/MgO and Ni/Al₂O₃ catalysts, respectively, for synthetic natural gas from syngas. In our previous study [18], we showed that mixed-oxide catalysts (Ni/Al_xFe_y) showed a better catalyst performance than the Ni/Al and Ni/Fe catalysts, with the Ni/Al₃Fe₁ catalyst being the best for hydrogenolysis of glycerol. Consequently, this work studied the stability and reusability of Ni/Al-Fe catalyst, in particular the Ni/Al₃Fe₁ catalyst, In addition, the effect of pressurized water on the physicochemical characteristics of the catalyst was asamined when the production of the control of the (XRD) n Niophysidosption of FESTAM, the descenning path principal on all through the residual for the acid nature of the reaction medium. Collected liquid products had a pH of around 4 as reported by 2. Results and Discussion Morales-Marin et al. [20]. This pH can be a consequence of the dissolved CO2 and/or the presence of 29 Lible axygenated compounds generated in the APH reaction of glycerol [20,24]. Arandia et al. [25] reported a significant metal leaching when the feedstock included acetic acid. Under these conditions, no effect is wine at the stability and veuse bility of Ni/Ala Fire catalysts for any appearance. Involves and veuse bility of Ni/Ala Fire catalysts for any appearance in volves and veuse bility of Ni/Ala Fire catalysts for any appearance in volves and veuse bility of Ni/Ala Fire catalysts for any appearance in volves and veuse bility of Ni/Ala Fire catalysts for any appearance in volves and veuse bility of Ni/Ala Fire catalysts for any appearance in volves and veuse bility of Ni/Ala Fire catalysts for any appearance in volves and veuse bility of Ni/Ala Fire catalysts for any appearance in volves and veuse bility of Ni/Ala Fire catalysts for any appearance in volves and veuse bility of Ni/Ala Fire catalysts for any appearance in volves and veuse bility of Ni/Ala Fire catalysts for any appearance in volves and veuse bility of Ni/Ala Fire catalysts for any appearance in volves and veuse of slive selivation by the development of the slive selection was studied at \$227 a Se and \$24 absolute bars use the life slive solvers line. deignized waterara feet for Johandeun ne Jeaurnes ive 4% ne faturation of John each (Interfatetal areaction) time) Figure 1 shows the glycarol conversion and carbon yield to gases and liquids. The carbon yield to gas or sliquide is calculated as the percentage of the crub on feeling to relicate states of the crub on feeling to relicate the converted to see, caliguide in Section 2.2, this is explained in depth Clearly Ni/Ala Fet catalystamoin to in 2001 and the catalystamoin to income caliguide. activityes veneuring sixe rups. Garbon vielent pases and liquid hand aluceral conversion this continue significantly when compared at 3 h and 9 h. Thus, the behavior of the catalyst was quite stable during a period of the rithod dition this satalyst shows more selectivity for living arroducts therefore an arroducts The walves of the servered nonversion and continuous teld to divide and see sea are between 2.15 w 5.3% c 237/c - 2.9 w 2.9 d 17 % = 1 9 % a respectively ver then stield ted invident 1 till 6 m and 9 h is in experimental extent because of the exercipating of interference of the control of the slightly high value increase after prolonged contact time.

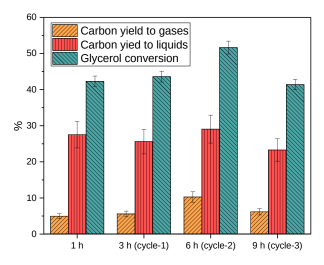
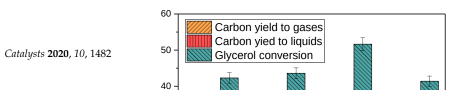


Figure 1. Clycerol conversion and carbon yield to gases and liquids.

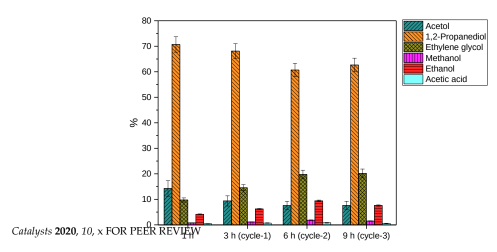
Consequently, Figure 2 shows the carbon selectivity to liquids for the different experiments. It is observed that the main liquid products are 1,2-propanediol, a liquid products are 1,2-propanediol, a liquid product are 1,2-propanediol product are 1,2-propaned

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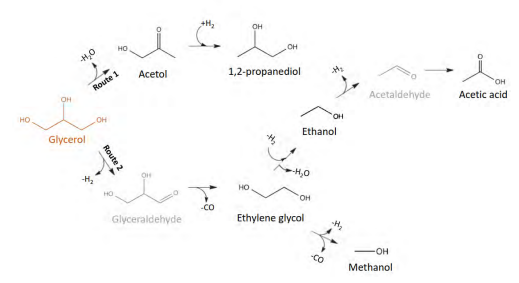


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d. In ad**i**ion, a sli Meanwhile the amount of ethylene col incre increase in the amount of ethanol and methanol was observed v h increas reaction ie, while 1,2-propanediol and acetic well kno that acid practically remained constant routes to obtain the liquid re two⊤n rmati products during the APH of glycer eme ıte 1 is cetol by the dehydration of glycerol and 1,2-propanediol prod ent h etol. This is the main route by su enatio in hydrogenolysis of glycerol. Rou the p tion c erald by the dehydrogenation of glycerol and then its consequent de Then, ethanol and methanol onyla fo can be generated by dehydrati carbonylation of ethylene glycol, respectively. Ethanol can also produce action action [42] 3]. According to the results obtained, it is possible that route 2 was favoured with an increase of reaction time and could occur after 6 h of reaction, corroborated with the results of glycerol conversion and carbon yield to gases and liquids.

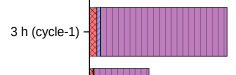


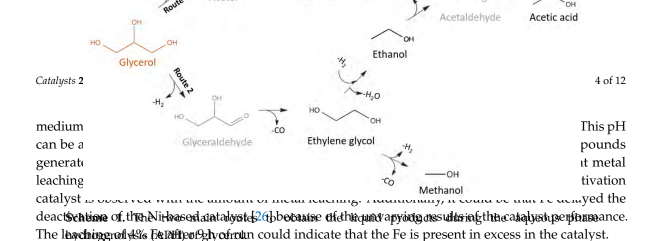
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Scheme 1. The two main ranters to obtain the liquid products during the advenus phase during p

In addition, the liquid products were submitted to ICP-OES analysis to determine if metal had been leached after the APH reaction of glycerol takes place. Trace amounts of Ni, Al and Fe were detected in the (diqueid) product (Figure 3). The % presented on Figure 3 represents the accumulated amount of leached metal. This is the percentage for each metal of total leached amount with respect to the initial amount in the catalyst. It is observed that Al in very low amounts is the only metal leached when H_2O is fed into the reactor. Fe is the metal that leached the most and then Ni. The explanation for this could be that the leaching took place during the APH reaction because of the acid nature of the reaction





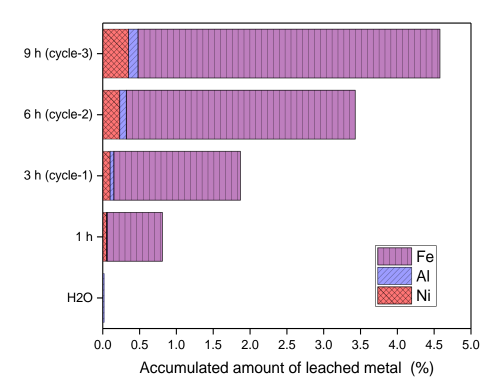


Figure 3. Metal leached after APH of glycerol takes place.

It is volk known about pretal deaching, metal capticly sainterings the adsurbious of preparities. Anthon for matine, each are sensidered the reasons for the catalyst deactivation for well in Metal particles, with the following section of the properties of the intering for the interior of the

2.2. Chisobserved that gluserol conversion and carbon yield to gases at 6 h show a slightly high value. Moreover, higher carbon selectivity to ethylene glycol is observed at 6 h than at lower time-on-stream. These two facts could be explained considering the obtained results: Fe is leached in higher proportion than Ni, the leached Fe could provide more active centres with Ni and these centres, rich in Ni, could favour the route 2. We do not have characterization results to demonstrate this, so it is just a hypothesis. Ni has the capability to break the C-C bond effectively towards ethylene glycol production [13].

2.2. Catalyst Characterization

The XRD patterns of fresh and used samples are presented in Figure 4. The NiO (bunsenite, JCPDS 00-001-1239), γ -Al₂O₃ (JCPDS 00-029-0063), NiAl₂O₄ (JCPDS 00-010-0339), Fe₂O₃ (JCPDS 00-001-1053), Fe₃O₄ (JCPDS 00-001-1111) and FeAl₂O₄ (JCPDS 01-086-2320) phases are observed in the calcined sample. Contrariwise, there are not differences between the XRD patterns of the reduced sample with

means that the water used to stabilize the system (at temperature and pressure of APH of glycerol) is the main factor for the boehmite formation. It is obtained by the reaction of Al₂O₃ with H₂O. In addition, the boehmite crystallite size was between 11 to 17 nm (Table 1).

Catalysts 2020, 10, 1482 metal, a high level of performance is obtained after 9 h of reaction time that could be due to excess of Fe in the catalyst as was cited above. A detailed analysis of Figure 4 does not show significant differences in the XRD patterns for the used catalyst for 1, 3 and 9 h. Because the tampsed promple towcopt thin the presence posting to propose promple of the catalyst for 1, 3 and 9 h. Because the tampsed promple for complete the proposition of the tampsed promple of the catalyst and for the proposition of the catalyst and the first and distinguished the catalyst and the first and the f

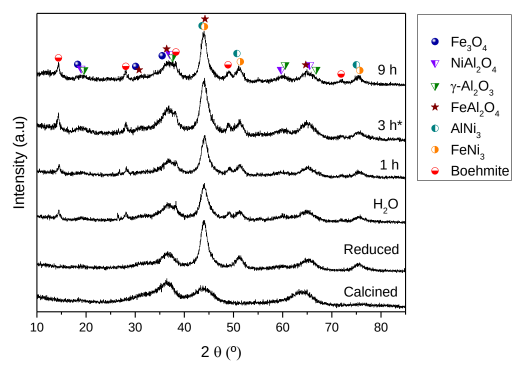


Figure 4. X-Ray Diffraction (XRD) patterns of fresh and used samples. * Raso et al. [18].

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Table 1 Textural properties FeNi: crystallite size and scanning transmission electron microscopy Table 1 shows the textural properties of fresh and used samples. The Brunauer, Emmett and (STEM) results of the samples. Boehmite crystallite size that Barrett-Joyner-Halenda (BFI) methods have been employed. It is observed that after reduction, the pore diagneter increased by around 59% applithe specific area declined in the samples.

Sample	(m^2/g)	(cm ³ /g)	(nm)	(nm)	(nm)	(nm)	
Calcined ⁶	222	0.211	3.93	-	-	-	_
Reduced	168	0.262	6.25	5	8.2	-	
H_2O	207	0.192	5.40	5	8.4	14	
1 h	210	0.189	4.75	6	8.5	11	
$3 h^6$	215	0.151	3.94	5	8.4	15	
9 h	210	0.153	4.18	6	8.2	17	

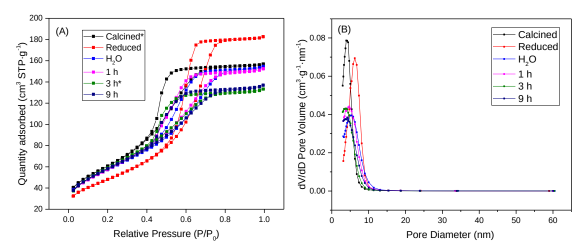
¹ The Brunauer, Emmett and Teller (BET) method. ² BJH adsorption method. ³ FeNi₃ crystallite size calculated from Scherrer equation. ⁴ Mean particle size of nickel-rich phase. ⁵ Boehmite crystallite size calculated from Scherrer equation. ⁶ Raso et al. [18].

The boehmite is observed before the APH reaction takes place (Figure 4: H_2O sample). This means that the water used to stabilize the system (at temperature and pressure of APH of glycerol) is the main factor for the boehmite formation. It is obtained by the reaction of Al_2O_3 with H_2O . In addition, the boehmite crystallite size was between 11 to 17 nm (Table 1).

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show, significant differences in the XRD patterns for the used catalyst for 1, 3 and 9 h. Because the samples present low crystallinity, it is not possible to propose which is the crystalline phase of the Fe that is despited. Regimpant of the check disconstituted and the present ALO. Amorphouse barranet Three the specific Three brains and the check disconstituted and the check disconstituted and the contract of the check disconstituted and the contract of the check disconstitution of h. Figure 5 is have a check disconstituted and the check disconstitution of th



Higure 5. (A) No adsorption description isotherms and (B) Barrett Joyner Halanda (BJH) adsorption pure size distribution of fresh and used samples.**Resortal [[18]].

Table 1. shows the textural, properties of thresh and used is amples in The Brunauer Emmett and Teller (BET) and Barrett-Jeyner-Halenda BH), methods have been employed. It is observed that after reduction, the pore diameter increased by around 59% and the specific area declined up to 24%, caused by the metallic not the nigration of the specific area of spent samples increased and there are almost no differences between them (H₂O, 1 h, 3 h and 9 h). This behavior could be due to the boshmite leaching from the catalyst surface. It is well known that γ-alumina could be hydrated to a poshmite or spibbsite phase under hydrothermal conditions, because hydroxides are thermodynamically more stables has under hydrothermal conditions, because hydroxides are thermodynamically more stables has posed and the reaction to perfect the stability of the catalyst [36]. Conversely, the pose diameter and volume decrease with the reaction to perfect the nation? In and the thirst more stables in page and the notion to the pose diameter and volume decrease with the reaction to perfect the nation? In and the thirst more stables in page and the nation of the nation o

Figure 6 howesther FEM image and particle action distribution of the reduction sicked sicker particle size of the complete is measured by image processing software. Around one hundred particles were measured per sample. It was observed that the mean diameter values for the reduction distribution of the particles were measured per sample. It was observed that the mean diameter values for the reduction distribution of the reduction distribution of the reduction o

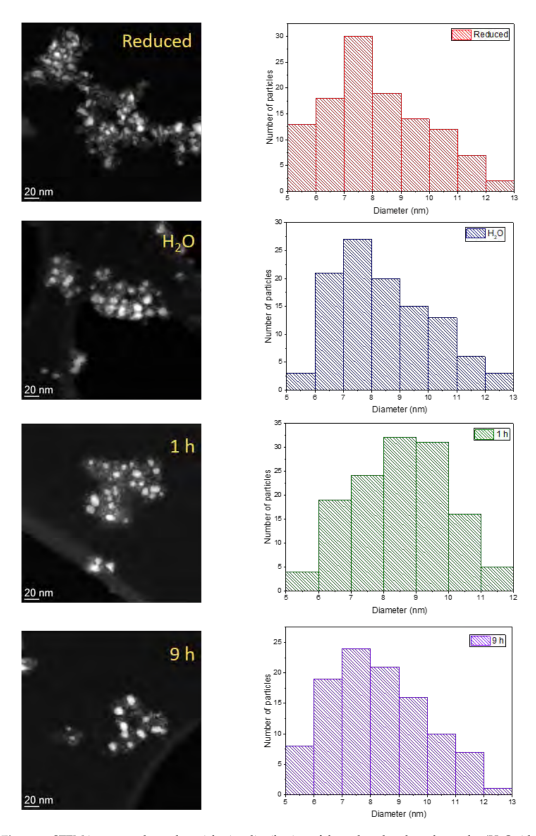


Figure 6. STEM images and metal particle size distribution of the reduced and used samples (H_2O , 1 h Figure 6. STEM images and metal particle size distribution of the reduced and used samples (H_2O , 1 h and 9 h.

Figure 7 shows the field emission scanning electron microscopy (FESEM) images of the reduced and used samples. It is observed that the morphology of Ni/Al₃Fe₁ catalyst changes after its use. There is almost no difference between the used samples with differences in treatment from just feeding water into the reactor until 9 h of reaction. The Ni/Al₃Fe₁ catalyst presented considerable morphology stability despite the hydrothermal conditions that occur during the reaction [18].

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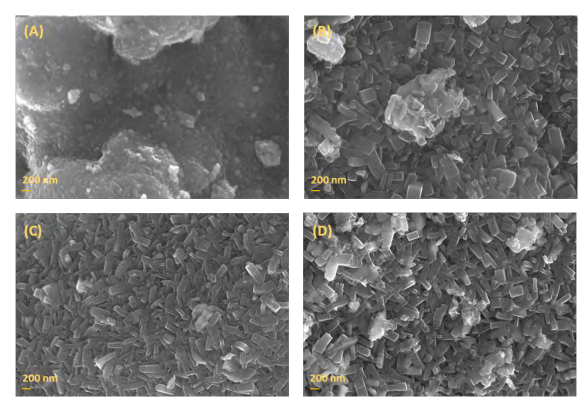


Figure 7: FESEM images of the reduced (A) and used samples: H2O(B), 1 h (O) and 9 h (D).

Single atom alloy catalysts were employed by Zhang et al. [31] in 1,2-propanediol production by glycerol hydrogenolysis. The PtCu single atom alloy catalysts howedlescellant catalytic performance. Single atom alloy catalysts are a promising area to be investigated due to the beneficial effect of metal dispersion and the synergistic effect when combining two active metals [32]. This is a relevant field that should be explored for Ni/Al-Fe catalysts in order to develop more active, selective and stable catalysts for 1,2-propanediol by glycerol hydrogenolysis.

3. Matterials and Methods

3:1: Eatalyst Preparation and Characterization

The Ni/Al-Fe catalyst with 28 molar % of Ni and 3/1 of molar ratio of Al/Fe was prepared by the co-precipitation method as reported earlier and the catalyst was named Ni/Al-Fe₄ [18]: It was calcined at 500 °C during 3 h: The catalyst was sieved to a mesh size of 160–315 µm: It was characterized by various methods such as N2-physisorption, X-ray diffraction (XRD), field emission scanning electron microscopy (FEEM), catalyst transmissions electron chief or the catalyst was prepared by the catalyst was prepared by the continued as the catalyst was sieved to a mesh size of 160–315 µm: It was characterized by various methods such as N2-physisorption, X-ray diffraction (XRD), field emission scanning electron microscopy (FEEM), catalyst was prepared by the catalyst was prepared by the continued at 500 °C during 3 h: The catalyst was prepared by the continued at 500 °C during 3 h: The catalyst was prepared by the continued at 500 °C during 3 h: The catalyst was prepared by the continued at 500 °C during 3 h: The catalyst was sieved to a mesh size of 160–315 µm: It was characterized by various methods such as N2-physisorption, X-ray diffraction (XRD), field emission scanning electron microscopy (FEEM), catalyst was prepared by the catalyst was prepar

3.2. Catalyst Tests 3.2. Catalyst Tests

The study of the Ni/Al₃Fe₁ catalyst stability was carried out in a continuous pressurized fixed bed The study of the Ni/Al₃Fe₁ catalyst stability was carried out in a continuous pressurized fixed reactor at 1 h of reaction and upon 3 successive runs at 227 °C and 34 absolute bar. The experimental bed reactor at 1 h of reaction and upon 3 successive runs at 227 °C and 34 absolute bar. The experimental system was developed and designed by PID (Process Integral Development Eng. & Tech, Madrid, Spain). It consists mainly of a stainless-steel tubular reactor (inner diameter of 9 mm) heated up by means an electric furnace and a micrometric valve, which regulates the pressure system. The fixed bed is constituted of a mixture of catalyst (2 g) and inert sand (5 g) with the same mesh size (160–315 µm) and is put inside the tubular reactor between quartz wool supports. The feeding was

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system was developed and designed by PID (Process Integral Development Eng. & Tech, Madrid, Spain). It consists mainly of a stainless-steel tubular reactor (inner diameter of 9 mm) heated up by means an electric furnace and a micrometric valve, which regulates the pressure system. The fixed bed is constituted of a mixture of catalyst (2 g) and inert sand (5 g) with the same mesh size (160–315 μm) and is put inside the tubular reactor between quartz wool supports. The feeding was an aqueous solution of 10 wt.% glycerol (purity: ≥99.5%, Sigma-Aldrich, St. Louis, MO, USA) in deionized water during the reaction of aqueous phase hydrogenolysis of glycerol, that was fed into the reactor by using a high-performance liquid chromatography (HPLC) pump (Gilson 307 HPLC Piston Pump, Champaign, IL, USA). The total flow was 1 mL/min with the mass of catalyst/glycerol mass flow rate ratio (W/m) of 20 g_{catalyst}.min/g_{glycerol}.

Prior to the start the experiments, the calcined catalyst was reduced in situ at 500 °C during 1 h using a H₂ stream of 100 cm³ (STP)/min. The reduction temperature was stablished from the catalyst characterization results by H2-TPR [18]. Then, before starting the reaction, deionized water with a flow rate of 1 mL/min was fed to stabilize temperature and pressure to reaction conditions for about 2 h. Hence, in this work the influence of the water on the physicochemical characteristics of Ni/Al₃Fe₁ catalyst was studied.

The reaction products (gas and liquid) exiting from the reactor is depressurised by means of the micrometric valve and arrives at the condensation system. The condensation system consists of four condensers and the first one was employed to collect the water used to stabilize the system. The other condensers were used to gather the liquid products. The exit gas mixture (N2, H2, CH4, CO2, CO, C₂H₆ and C₃H₈) was analysed online by means of an Agilent 490 Micro-GC equipped with Thermal Conductivity Detectors (TCD) (Santa Clara, CA, USA). The liquid products (methanol (MeOH), ethanol (EtOH), acetol, acetic acid, 1,2-propanediol (1,2-PDO), ethylene glycol (EG) and non-reacted glycerol) were analysed offline with an Agilent 7820A GC equipped with a Flame Ionization Detector (FID) and a HP-FFAP Agilent 19091F-105 capillary column (Santa Clara, CA, USA). In addition, the liquid products were submitted to an inductively coupled plasma optical emission spectrometry (ICP-OES) analysis to study metal leaching, using a Thermo Elemental IRIS Intrepid Radial with an automatic injector (Markham, ONT, Canada). More details of the experimental rig can be found in our previous study [18].

The stability of the catalyst was explored in repeated experiments for 3 h in order to test its reuse. Reusability of the catalyst was tested using the reaction conditions above but with the previous and only catalyst reduction in the first cycle. 3 total cycles were analyzed for 3 h (9 h in total).

The catalytic performance was calculated according to expressions (1)–(3) below [18].

The global glycerol conversion was calculated as follows:

Glycerol conversion (%) =
$$\frac{n_{glycerol}^{in} - n_{glycerol}^{out}}{n_{glycerol}^{in}} \times 100$$
 (1)

where $n_{glycerol}^{in}$ and $n_{glycerol}^{out}$ are the moles of glycerol fed and the moles of unreacted glycerol in the exit liquid, respectively.

The carbon yield to liquids and carbon yield to gases were defined as follows:

Carbon yield to liquids (%) =
$$\frac{n_{\text{MeOH}} + 2n_{\text{EtOH}} + 2n_{\text{Acetic acid}} + 3n_{\text{Acetol}} + 3n_{1,2-\text{PDO}} + 2n_{\text{EG}}}{3 \times n_{\text{glycerol}}^{\text{in}}} \times 100$$
 (2)

Carbon yield to liquids (%)
$$= \frac{n_{MeOH} + 2n_{EtOH} + 2n_{Acetic\ acid} + 3n_{Acetol} + 3n_{1,2-PDO} + 2n_{EG}}{3 \times n_{glycerol}^{in}} \times 100$$
 (2)
$$Carbon\ yield\ to\ gases\ (\%) = \frac{n_{CO} + n_{CO_2} + n_{CH_4} + 2n_{C_2H_6} + 3n_{c_3H_8}}{3 \times n_{glycerol}^{in}} \times 100$$
 (3)

where n_i are the moles of each i product, i being the liquid or gas products.

The carbon selectivity to liquid products was defined as the percentage ratio of carbon in a liquid product to the total carbon in all the analysed liquid products. Unreacted glycerol was not considered. Catalysts 2020, 10, 1482

There is not total coincidence between the glycerol conversion and the addition of the carbon yield to products (gas and liquid) due to errors in analysing and collecting the samples. An experiment with a value of carbon deficit below o% was considered a reliable test, as reported by other authors [7,18,23]. The carbon deficit was defined as follows:

Carbon deficit = Glycerol conversion – (carbon yield to gases + carbon yield to liquids)

4. Conclusions

In this study, APH of glycerol under Ni/Al $_3$ Fe $_1$ catalyst was performed during 9 h and the influence of the pressurized water on the physicochemical characteristics of the catalyst was analyzed. Structural stability and reusability of the Ni/Al $_3$ Fe $_1$ catalyst were verified under the reaction conditions. All samples present approximately the same FeNi $_3$ crystallite size from the reduced to 9 h samples. The morphology of the catalyst remains stable after successive reuses under severe hydrothermal conditions. The carbon yield to gases and liquids and glycerol conversion did not change significantly when compared at 3 h and 9 h. Although trace amounts of Ni, Al and Fe were detected in the liquid product, this did not cause deactivation of the catalyst under the operating conditions. In addition, the boehmite was formed by the reaction of Al $_2$ O $_3$ with H $_2$ O under the operation conditions, prior to the APH of glycerol. Meanwhile, the metals leaching occurred during the APH of glycerol.

Author Contributions: Conceptualization—L.G. and M.O.; Methodology—R.R. and J.R.; Software—R.R.; Validation—R.R.; Formal analysis—R.R. and J.R.; Investigation, R.R.; Writing—original draft preparation—R.R.; Writing—review and editing—R.R., L.G., J.R., M.O. and J.A.; Visualization—R.R.; Supervision—L.G. and J.A.; Project administration—L.G.; Funding acquisition—L.G. and J.A. All authors have read and agreed to the published version of the manuscript.

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