

Catalytic oxidative desulphurization of pyrolytic oils to fuels over different waste derived carbon-based catalysts

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Abstract

In this work, we reported the conversion through carbothermal process of two catalysts produced by pyrolyzing exhausted coffee and waste tires. We tailored the surface with anchored iron nanoparticles through a facile carbothermal route and tested them for catalytic oxidative desulphurization of high sulphur content oil derived from tires pyrolysis. We studied their activity in a biphasic system under different conditions reaching a desulphurization of up to 60 % by using an oil with a sulphur concentration of up to 7139 ppm. The extensive characterization proved the reliability of those materials as promising catalysts for upgrading of sulphur rich drop-in fuels.

Keywords: biochar; drop-in fuel; tires; pyrolysis; biphasic catalysis

1.Introduction

Increased accountability of European Union with regards to environmental issues has represented a formidable driving force for the development of sustainable processes[1]. As a consequence, tires landfilling was forbidden with the European Council Directive 1999/31/CE due to its dangerous impact on the environment and human health[2, 3]. Furthermore, sustainable and circular economy principles have growth leading to the exploration of alternatives end-life tires processes[4].

The simpler methods are their use as inert filler for constructions[5, 6] and as road asphalt[7] or for the direct production of raw materials such as rubber[8-10] and textiles[11]. Alternatively, thermochemical routes are cost-effective approaches for end-life tires treatment avoiding the complexity of alternative management routes [12] such their conversion through pyrolytic techniques[13, 14].

Oil recovered from pyrolysis of tires is a very attractive product stream due to their composition that is very close to an engine fuel[15]. Nonetheless, their sulphur content is generally very high avoiding their for real field applications and requires an upgrading process to met the fuel regulation requires. Among the all upgrading procedure, catalytic oxidative desulphurization[16] is very interesting for meeting the sulphur limits of common fuels [17] under the principles of green chemistry[18]. Plenty of different approached are described in literature based on transition metal-based catalysts (i.e. cobalt [19], tungsten [20] and manganese [21] or polyoxometalates [22]) using air [23, 24] or peroxides as oxidant agents [25-27], in homogenous or biphasic systems [28-31]. Iron-based catalysts are very promising due to their low cost price and to the ability to enhance oxidative power of oxygenated water [32] through Fenton mechanism [33]. This procedure achieved remarkably performances in the degradation of benzothiophene [34] and desulphurization of diesel-like fuels [35] by using nanostructured catalysts. Nonetheless, these systems were produced by using unfriendly procedures and lead to the merely deposition of the iron nanoparticles onto the support surface. Carbothermal route could represent a simple and facile alternative to produce iron nanoparticles anchored onto carbon support through thermochemical conversion of iron salts to metal iron nanoparticles[36, 37]. In this study, we exploited the synergist effect of the use of waste-derived carbon support for the upgrading of waste-derived drop-in fuel. The integration of waste valorization and alternative fuels production could represent a first step for a deep re-think of sustainable platforms. Here, we reported the development of two nanostructured iron-based catalysts produced though carbothermal conversion of exhausted coffee residues biochar (CC) and carbon recovered from the very same tire pyrolysis (TC). Nanostructured catalysts were used to upgraded oil from pyrolysis of tires though a biphasic catalytic desulphurization in a watery biphasic system. The catalytic activity and catalyst structures were widely analyzed and compared achieving good results in desulphurization of high sulphur content oils. We also evaluated the effect of microstructured (CC) and nanostructured (TC) support in the catalytic performances.

2. Materials and methods

2.1 Materials

Fe (NO₃)₃ nonahydrate EtOH (>98%), HNO₃ (65%) and H₂O₂ (30% v/v) were purchased by Sigma Aldrich. Coffee powder was collected from Bar Katia (Turin, Italy) supplied by Vergnano (Arabica mixture). A Pirelli branched tire was used for this study.

61 2.2 Methods

62 2.2.1 Pyrolysis

63 Coffee was collected and dried at 105°C prior the pyrolytic process. Afterwards, 100 g were
64 pyrolyzed using a vertical furnace and a quartz reactor, heating rate of 15 °C/min and kept at 800°C
65 for 30 min accordingly with previous studies[38-41].

66 Tire was chopped in small pieces and pyrolyzed using a tubular furnace (Carbolite TZF 12/65/550) in
67 nitrogen by using a heating rate of 15 °C/min and kept at 550°C for 30 min. Solid and liquid fractions
68 were collected.

69 A thermogravimetric analysis (TGA) of the oil was performed from 25 to 900 °C using a TGA
70 Mettler Toledo model 1600 in two different atmospheres (Argon and air) with a gas flux of 50 ml/min
71 with a heating ramp of 10 °C/min to describe a possible oil composition.

72 2.2.3 Preparation and characterization of catalysts

73 Recovered carbon from tire and biochar produced by pyrolyzing exhausted coffee were used as
74 starting materials for carbothermal process. 50 g of each of them were suspended in 250 mL of
75 deionized water together with $\text{Fe}(\text{NO}_3)_3$ (weight of Fe/ weight carbon precursor of 1:10). The solution
76 was stirred for 10 min and dried in a ventilated oven at 105°C overnight. The dried materials were
77 used without any additional purifications. Carbothermal process was run by using a tubular furnace
78 (Carbolite TZF 12/65/550) in nitrogen atmosphere with a heating rate of 15 °C/min and kept at 800°C
79 for 30 min. The solids recovered were analyzed prior and after carbothermal processes by using
80 several techniques.

81 Morphology was studied by using a field emission scanning electron microscope (FESEM, Zeis
82 SupraTM40, Oberkochen, Germany). The microscope was equipped with an energy dispersive X-ray
83 detector (EDX, Oxford Inca Energy 450, Oberkochen, Germany) that was used to explore the
84 elemental composition of catalysts.

85 Raman spectra were collected by using Renishaw® Ramanscope InVia (H43662 model,
86 Gloucestershire, UK). Signals were fitted according to methodology proposed by Tagliaferro et
87 al.[42]

88 Surface of catalysts was investigated by using X-ray photoelectron spectroscopy (XPS). XPS
89 spectrometer was a PHI 5000 Versaprobe Physical Electronics, Chanhassen, MN, USA) scanning X-
90 ray photoelectron spectrometer (monochromatic Al K-alpha X-ray source with 1486.6 eV energy, 15
91 kV voltage, and 1 mA anode current) to investigate surface chemical composition.

Specific surface area of the samples was measured by means of N₂ sorption at -196°C on a micrometrics Tristar II instrument (Micromeritics Instrument Corporation, USA). Brunauer–Emmett–Teller (BET) model was applied.

2.2.3 Catalysis

Catalytic tests were run in a biphasic systems as sketched in figure 1.

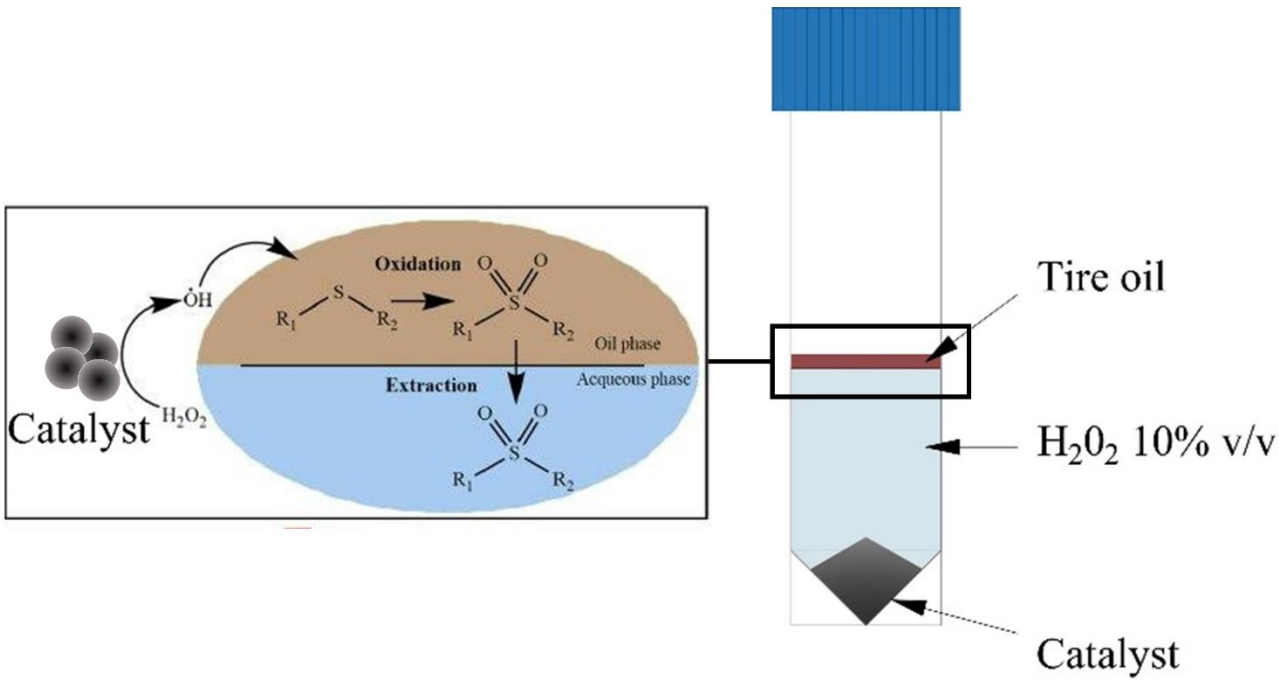


Figure 1. Scheme of catalytic set-up

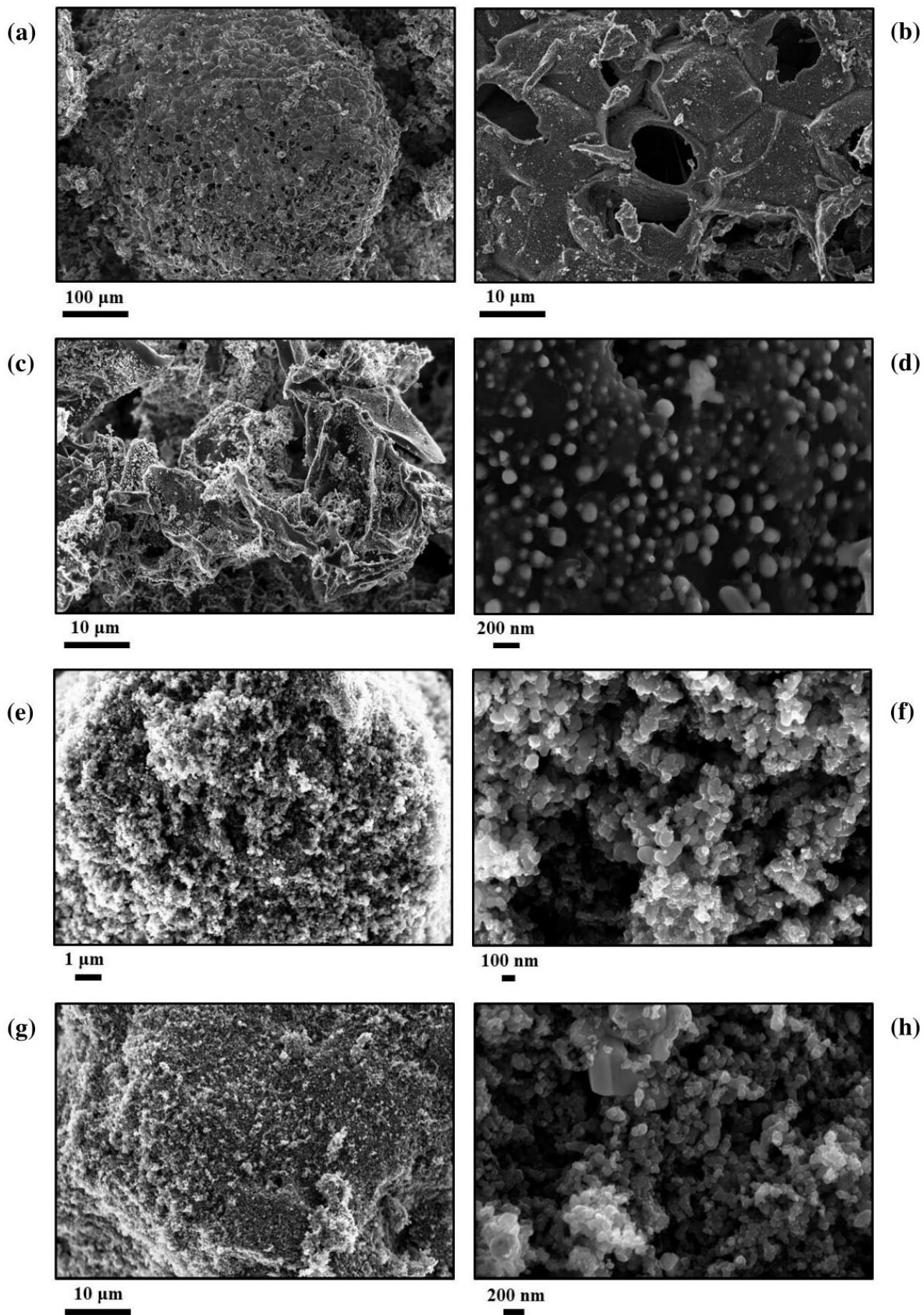
Catalysts were suspended in a 15 mL of a watery phase of H₂O₂ (10 v/v%) and 2 g of oil recovered from pyrolysis of tires were added. The sealed vials were put in an oil bath at different temperatures (60, 80 and 100 °C), for different time (2, 4 and 6 h) and by using several catalysts loading. After the reaction, they were cooled down at room temperature and 10 mL of deionized water were added. Oily phases were collected and purified by suspended particles by using a centrifuge (1000 rpm for 10 min). Recovered catalysts were collected, washed with acetone, dried at 105 °C overnight and further analyzed.

Sulphur concentration was determined by ICP (ICP-OES Perkin Elmer Optima 2000 DV) after acidic digestion in microwave oven (ETHOSUP Milestone) by using H₂O₂ 30 v/v% in HNO₃ 65 wt.% (ratio 1:9) with a power of 800W for 35 min.

109 **3. Results and discussion**

110 **3.1 Characterization of the catalysts**

111 Solid recovered from pyrolytic conversion of both exhausted coffee powder and waste tires were used
112 as starting materials for carbothermal synthesis of catalysts[43, 44]. This facile approach has been
113 used to tailor the surface of carbon from pyrolysis of exhausted coffee and waste tires with metal
114 nanostructures as those shown in figure 2.



116 **Figure 2.** FESEM captions of a-b) biochar from exhausted coffee, c-d) CC prior catalysis, e-f)
117 carbonized tires and g-h) TC prior catalysis

118 The surface of the coffee biochar prior carbothermal process (figure 2 a-b) was structured in
119 micrometric sponge-like aggregates whose diameter is around 20 to 100 μm . Coffee biochar shows a
120 very low specific surface area of up to $0.14 \text{ m}^2/\text{g}$ mainly due to the presence of micrometric pores
121 formed during the release of volatile organic matters from the inner core of biomass particles.
122 Contrary, carbon recovered from pyrolysis of tires showed a structure closely related to carbon black
123 used for tires production[45]. This material was composed by of a complex agglomeration of carbon
124 nanoparticles (similar to carbon black structure) with a specific surface area of up $37.7 \text{ m}^2/\text{g}$.
125 Carbothermal synthesis of CC induces the formation on the carbonaceous surface of semispherical
126 iron particles with diameters ranging from 50 to 150 nm (figure 2 c-d). CC surface area increased of
127 up to $46.2 \text{ m}^2/\text{g}$ due to the presence of nanoparticles together with the activation of carbon matrix, as
128 described by Wang et al[46]. Similarly, TC displayed spherical iron nanoparticles anchored to carbon
129 together with submicrometric iron aggregate (figure 2 h). TC showed a specific surface area
130 increment of up to $53.9 \text{ m}^2 / \text{g}$ mainly ascribed in this case to the presence of iron species. The more
131 Comparing structures reported in figure 2d and figure 2 h, we observed that iron nanoparticles are
132 fully exposed on surface of CC due the bigger size of carbon support compared with growing iron
133 nanoparticles. Considering TC, carbon black particles were much smaller compared with iron
134 nanoparticles and were unable to provide an anchoring surface as in the case of CC partially covering
135 the nanostructures formed.

136 The elementary compositions of the CC, TC and their precursors were estimated by using EDX
137 analysis as shown in Table 1.

138 **Table 1.** Elementary composition of pyrolyzed waste coffee obtained through EDX analysis, I_D/I_G and surface area before and after deposition of Fe
 139 nanoparticles.

	Elemental analysis (wt.%)											I_D/I_G	Surface area (m ² /g)
	C	O	Mg	P	K	Ca	Fe	Si	S	Zn	Na		
Coffee biochar	86.9	8.9	0.6	0.5	1.8	1.3	Not detected	Not detected	Not detected	Not detected	Not detected	2.13	0.1
Tires char	91.9	4.8	Not detected	Not detected	Not detected	Not detected	Not detected	0.2	1.0	2.2	Not detected	2.04	37.7
TC	70.9	7.8	Not detected	Not detected	Not detected	1.5	19.8	0.5	1.3	1.5	Not detected	2.72 ^b	53.9
TC^a	72.8	4.4	Not detected	Not detected	Not detected	Not detected	21.1	0.2	0.5	Not detected	Not detected		52.1
CC	62.2	11.6	0.6	0.3	3.5	0.7	21.2	Not detected	Not detected	Not detected	Not detected	2.04 ^b	46.2
CC^a	60.5	15.8	0.7	1.1	1.2	0.7	19.5	Not detected	Not detected	Not detected	0.5		44.9

140 a) Catalysts after third catalytic cycle, b) I_D/I_G was unchanged before and after third catalytic cycle since the Raman spectra did not show any
 141 appreciable differences

Coffee biochar showed an amount of carbon as carbon recovered from tires together with an appreciable amount of oxygen that could be associated to both residual group on carbon surface and to inorganic species. Inorganic species in coffee derived biochar derived from the biological fixation routes that originates the feedstock[47]. Additionally, carbon recovered from tires displayed a sulphur content of up to 0.5 wt.% due to the not complete conversion of sulphur into organic or volatile organic matters as previously described by Undri et al[48]. Furthermore, TC precursors showed a negligible native iron content estimated in less than 0.1 wt.% by analysis its ash content. After, carbothermal process TC showed a totally disappearing of sulphur due to its oxidation with formation of SO_x . The final [Fe] was attested in both cases around to 20 wt.% (19.8 wt.% and 21.2 wt.% for TC and CC respectively).

A further investigation of carbon structure of CC, TC and their precursors was run by using Raman spectroscopy as reported in figure 3.

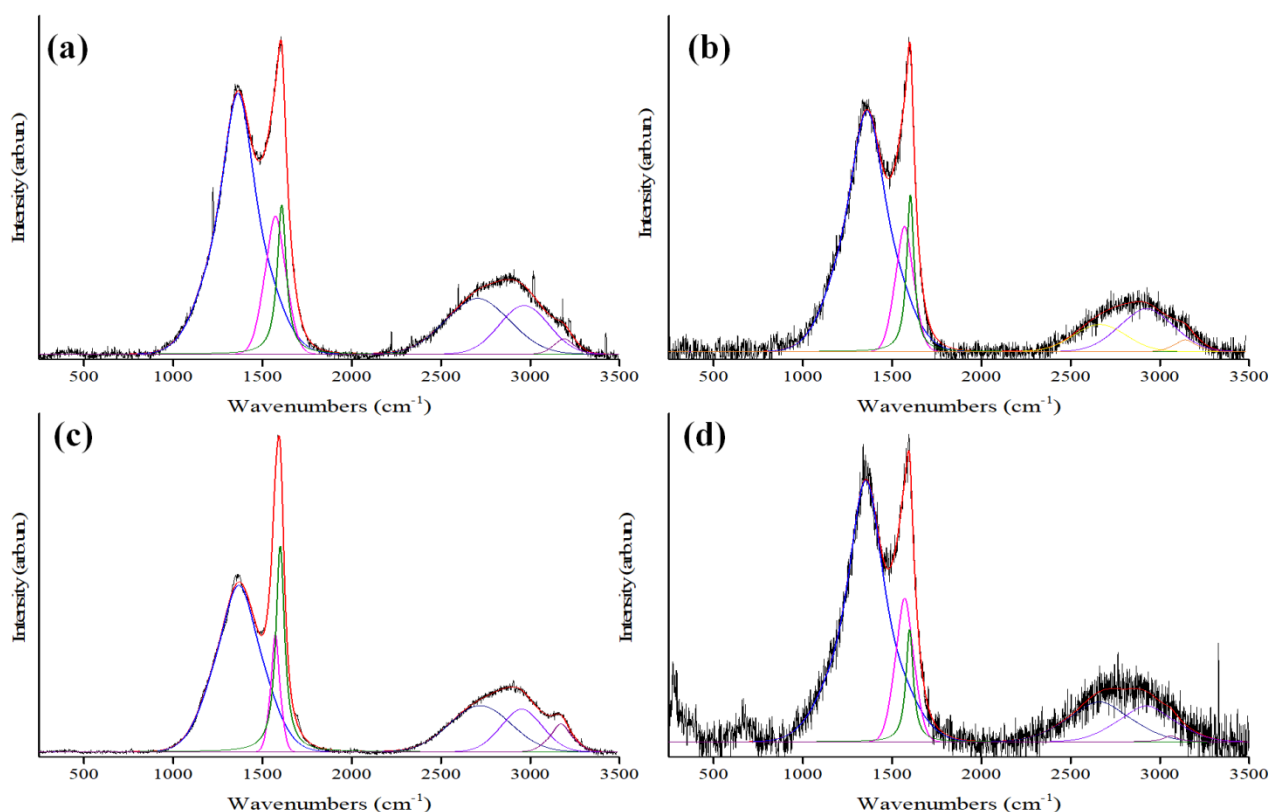


Figure 3. Raman spectra of a) biochar from exhausted coffee a coffee, b) CC prior catalysis, c) carbonized tires and d) TC prior catalysis. In black was reported the original Raman spectra and in red the fitted ones accordingly with the procedure proposed by Tagliaferro et al[42].

Raman spectroscopy is a powerful tool to estimate the carbon degree of graphitization [49]. Particularly, the area ratio of D and G peaks in the region from 1300 to 1700 cm^{-1} (I_D / I_G) is possible investigated the carbonaceous structure disorder. Both coffee derived biochar and carbon recovered from waste tires were highly disordered materials with a I_D/I_G of up to 2.13 and 2.04 respectively[50]. After carbothermal process, CC and TC showed a decrement of I_D/I_G down to 2.04 while TC showed an increment of up to 2.72. This different behavior could be ascribed to the intrinsically differences between the two precursors. CC was produced by using a material that retained a relevant amount of residual functional groups and sp^3 carbon content while carbon recovered from carbonization of tires was mainly composed by original carbon used to produce the neat tires. Considering CC, carbothermal process polished partially the carbon surface with a decrement of imperfections of carbonaceous structure while those are magnified for TC that was more ordered. This was also confirmed by the analysis of XPS spectra reported in the figure 4.

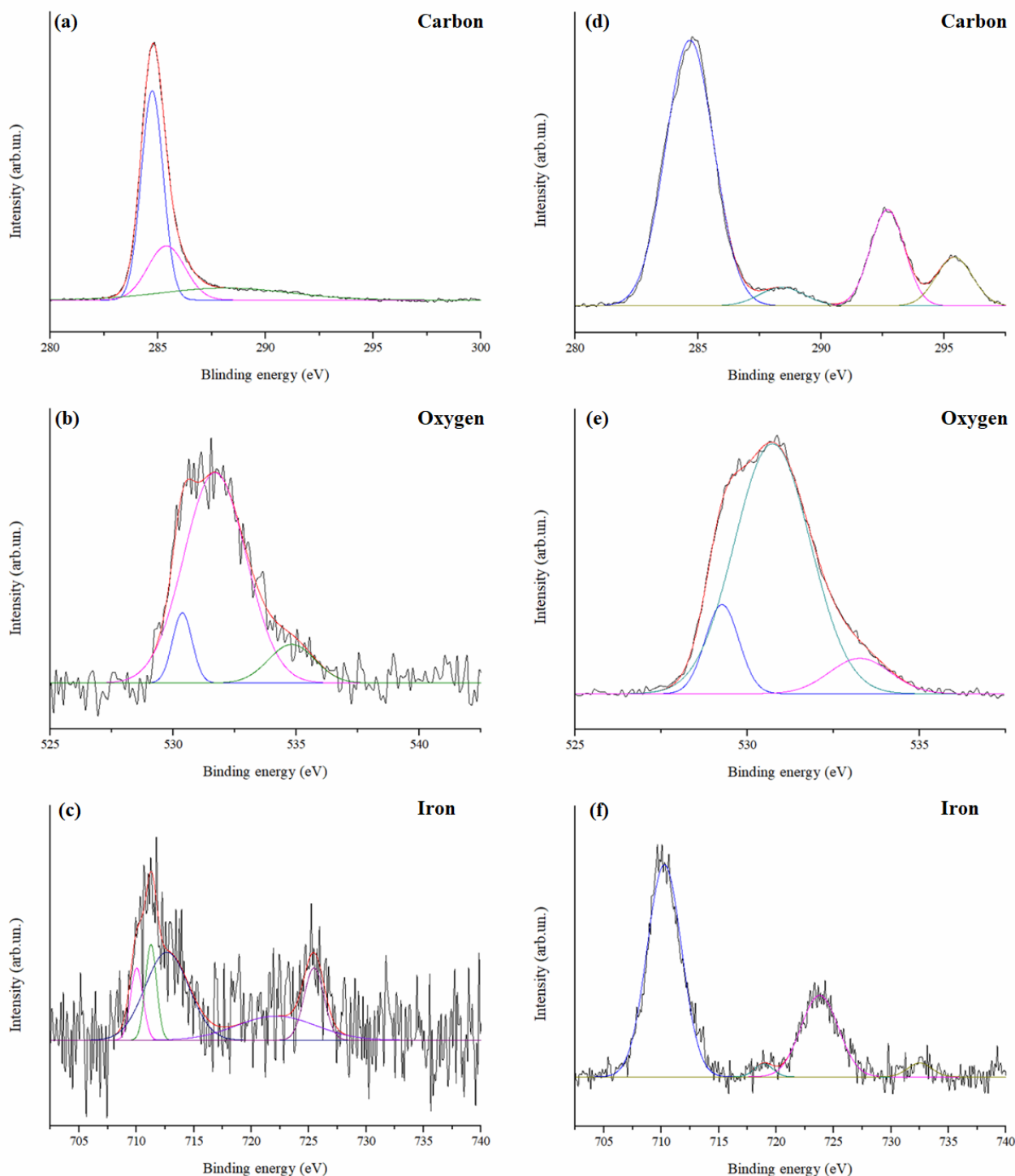


Figure 4. XPS spectra of CC (a-c) and TC (d-f). Carbon, oxygen and iron region are here displayed.

XPS spectra of CC (figure 4 a) showed a high amount of sp^2 carbon preponderance (peak 284 eV) that reach the 68.5% of the total amount while sp^3 represented only the 17.5 % (peak 284.9 eV). TC was composed by only sp^2 carbon that was highly functionalized by oxygen functionalities proving the disorder increment observed by Raman spectroscopy. Those oxygen functionalities displayed peaks at 529, 531 and 533 eV. Peak at 531 eV represented up to 79% for both CC and TC and it was

associated with the presence of oxygen linked to quinoid carbons. The peak at 529.3 eV with a relative area of around 12 % was instead attributed to an oxygen bound to an iron in iron oxide[51]. Furthermore, the presence of Fe-OH residue on the particle surface was proved by the peak at 533.3 eV that represent the 8.9% of the total area[52]. The small area underlying this last peak shows that these groups exist only on the surface of the Fe nanoparticle and render the metal oxide layer partially defective. The formation of this defects could be ascribed to the presence of FeOOH formed during air passivation of iron nanoparticles as reported by Ponder et al.[53] at room temperature.[54]. Accordingly to figure 4 c and f, the spectrum of iron was shown and the presence of metallic iron and iron oxide across the peaks at 710.5 and 712.2 eV was detected. Iron oxidation state was in agreement with the peak at 530.0 eV of the oxygen spectrum, typical of a metal oxide. From the conclusions extrapolated from the carbon characterization before and after the carbothermic process, it was possible to obtain a model of the iron nanoparticle supported on a carbonaceous matrix. As can be seen from figure 5, the hypothesized iron nanoparticle model was based on a metallic iron core whose external surface was made by a layer of Fe₂O₃ and FeOOH sites generated through passivation attached to the surface of the pyrolytic carbon through a very thin layer of iron carbide accordingly to Li et al[55]. CC showed a Fe(III)/Fe(0) ratio of 55.7% while TC showed a value of 53.3%.

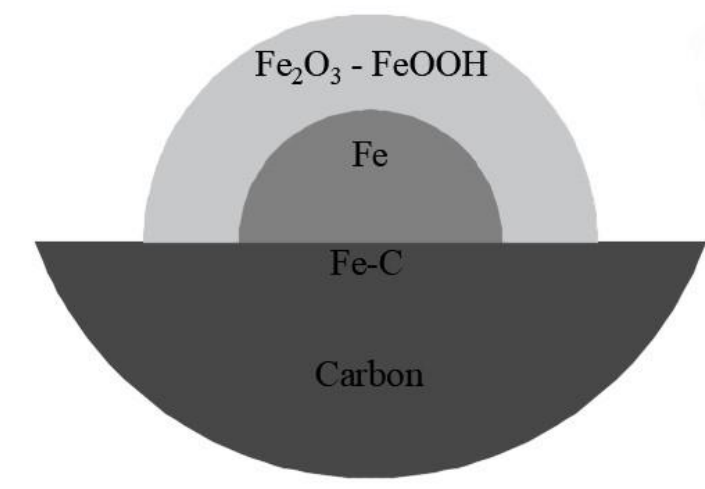


Figure 5. Hypothesized structure of iron particles anchored onto carbon support after carbothermal reduction and air passivation.

3.2 Preliminary considerations about oil recovered from pyrolysis of waste tires

Pyrolysis of waste tires was run at 550 °C accordingly with Ucar et al.[56] trying to maximizing the liquid yield. Nonetheless, the temperature adopted promoted the highest polycyclic sulphur aromatic compounds content as reported by Williams et al.[57]. In our study, oil recovered from the exhausted tires pyrolysis showed a concentration of up to 7139 ppm that greatly exceed the limit for use as drop-

in fuel. Nonetheless, the TGA profiles of oil shown in figure 6 confirmed a composition close to a conventional fuel, rich in C₆-C₁₀ hydrocarbons, accordingly with the study reported by Kök et al.[58].

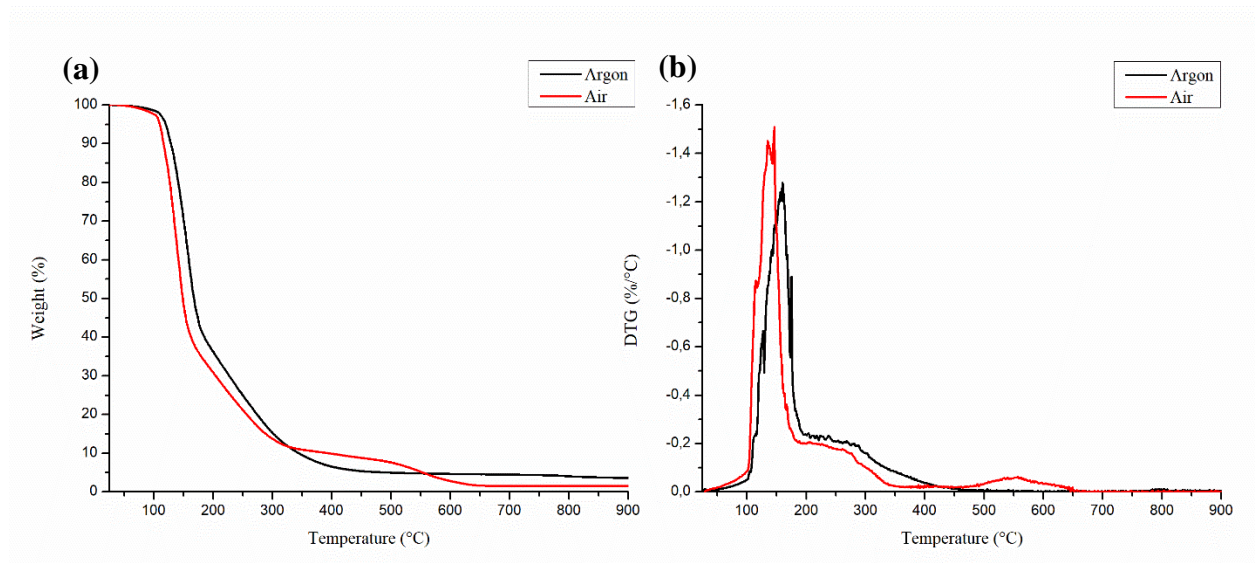


Figure 6. TGA and DTG of tire oil at Ar and air atmosphere from 25°C to 900°C.

Furthermore, the weigh losses between 500°C and 600°C could be attributed to the presence of polycyclic aromatic hydrocarbons which are more resistant to oxidation than other hydrocarbons.

3.3 Catalytic activity of CC and TC in oxidative desulphurization of oil recovered from pyrolysis of waste tires

Catalytic oxidative desulphurization of high sulphur content oil recovered from pyrolysis of tires was study by using CC and TC and varying several parameters as shown in table 2.

Table 2: Catalytic performances of CC and TC during oxidative desulphurization of high sulphur content oil recovered from pyrolysis of tires.

Entry	Catalyst	T (°C)	T (h)	Desulphurization ^(a) (%)	[S] (ppm)	Loading ^(b) (wt.%) / TON ^(c)
1	-	100	6	33.2	4769	-
2	CC	100	2	47.5	3747	5/219
3		100	4	52.3	3406	5/241
4		100	6	63.7	2592	5/293
5		60	6	36.4	4540	5/168
6		80	6	52.0	3425	5/240
7		100	6	49.0	3639	2/565

8	TC	100	6	40.8	4223	10/94
9		100 ^(d)	6	39.2	3747	5/181
10		100	2	57.5	3036	5/283
11		100	4	58.1	2989	5/286
12		100	6	56.1	3132	5/276
13		60	6	34.0	4709	5/167
14		80	6	36.1	4564	5/177
15		100	6	53.7	3305	2/660
16		100	6	52.2	3415	10/128
17		100 ^(d)	6	42.3	4116	5/169

(a) Desulphurization= $100 - (100 \times \text{final sulphur concentration} / \text{initial sulphur concentration})$, (b) loading= $100 \times (\text{weight of iron} / \text{weight of oil})$, ^(c)TON= mmol of removed sulphur/mmol of accessible iron sites ^(d)3rd cycle of desulphurization.

Comparing the removal efficiencies of the catalysts and the H₂O₂ (desulphurization of up to 33.2) under the same operating conditions (6h at 100 ° C with 5% g Fe / g oil) was evident the improvement induced by CC and TC that reached a desulphurization of up to 63.7 % and 53.7%. CC displayed the better performances compared with TC according to TON values. This was reasonably due to the effect of different carbon matrix. CC present iron nanoparticles fully exposed on the surface while TC was characterized by iron particles surrounded by nanometric carbon particles. Accordingly, TC activity could deplete by the hindrance of carbon matrix with lower iron nanoparticles accessibility.

By examining the entries 12, 13 and 14, an improvement in sulfur removal at high temperatures can be seen due to an evident kinetic boost. TC desulphurization rose from 36.1% at 80 ° C of up to 56.1% at 100 ° C. Similarly, temperature improved the CC desulphurization activity from 36.4% to 63.7% of. Temperature effect was scarcely influenced by the support but it was more related the activation energy together with phase transfer equilibria [59].

The catalyst loading is the other key parameter of the investigated. Catalyst loading of 2 % induced a reduction of desulphurization compared to the use of 5% for both CC and TC. Surprisingly, a catalyst loading of up to 10 wt.% lead to a decrement of desulphurization at 100°C after 6 h down to 40.8 % for CC while TC was unaffected. This was reasonably due to aggregation phenomena of micrometric particles of CC that did not occurred by using TC composed by nanometric carbon particles. Additionally, the original high sulphur content represented an obstacle to an efficient removal by using a biphasic approach with the extracted sulphone that rapidly reached the saturation of watery phase. Nonetheless, CC and TC displayed catalytic performances very close to systems

operating on tires derived oil[28, 60] even if none of previously published research based on biphasic systems treated oil with a sulphur content high such the one used in the present study.

Both CC and TC showed a reduction of activity after the third catalytic cycle down to 39.2% and 42.3% respectively. This was not related to iron leaching as proved by iron content retention as shown in table 2 but it was ascribed by a sized increment of iron particles as shown in figure 7.

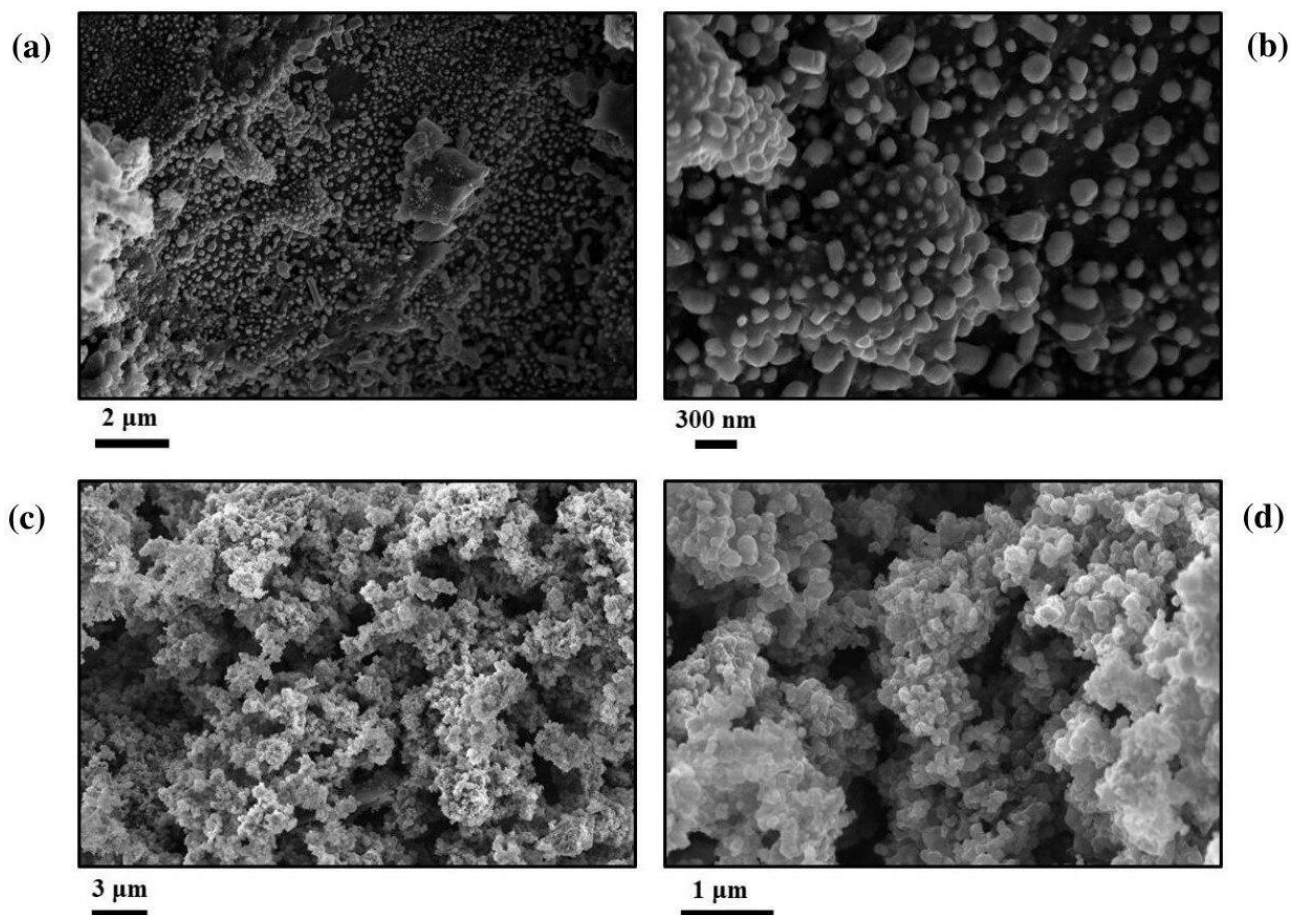


Figure 7. FESEM captions at different magnifications of CC (a-b) and TC (c-d) after three catalytic cycles.

CC showed an increment of particles average diameter of up to 100-300 nm while TC shows iron aggregated covered by spherical carbon nanoparticles.

However, the XPS spectra of recycled catalysts (figure 8) support the observed loss of reactivity due to the composition change on the catalyst surface.

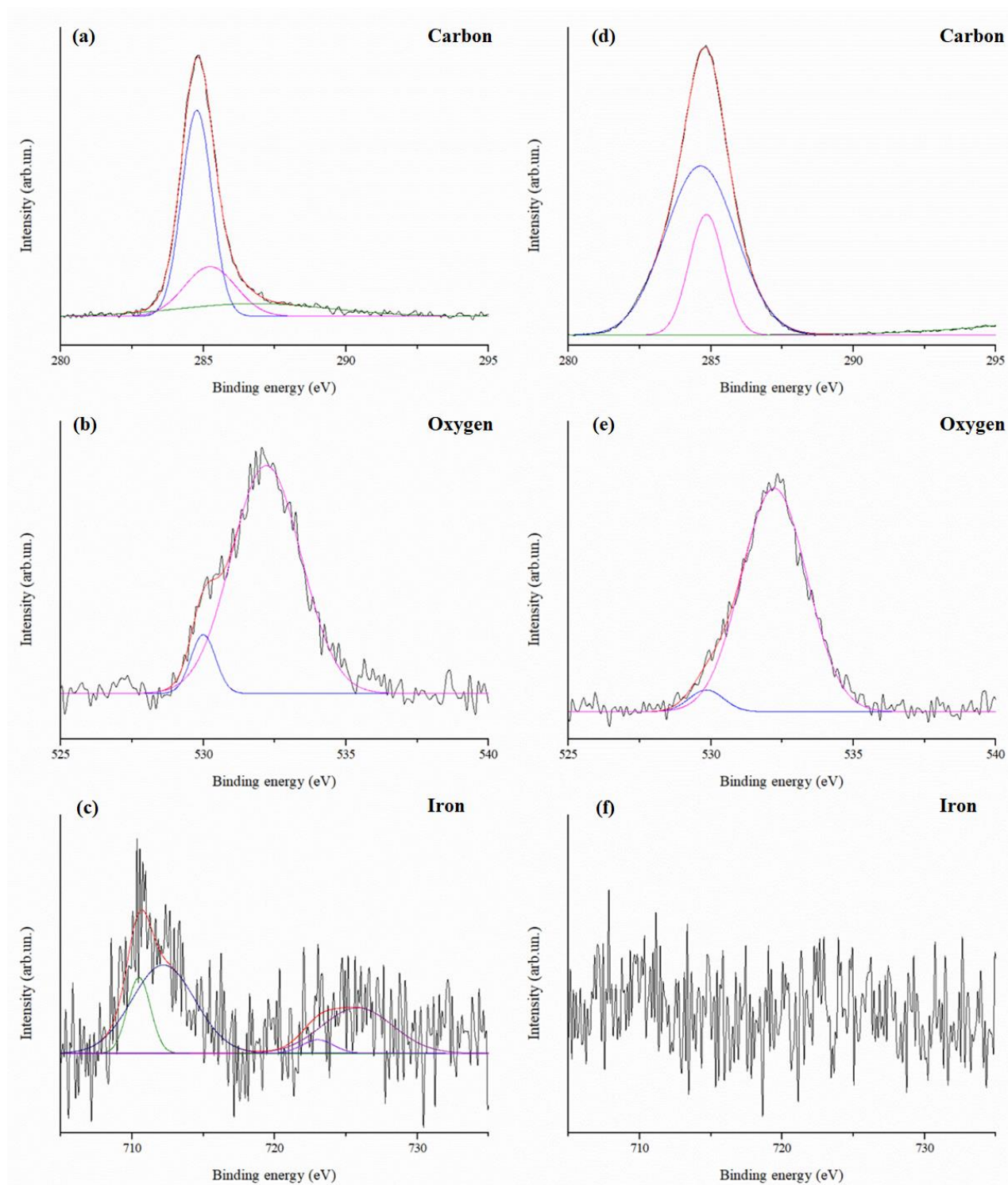


Figure 8. XPS spectra of CC (a-c) and TC (d-f) after the 3rd catalytic run. Carbon, oxygen and iron region are here displayed.

The loss of reactivity was also due to the increasing of the thickening of the crystal lattice of the iron oxide which reduces the tension on the same thus relaxing the orbitals. About each spectrum shown in figure 8, the iron signal of TC (figure 8 f) was characterized by a high noise level which did not

allow the detection of any peak. A fitting analysis of the iron spectrum of CC was performed although the large noise because of the micrometric size of the unpacked material from this type of catalyst. In particular, five peaks were distinct (the last two are replicas) while the three initial peaks are identifiable to a metallic iron, Fe-OH and iron oxide. About oxygen spectra, the position of the oxygen peaks has remained unchanged, so the catalyst continues to present oxides of iron and oxygen of a quinoidic nature. While the signal of carbon (figure 8 a), the spectrum of CC a showed the insertion of a new peak at 285.3 eV which is associated with the formation of a C-OH bond caused by the action of H₂O₂. The other two peaks at 284 and 288.1 eV of area, respectively, of 56.8 and 18.9% of the total, mark, as previously seen, the presence of C sp² and C of a quinoidic nature.

We supposed that the original structures of CC and TC underwent to different deactivation process as such reported in figure 9

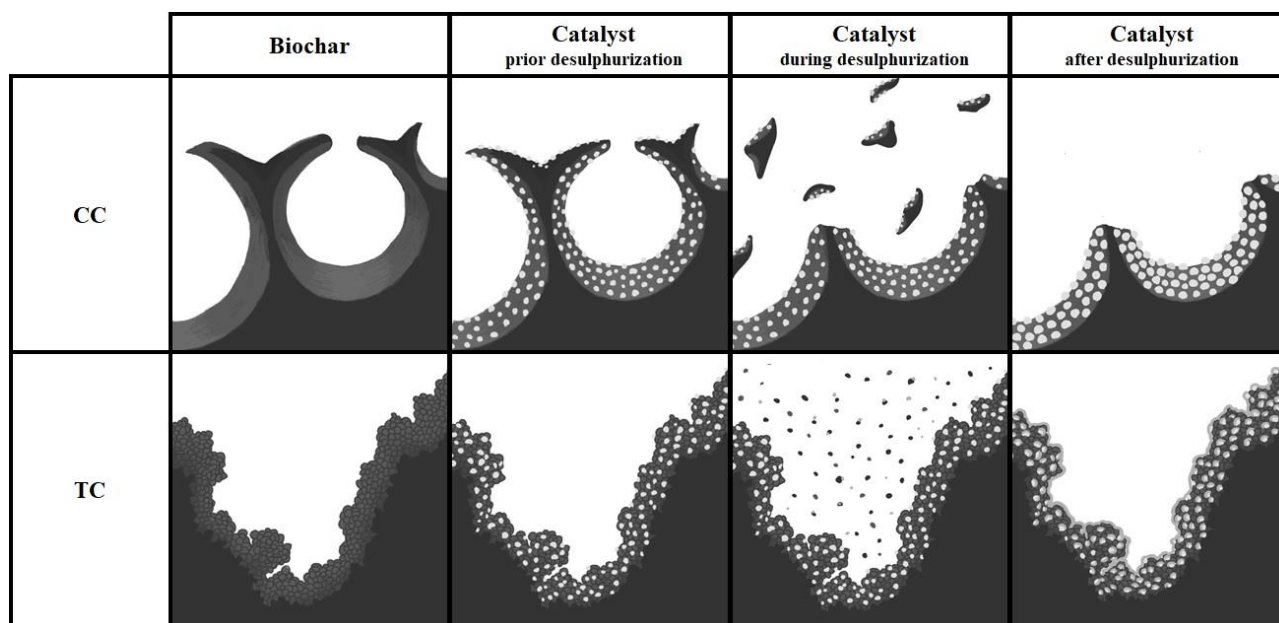


Figure 9. Evolution of the catalysts during the catalytic oxidative desulphurization of oil from tires pyrolysis.

According with FESEM analysis (figure 8 b), iron nanoparticles anchored onto CC increased their sized with a decrement of available catalytic sites on the surface. TC deactivation was mainly due to the covering of the iron nanoparticles by the spherical carbon particles of the support. This proved lower anchoring of iron nanostructures to TC than CC. The surface flatterring of CC was caused by the unpacking of micrometric carbon particles so modifying the surface morphology, as shown in figure 7 a.

4. Conclusions

In this work we reported the production and catalytic desulphurization of high sulphur content oil recovered from pyrolysis of tires by using biphasic catalytic oxidative desulphurization. Catalytic systems proposed were based on conversion of two waste streams through a facile carbothermal process capable to produce highly active heterogeneous catalysts characterized by nanostructured iron-based material. Air passivation after the carbothermal processes was a key step for achieving a hemispherical external layer of Fe_2O_3 and FeOOH on an iron-based core. Thanks to the special geometry of the catalytic system created, CC showed a desulphurization ability of up to over 60% by using high sulphur content oil at the best conditions (at 100°C with a residence time of 6 hours). The morphological characteristics of the pyrolyzed waste streams were fundamental to describe the behavior of the catalytic system during the desulfurization processes. TC was successful in maintaining constant the desulphurization rate at different conditions thanks to submicrometric structure which increases the phase transfer between the reactants in the biphasic system. While the CC shows a micrometric structure, covered by nanometric iron-based particles, which tends to influence more the kinetics of the catalytic system. Despite this, iron-based materials produced by carbothermic processes could be considered a great improvement in Fenton reactions for oxidative desulphurization thanks to easier separation through magnetic proprieties and absence of leaching into reaction solution.

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