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*Original Citation:*

*Availability:*

This version is available <http://hdl.handle.net/2318/1687811> since 2019-01-24T14:13:19Z

*Published version:*

DOI:10.1016/j.jenvman.2018.10.001

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(Article begins on next page)

1 **REMOVAL OF QUEBRACHO AND TARA TANNINS IN FUNGAL BIOREACTORS:**  
2 **PERFORMANCE AND BIOFILM STABILITY ANALYSIS**

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24

1 **Abstract**

2 Tannins are polyphenolic compounds produced by plants that are used in the vegetable tanning of  
3 leather at industrial scale. Quebracho tannin and Tara tannin are intensively used by the tanning  
4 industry and are two of the most recalcitrant compounds that can be found in tannery wastewaters.  
5 In this study two reactors fed with Quebracho tannin and Tara tannin, respectively, were inoculated  
6 with polyurethane foam cubes colonized with a fungal strain biofilm of *Aspergillus tubingensis*  
7 MUT 990. A stable biofilm was maintained in the reactor fed with Quebracho tannin during 180  
8 days of operation. Instead, biofilm got detached from the foam cubes during the start-up of the  
9 reactor fed with Tara tannin and a bacterial-based suspended culture was developed and preserved  
10 along the operational period (226 days). Soluble chemical oxygen demand removals up to 53% and  
11 90% and maximum elimination capacities of 9.1 g sCOD m<sup>-3</sup> h<sup>-1</sup> and 37.9 g sCOD m<sup>-3</sup> h<sup>-1</sup> of  
12 Quebracho and Tara tannins, respectively, were achieved in the reactors without the addition of co-  
13 substrates. Next generation sequencing analysis for bacteria and fungi showed that a fungal  
14 consortium was developed in the reactor fed with Quebracho tannin while fungi were outcompeted  
15 by bacteria in the reactor fed with Tara tannin. Furthermore, Quebracho and Tara tannins were  
16 successfully co-treated in a single reactor where both fungi and bacteria were preserved.

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25 **Keywords:** recalcitrant compounds; tannins removal; non-sterile conditions; fungal reactor;  
26 biomolecular analyses.

## 1   **1   Introduction**

2       Tannins are abundantly applied in leather tanning industry due to their prerogative of  
3 precipitating proteins (Lorenz et al., 2014). In fact, tannins bind to the collagen proteins of the  
4 animal skin making leather more durable and not putrescible. Tannins are water-soluble  
5 polyphenolic compounds produced by plants (Barbehenn and Peter Constabel, 2011) and could be  
6 subdivided into hydrolysable (gallotannins and ellagitannins), condensed (or proanthocyanidins)  
7 and complex tannins (Khanbabae and Ree, 2001). Tara tannin (TT) is obtained from the fruit pods  
8 of *Caesalpinia spp.*, which principal components are hydrolysable tannins based on a galloylated  
9 quinic acid structure (Sciences et al., 2007). Quebracho tannin (QT) is obtained from the wood of  
10 *Schinopsis spp.* and is composed mainly of condensed tannins that are more difficult to be  
11 biodegraded than those found in TT given that hydrolysable tannins are easier to be biodegraded  
12 (He et al., 2008). Conventional activated sludge systems (Li et al., 2009) and anaerobic digestion  
13 (Mannucci et al., 2010) are not effective to treat tannins due to their low biodegradability and high  
14 soluble chemical oxygen demand (sCOD) (Lofrano et al., 2013). In fact, high concentration of  
15 tannins can inhibit the biological treatment (Munz et al., 2009). Then, tannins are usually removed  
16 by means of chemical processes in tannery wastewater treatment plants (WWTPs).

17       Although the leather tanning industry is known to be of prime economic importance in many  
18 countries, the concern about the environmental impacts related to the production of leather and the  
19 release of various recalcitrant pollutants in tannery wastewaters (Lofrano et al., 2013; Romer et al.,  
20 2011) has been increasing in the last decades. Current biological wastewater treatments are  
21 engineered biological ecosystems based on bacteria, which are ineffective in the removal of several  
22 recalcitrant compounds such as tannins (Mannucci et al., 2010). Fungi play a role in the  
23 biodegradation of phenols, chlorinated phenolic compounds, chlorinated alkanes and alkenes,  
24 polycyclic aromatic hydrocarbons, petroleum hydrocarbons and other emerging contaminants  
25 (Harms et al., 2011). Although fungi represent a promising biological resource in environmental

1 biotechnology, they have rarely been applied in wastewater treatment due to a lack of knowledge  
2 regarding the optimal process conditions and due to their lack of stability under non-sterile  
3 conditions (Espinosa-Ortiz et al., 2016). Most research in wastewater treatment with fungi has been  
4 focused on the degradation of pharmaceuticals, dyes and pesticides at lab-scale, with important  
5 limitations and, above all, mostly under sterile conditions (Svobodová and Novotný, 2017). In fact,  
6 the main operational concerns are related to bacterial contamination and to the robustness of the  
7 bioprocess in the long-run since fungi are easily outcompeted by bacteria. Nowadays, operation of a  
8 fungal-based bioreactor able to maintain stable fungal growth and performance, under sterile and  
9 non-sterile conditions, is still challenging. Furthermore, the fungal degradation of most recalcitrant  
10 compounds frequently requires the supply of an external carbon source (co-substrate), since  
11 recalcitrant compounds might not be suitable substrates for fungal growth (Palli et al., 2016).  
12 Nevertheless, among recalcitrant compounds present in tannery wastewaters, tannins represent a  
13 potential carbon source for fungi, despite their antimicrobial properties (Mingshu et al., 2006).  
14 Alternative biological treatment processes able to effectively remove this fraction could lead to  
15 environmental and economical advantages (Giaccherini et al., 2017). Moreover, *Aspergillus* spp.  
16 and *Penicillium* spp. have been isolated in tannery wastewaters and reported as microorganisms  
17 capable to biodegrade tannins (Murugan et al., 2007). In particular, *Aspergillus* (section Nigri) can  
18 grow on tannic acid (TA) as carbon source and was found in tannery wastewaters with high QT  
19 concentration (León-Galván et al., 2010). Based on this findings, it can be hypothesised that fungi  
20 would be able to grow in the recalcitrant fraction of wastewater containing a high concentration of  
21 tannins, such as tannery wastewater. Most of the research reported in the literature about the  
22 biodegradation of natural tannins and polyphenolic compounds by fungi has been performed in petri  
23 dishes or flasks using tea by-products (Ni et al., 2015), tannin extracts (Belmares et al., 2009) or on  
24 TA (Van Diepeningen et al., 2004). To the authors' knowledge, no previous experiments have been  
25 reported with continuously fed bioreactors (under sterile or non-sterile conditions) for the removal  
26 of QT, TT or other natural tannins with fungi.

1           The aims of the present work were i) to demonstrate the proper performance ii) to evaluate the  
2 stability and iii) to assess the bacterial and fungal diversity of a novel fungal bioreactor  
3 configuration operated under non-sterile conditions to reach an effective removal of Tara and  
4 Quebracho tannins. TT was selected as a representative of hydrolysable tannins and QT as a  
5 representative of condensed tannins, among the natural tannins applied as tanning agents. The  
6 evolution of the system was evaluated by means of combined physical-chemical analyses and  
7 molecular techniques for bacteria and fungi. Almost no works in literature have analysed both  
8 communities and their relationship during the treatment of tannins in bioreactors.

## 9   **2   Materials and methods**

### 10   **2.1   Fungal strain, immobilization on supports and reagents**

11           Based on previous tests (Tigini et al., 2015) and literature research, *Aspergillus tubingensis*  
12 MUT 990, a black *Aspergillus* belonging to the section Nigri, was chosen as the inoculum for the  
13 present study. This fungal strain was originally isolated from commercial TT powder and is  
14 preserved at Mycotheca Universitatis Taurinensis (MUT) on malt extract agar (MEA) (agar 20 g,  
15 glucose 2 g, malt extract 2 g, peptone 0.2 g, water up to litre) at 4°C. The selected fungal strain was  
16 inoculated on 20 generic MEA plates (150 mm diameter) and incubated at 25°C in the dark for  
17 seven days. After incubation, the fungal colonies were cut to obtain a homogenate. For each cm<sup>2</sup> of  
18 mycelium, 1 mL autoclaved distilled water was added (Anastasi et al., 2012). Polyurethane foam  
19 (PUF) cubes were chosen as carriers (Spina et al., 2012). PUF carriers were cubes of about 2 cm per  
20 side (density 25 kg m<sup>-3</sup>, pores/volume ratio: 0.97 and specific surface 600 m<sup>2</sup> m<sup>-3</sup>), which were  
21 inserted in six 1L-Erlenmeyer flasks (35 cubes each) containing 500 mL of GLY medium (5 g L<sup>-1</sup>  
22 of glucose and 1.9 g L<sup>-1</sup> of yeast extract) and autoclaved at 121°C for 30 min. Afterwards, 10 mL of  
23 fungal homogenate was added to each flask. Flasks were shaken in the dark for seven days at 110  
24 rpm and 25°C. This procedure allowed the immobilisation of the selected fungal strain onto the  
25 PUF. The QT and TT commercially used for tanning were kindly provided by Chimont

1 International Spa, Montopoli (Italy). All other reagents used in the present study were of analytical  
2 grade (Sigma-Aldrich).

### 3 **2.2 Analytical methods**

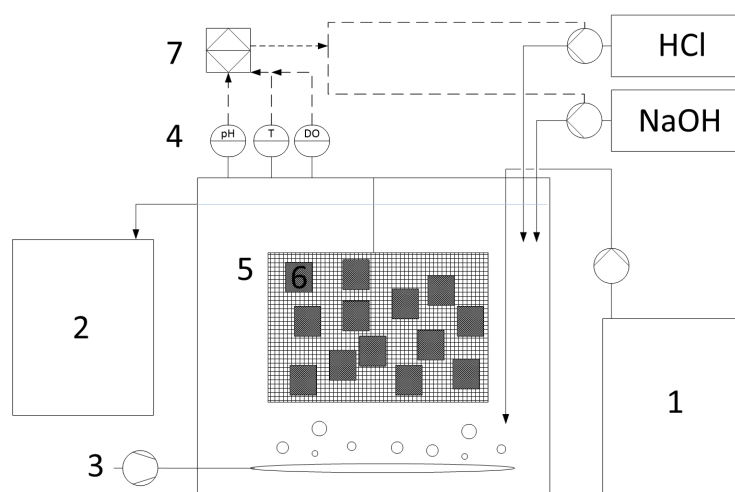
4 The concentrations of dissolved organic carbon (DOC), COD and sCOD in the liquid samples  
5 collected from the system were determined using a TOC-TN analyser (TOC-L series, Shimadzu  
6 Analyser, Japan) and Lovibond COD cuvettes, respectively. The COD and sCOD measurements  
7 were performed immediately after sampling. DOC was measured once the samples were filtered  
8 and frozen at -20°C. DOC and sCOD were measured after the samples were filtered with 0.45 µm  
9 acetate filters.

10 Dry mass content in the colonized PUF cubes was periodically determined. The procedure  
11 followed consisted of drying a colonized PUF cube collected from the reactors at 105°C until  
12 constant weight ( $W_1$ ). Then, the dried colonized PUF cube was submerged into a commercial  
13 solution of sodium hypochlorite and then in ultrapure water to obtain a clean PUF. The clean PUF  
14 was dried again at 105°C until constant weight ( $W_2$ ). The difference between  $W_1$  and  $W_2$  was the  
15 dry mass contained in 1 cm<sup>3</sup> of colonized PUF cube. Total suspended solids (TSS) in the liquid  
16 influent and effluent from both reactors were measured according to standard methods (APHA-  
17 AWWA-WPCF, 2005).

### 18 **2.3 Experimental set-up**

19 Two submerged, packed bed reactors were designed, built and installed at the Autonomous  
20 University of Barcelona (GENOCOV laboratories). Each reactor consisted of a 5 L vessel (4 L of  
21 effective volume and 1 L of headspace) equipped with pH control. NaOH (1M) and HCl (1M)  
22 solutions were dosed to maintain a pH of  $5.8 \pm 0.2$  during the reactors operation. The acidic  
23 environment is generally more favorable for growth and metabolic activities of fungal strains and in  
24 particular for *Aspergillus*, as reported by Mohan et al. (2013).

1 Air was injected through a stone diffuser and manually controlled with a rotameter (100 NL  
2  $\text{h}^{-1}$ ). Outlet air was filtered at 0.2  $\mu\text{m}$  for safety reasons. The air flow allowed for complete mixing  
3 inside the reactor. Inside the vessel, a submerged plastic cylindrical cage was inserted containing  
4 100 immobilised PUF cubes. The cage had a radius of 5 cm and a square grid with 1.25 cm sides.  
5 Probes for continuous pH, dissolved oxygen (DO) and temperature measurements were located  
6 inside the vessels. DO, temperature and pH were measured with a galvanic DO sensor (Oxi 340i  
7 with CelloX 325, WTW, Germany) and pH electrodes (53 33, Crison, Spain). Temperature was  
8 maintained at  $23 \pm 2^\circ\text{C}$  (room temperature) and the DO was maintained above  $6.5 \text{ mg L}^{-1}$  to avoid  
9 oxygen limitation. Liquid samples were also collected from the reactors and feeding tanks three  
10 times per week to analyse, sCOD and DOC. A computer and a SCADA system developed in-house  
11 were used for data acquisition and pH regulation. A schematic of the reactor is shown in Figure 1.



12  
13 **Figure 1.** 1) feeding tank; 2) discharge tank; 3) aerator and air sparger; 4) pH, temperature and DO probes;  
14 5) plastic cage containing PUF carriers; 6) PUF cubes; 7) computer.

## 15 2.4 Process operation

16 As previously mentioned, tests were carried out in two submerged, packed bed reactors run  
17 in parallel. Both reactors were fed with medium solution containing  $1 \text{ g L}^{-1}$  tannin (each reactor  
18 was fed with a different tannin),  $0.1 \text{ g L}^{-1} \text{ NH}_4\text{Cl}$  and  $0.01 \text{ g L}^{-1} \text{ KH}_2\text{PO}_4$  dissolved in tap water. QT  
19 and TT were selected as the most representative recalcitrant compounds of condensed and



1 hydrolysable tannins contained in tannery effluents (Romer et al., 2011; Tilli et al., 2010). The  
2 reactor degrading QT (RQ) was initially fed with a sCOD concentration of  $1433 \pm 56 \text{ mg L}^{-1}$  (1 g  
3 QT L<sup>-1</sup>) while the reactor degrading TT (RT) was initially fed with a sCOD of  $786 \pm 67 \text{ mg L}^{-1}$  (1 g  
4 TT L<sup>-1</sup>). Both concentrations of COD corresponded to the same concentrations of soluble COD  
5 (sCOD) due to the absence of particulate COD (pCOD) in the medium. In RQ the biofilm was  
6 stable and active for five months of operation under non-sterile conditions leading to an average  
7 outlet TSS concentration of  $0.03 \text{ g L}^{-1}$  which means that the outlet COD was mainly sCOD. The  
8 pCOD was less than 4% of the total COD and likely composed of detached biomass. In addition,  
9 the ratio between sCOD and DOC at the reactor inlet was determined to be  $3.02 \pm 0.15$ .  
10 Unfortunately, DOC measurements in the effluent were not reliable due to an unexpected  
11 coagulation phenomenon that occurred after samples storage at  $-20^{\circ}\text{C}$ . The hydraulic retention time  
12 (HRT) and tannin load (measured as volumetric organic loading rate or OLR) were varied during  
13 the operation in both reactors to study their performance in terms of elimination capacity (EC) and  
14 removal efficiency (RE). HRT values ranging from 52h to 9h were set by modifying inlet and outlet  
15 flows with peristaltic pumps (300 series, Watson and Marlow, USA). Tannin concentrations  
16 ranging from 0.1 to  $8 \text{ g L}^{-1}$  were also tested by changing the tannin concentration in the medium  
17 (see Table 1 and Table 2).

18 Despite the fact that both reactors were run in parallel, the operating conditions tested were  
19 different due to the completely different behaviour of the bioreactors. This fact led to design an  
20 independent experimental plan per reactor. Regarding to the RQ, different QT concentrations were  
21 tested to optimise the RE (Table 1) since experiments performed under different HRT in RQ were  
22 not effective (data not shown). Then, only data corresponding to the lowest and highest HRT tested  
23 in the RQ are presented. On the other hand, during the start-up of the RT the biofilm got completely  
24 detached from the PUF cubes and consequently a second start-up was required. Since similar results  
25 were obtained in the first start-up and the second one (see Section 3), the cage was removed and

1 different HRT and TT concentrations were tested. Then, the RT performance and the inhibition  
 2 caused by the presence of TT on the suspended biomass could be assessed (Table 2). After 154 days  
 3 of operation, the simultaneous treatment of QT and TT was tested in RQ (see Table 1) which was  
 4 fed with a medium containing 1 g TT L<sup>-1</sup> and 0.1 g QT L<sup>-1</sup> (corresponding to a total inlet sCOD of  
 5 879 ± 63 mg L<sup>-1</sup>).

6

7 **Table 1.** Operating conditions of RQ fed with QT.

<b>Time (days)</b>	0	56	84	100	106	113	141	154
<b>HRT(h)</b>	28	52	52	52	28	28	28	28
<b>QT concentration (g QT L<sup>-1</sup>)</b>	1	1	0.5	1	0.5	0.2	0.1	0.1*
<b>OLR** (g COD m<sup>-3</sup> h<sup>-1</sup>)</b>	51.1	27.5	12.9	27.5	24.0	7.2	3.6	31.0
<b>COD (mg O<sub>2</sub> L<sup>-1</sup>)</b>	1,433	1,433	672	1,433	672	202	104	869

8 *(\*) the composition of the medium was 1 g TT L<sup>-1</sup> and 0.1 g QT L<sup>-1</sup> (\*\*) OLR is the volumetric organic loading rate*

9

10 **Table 2.** Operating conditions of RT fed with TT.

<b>Time (days)</b>	0	150	155	159	164	169	174	179	187	201	211
<b>HRT(h)</b>	28	52	28	16	12	9	52	28	28	28	28
<b>TT concentration (g TT L<sup>-1</sup>)</b>	1	1	1	1	1	1	1	1	2	8	1
<b>OLR* (g COD m<sup>-3</sup> h<sup>-1</sup>)</b>	27.9	15.0	27.9	48.9	65.2	87	15.0	27.9	55.9	223.9	15.0
<b>COD (mg O<sub>2</sub> L<sup>-1</sup>)</b>	786	786	786	786	786	786	786	786	1,570	6,270	786

11 *(\*) OLR is the volumetric organic loading rate*

## 12 **2.5 Adsorption of tannins on the fungi mycelium**

13 The adsorption capacity of QT and TT on the mycelium was determined in 1-week flask  
 14 tests. The flasks were filled with 100 mL of tannins (QT and TT separately) plus two colonized and

1 autoclaved PUF cubes (with the same liquid/PUF ratio obtained in the bioreactors) and stored with  
2 agitation in an orbital shaker for one week. sCOD was measured per triplicate after 24 h, 72 h and  
3 168 h (less than 15% of the total volume of the liquid was sampled). Flasks without PUF were used  
4 as control. The adsorption of QT and TT to the biomass was assessed considering that part of the  
5 sCOD was adsorbed to the flasks and clean PUF cubes (adsorption obtained in control flasks) and  
6 part of the sCOD was adsorbed to the inactivated biomass. Indeed, the autoclaved biomass  
7 generally show higher adsorption yields with respect to the living one, because of the heat  
8 inactivation cause the cell wall breakdown and, thus, the exposition of the inner binding sites  
9 (Casieri et al., 2008). It must be pointed out that the measure of COD also was substituted by the  
10 measure of sCOD to avoid the effect of the fungal biofilm detachment.

## 11 **2.6 Study of tannins biodegradability with respirometric tests**

12 To estimate the biodegradable sCOD fractions (sbCOD) of QT and TT, a medium solution  
13 containing 10 g tannins L<sup>-1</sup> was prepared, filtered (filter pore of 0.45 µm) and spiked in a  
14 respirometer (MARTINA, Spes Srl, Italy) containing activated sludge. An LFF mode was set to  
15 perform the respirometric tests which means that the oxygen concentration was measured in the  
16 liquid phase (L) and that both liquid and gas phases were flowing (FF). The activated sludge used to  
17 run the respirometric tests was obtained from Cuoiodapur tannery WWTP (Pisa, Italy) to ensure its  
18 acclimation to tannins. A conventional procedure described elsewhere (Andreottola and Esperia,  
19 2001) was applied for COD fractioning.

## 20 **2.7 Metagenomic DNA analysis**

21 Fungi and bacteria communities were analysed from the biofilm developed in the PUF cubes  
22 contained in RQ after 50 days of operation (RQ-28h). Suspended fungi and bacteria communities  
23 developed in RT were also analysed under steady-state conditions on days 159 (RT-28h) and 174  
24 (RT-9h), corresponding to HRTs of 28 h and 9 h, respectively.

1 As mentioned in section 2.3, different biomass was developed in each reactor depending on  
2 the tannin used as substrate. Then, the biofilm immobilized in a PUF cube (RQ) as well as the  
3 suspended biomass (RT) were collected and washed with a phosphate buffered saline solution (0.01  
4 M P-PO<sub>4</sub><sup>3-</sup>). Afterwards, the samples were centrifuged at 8,000 rpm (Thermo Scientific Hareus  
5 Pico17, USA) and the supernatants were discharged. The total DNA of the pelleted biomass was  
6 extracted using a MoBio PowerBiofilm® DNA Isolation Kit (MoBio Laboratories, Inc., Carlsbad,  
7 CA) according to the manufacturer instructions. DNA quality and quantity were measured using a  
8 NanoDrop® spectrophotometer (Thermo-Scientific, USA). A total of 50 ng of DNA was used for  
9 the production paired-end libraries and for the sequencing of the 16S and ITS2 fragments. The  
10 primers used are reported in Table 3. The resulting paired-end libraries were purified using the  
11 Mag-Bind RXNPure Plus magnetic beads (Omega Bio-tek) and pooled in equimolar amounts  
12 according to the Qubit dsDNA high-sensitivity (HS) assay. The pools were sequenced using  
13 Illumina MiSeq PE300 run. The raw data obtained from sequencing was de-multiplexed, and the  
14 sequences of the primers were removed. The fastq files were assembled using a PEAR assembler  
15 (Zhang et al., 2014), followed by a quality check using BBmap tools and FastQC (Bushnell, 2014).  
16 Sequences with a length lower than 300 bp and with a Phred quality score lower than 20 were  
17 removed. Chimera sequences were removed using UCHIME (Edgar, 2016). The 16S sequences  
18 were analysed using QIIME pipeline (Caporaso et al., 2011) in order to obtain the Operational  
19 Taxonomic Units (OTUs) and to analyse the alpha diversity. The Ribosomal Database Project  
20 (RDP) 16S and the UNITE ITS database were used as reference databases in the selection step of  
21 the OTUs, performed by the SortMeRNA methods with a 97% threshold. The OTUs identified with  
22 low-confidence, related to less than 0.1% of the reads (0.1% is the estimate index miss-assignment  
23 on Illumina Miseq), were removed. The numbers of high-confidence sequences were normalised by  
24 using the Deseq2 method. The Shannon, Chao and Simpson alpha-diversity indexes were calculated  
25 by using QIIME and Mothur to characterise the microbial diversity.

1

2 **Table 3.** Primers used to amplify 16S and ITS2 regions to sequence bacterial and fungal rRNA.

---

16 S	515F: 5'- GTG CCA GCM GCC GCG GTA A -3'
	909R: 5' - CCG TCA ATT YHT TTR AGT - 3'
ITS2	ITS4: 5' - TCCTCCGCTTATTGATATGC - 3'
	ITS3: 5' - GCATCGATGAAGAACGCAGC - 3'

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3

## 4 **2.8 Statistical analysis**

5 The standard deviation of sCOD for the inlet solutions of QT was 0.04% and the standard  
6 deviation of sCOD for the inlet solutions of TT was 0.08%. The sCOD in the outlet was measured  
7 in triplicate only in few samples since the reactor was running as a long term experiment and not as  
8 a flask test. The standard deviation of the outlet samples for both reactors were less than 5%.

9

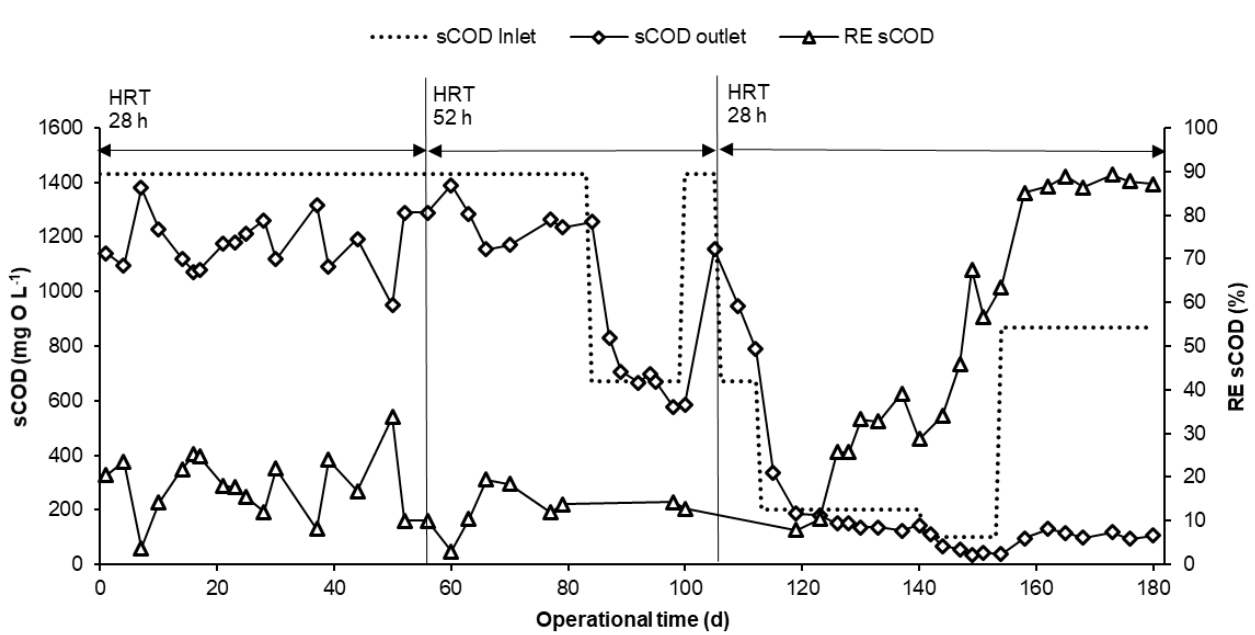
## 10 **3 Results and discussion**

### 11 **3.1 Performance of Quebracho tannin degrading reactor**

12 Despite some oscillations occurred in the first 56 days, the sCOD-RE was ranging from 4%  
13 to 34% which was in average ( $17\% \pm 7\%$ ) higher than that determined from respirometric tests  
14 carried out with an activated sludge sample collected from a tannery WWTP (sCOD-RE of 8%) as  
15 shown in Figure 2. This difference was related to the fungal biomass inoculated, which was more  
16 effective in QT removal than the autochthonous species of the activated sludge from the tannery  
17 WWTP. It may also depend on the chosen operating conditions of the reactor (acidic pH vs neutral  
18 pH), which could have influenced the performance of the inoculated fungal strain and the microbial  
19 communities developed during reactor operation. The operating conditions chosen generated  
20 enough selective pressure in the long-term to drive the evolution of the ecosystem towards an  
21 increase in the diversity of fungi, as described in a subsequent section.

1 As shown in Table 1, after a start-up phase of 56 days, the HRT was increased from 28 to 52  
 2 h maintaining the inlet COD concentration constant in order to verify if a longer contact time could  
 3 improve the low RE obtained during this phase.

4 However, this strategy did not enhance the QT-RE, which pointed at limited removal  
 5 probably due to biomass inhibition and not due to low QT biodegradation kinetics, as was initially  
 6 hypothesized. The average RE obtained during this period was 12%. From day 100 on, the inlet  
 7 concentration of QT was progressively reduced from 1 to 0.1 g QT L<sup>-1</sup> (Table 1). As shown in  
 8 Figure 2, as soon as the inlet concentration was reduced to 202 g COD m<sup>-3</sup> (equivalent to 0.2 g QT  
 9 L<sup>-1</sup>), the RE started to increase, thus confirming the inhibition of process culture by the presence of  
 10 QT. After 141 days of operation, a further inlet QT concentration decrease was set which allowed  
 11 reaching a sCOD-RE increase up to 53% ± 13% in the steady-state and an optimal EC of 9 g sCOD  
 12 m<sup>-3</sup> h<sup>-1</sup>. This result confirmed that the biomass was inhibited by QT, which is in agreement with  
 13 previous works performed at lab-scale with fungi and bacteria using antifouling coatings based on  
 14 tannins (Bellotti et al., 2012; He et al., 2008).



15  
 16 **Figure 2.** Inlet and outlet sCOD and sCOD removal percentage during continuous treatment in bioreactor  
 17 RQ.

1 It is worth mentioning that some PUF cubes were collected from the reactor after 24 h of  
2 operation and cut into two halves. The cross section showed that diffusion and adsorption of tannins  
3 occurred from the external layers of the PUF cube to the inner layers. In order to assess the role of  
4 adsorption in COD removal, the adsorption of QT on mycelium was estimated with flask tests using  
5 colonized and autoclaved PUF cubes at the same volumetric ratio of liquid/PUF set in the reactor.  
6 After 72 h, the adsorption equilibrium was reached with a total sCOD adsorbed of 31%. Then,  
7 adsorption was considered as a relevant phenomenon only at the beginning of the reactor operation  
8 as well as right after each change of influent load (either flow rate or inlet concentration changes).  
9 However, adsorption was considered negligible with respect to the observed COD removal once the  
10 system reached a new steady-state (at least 5-6 days after each stepwise inlet change performed at  
11 28 h HRT).

12 The biomass content of the reactor was estimated on the basis of the dry mass inside PUF  
13 cubes. The dry mass inside PUF increased for 100 days from an initial value of 0.03 g to 0.1 g of  
14 dry mass per cube. Afterwards, dry mass content in each PUF cube increased slowly until the end of  
15 the experiment when a dry mass content of 0.12 g per PUF cube was obtained. The sCOD balance  
16 during the first 100 days of operation (considering the amount of biomass grown in that period) lead  
17 to an observed yield coefficient of 0.19 g COD g<sup>-1</sup> COD. This value was lower than the value  
18 reported in the literature for *Aspergillus niger* (this strain belongs to section Nigri) grown in glucose  
19 which is 0.37 g dry mass g glucose<sup>-1</sup> (which would correspond to approximately 0.51 g COD g<sup>-1</sup>  
20 COD) (Viniestra-González et al., 2003). This result was somehow expected since the higher  
21 complexity of tannins compared to glucose probably caused less growth efficiency.

### 22 **3.2 Performance of Tara tannin degrading reactor**

23 Start-up of RT was performed as the start-up of RQ. However, detachment of the fungal  
24 biofilm and growth of the biomass in suspended form were observed during the first two weeks. A  
25 second start-up was performed with fresh *Aspergillus tubingensis* MUT 990 to verify the

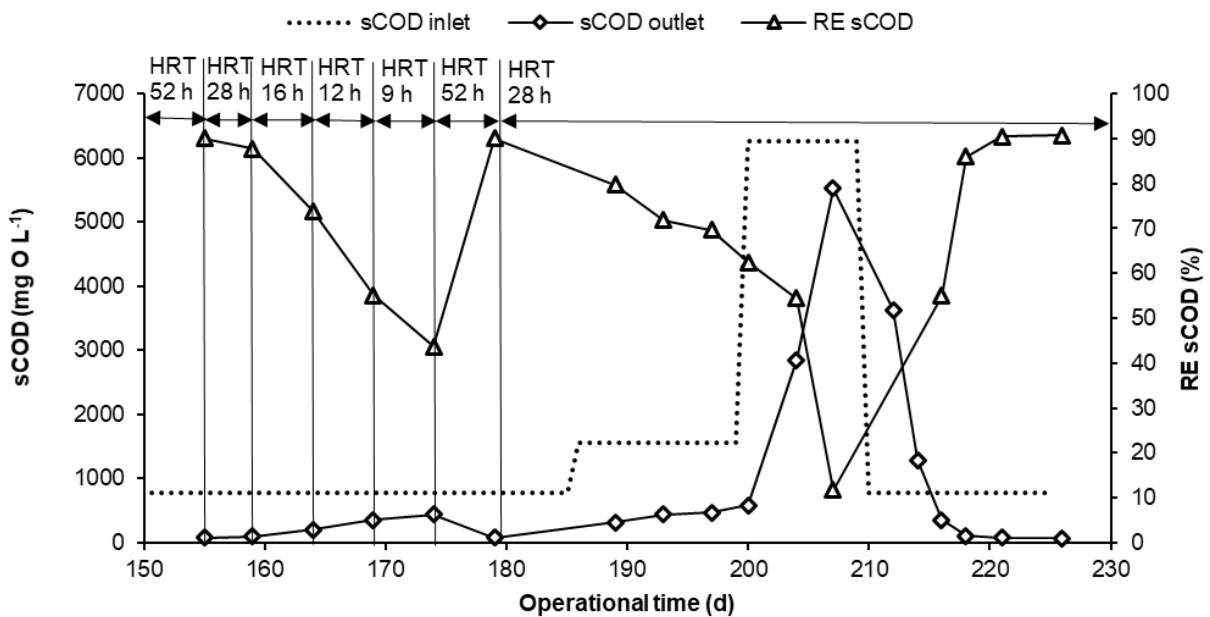
1 repeatability of the observed phenomenon. The same results were obtained; after two weeks the  
2 PUF cubes were almost devoid of biofilm (the dry mass inside the PUF cubes was below the  
3 detection limit of the measurement procedure). The fungal biofilm detachment may have been  
4 caused by bacterial proliferation, which could create competition for the substrate and also damage  
5 the mycelium (Espinosa-Ortiz et al., 2016). Since the biomass was in suspended form, the support  
6 medium was removed, and the reactor became a standard continuous stirred tank reactor (CSTR).  
7 Despite the detachment, the performance of the reactor was promising. After 20 days of operation, a  
8 stable  $90 \pm 2\%$  sCOD-RE was achieved (which corresponded to an EC of  $40 \text{ g sCOD m}^{-3} \text{ h}^{-1}$ ),  
9 much higher than the sCOD-RE of 19% found with respirometric tests using activated sludge from  
10 a tannery WWTP. Though TT (as other common tannins) has toxic or inhibiting effects on various  
11 organisms, some bacteria and yeast have the ability to degrade it. Since gallotannins, one of the  
12 main component of TT, are among the easiest tannins to be biodegraded, it is reasonable to think  
13 that there was a fast development of naturally selected suspended biomass in RT with an active role  
14 in TT biotransformation (Mingshu et al., 2006). The hypothesis that bacteria were the main agent  
15 responsible for TT degradation was further confirmed in the microbial diversity analysis (see  
16 Section 3.4).

17 As mentioned in Section 2 (Materials and Methods), the reactor performance was evaluated  
18 under different tannin loads. Firstly, the HRT was reduced stepwise from 52 h to 9 h while  
19 maintaining the inlet TT concentration (experiments beginning after 154 days of operation).  
20 Secondly, the HRT was set at 28 h (experiments beginning after 179 days of operation), and the  
21 inlet concentration was increased stepwise up to  $8 \text{ g L}^{-1}$  of TT (Table 2).

22  
23 Performance was assessed once the steady-state was reached at each step (i.e. at least four times the  
24 HRT). It was observed that at an HRT of 9 h, the biomass was almost washed out and the sCOD-RE  
25 dropped to 43% (Figure 3). In addition, the ratio between sCOD and DOC in the inlet and outlet



1 were  $2.80 \pm 0.30$  and  $2.69 \pm 0.44$ , respectively, which was another indicator of the large degree of  
 2 mineralisation of TT. Similar results were obtained in terms of DOC (data not shown). As  
 3 mentioned above (Section 2), the performance of the RT was also assessed by setting different inlet  
 4 concentrations in the medium (1, 2, and 8 g TT L<sup>-1</sup>). Inhibition was observed mainly over an inlet  
 5 TT concentration of 2 g L<sup>-1</sup>, in which the RE decreased from 90% to 62%. The system was almost  
 6 completely inhibited at 8 g L<sup>-1</sup>. No toxicity was observed, since the RE recovered after day 210 to  
 7 the initial RE of 90% when the inlet concentration was again reduced to 1 g L<sup>-1</sup>. In fact, the reported  
 8 minimum inhibitory concentrations (MICs) of TT for most bacteria is up to 1 g TT L<sup>-1</sup> (Aguilar-  
 9 galvez et al., 2014). The optimal operating conditions of the system were obtained at 1 g TT L<sup>-1</sup>.



10

11 **Figure 3.** Inlet and outlet sCOD and sCOD-RE profiles obtained from RT performance from day 150 until  
 12 the end of the operation. Vertical lines show the different HRT set during the operation.

### 13 3.3 Simultaneous treatment of TT and QT

14 After the previous tests, RQ was fed with a medium composed of TT (1 g L<sup>-1</sup>) and QT (0.1 g  
 15 L<sup>-1</sup>) (corresponding to a total inlet sCOD of  $879 \pm 63$  mg L<sup>-1</sup>) beginning on day 154 (Figure 2). It  
 16 was also inoculated with the bacteria developed in RT. Since the inhibition caused by QT was  
 17 strong even at a relatively low COD concentration, it was hypothesised that QT contributed to

1 biofilm preservation from bacterial growth. The RE of COD after 4 days of simultaneous QT and  
2 TT feeding was stable at approximately 88% on average. No synergistic effects were detected when  
3 comparing the weighted individual RE during simultaneous treatment with the RE for QT and TT  
4 obtained when fed individually during RQ and RT operation. The simultaneous feeding test lasted  
5 26 days, and the biofilm did not get detached from the PUF cubes, maintaining a stable removal  
6 performance similar to the previous test phase with QT only. Fungi were shown to be more tolerant  
7 to condensed tannins (QT) compared to bacteria and, as previously observed in literature  
8 (Mutabaruka et al., 2007), the presence of condensed tannins could increase the ratio between fungi  
9 and bacteria in a microbial ecosystem. In the literature, TA has been demonstrated to be an efficient  
10 bacterial quorum quenching molecule (QQM) (Patel et al., 2017), and others tannins also show  
11 quorum-sensing inhibitory activity (Yang et al., 2016). Moreover, QT is known for its antifouling  
12 features (Bellotti et al., 2012). Thus, it is possible that QT could also act as an efficient QQM as one  
13 of its mechanisms of action against bacteria. Despite the fact that inhibition of tannins increases  
14 with the concentration, the inhibition mechanism is still partially unclear. These results, together  
15 with the selected operating conditions, could be used as a possible strategy to increase fungal  
16 competitiveness over bacteria and favour the development of a fungal-bacterial consortium that is  
17 symbiotically effective in recalcitrant compounds removal. It is important to highlight that in a co-  
18 culture, microbial interaction could promote alternative intermediate compounds or metabolites and  
19 therefore different degradation pathways of the target pollutant and possible different end products  
20 (Bertrand et al., 2014).

### 21 **3.4 Microbial community analysis**

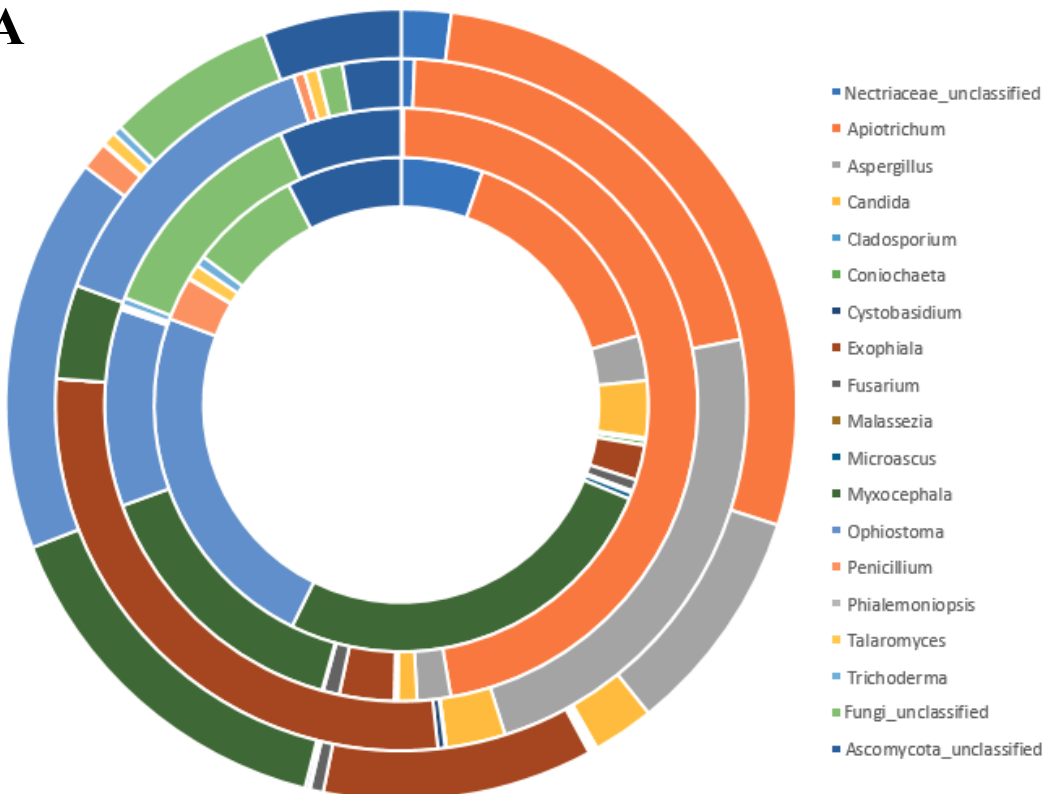
22 The total number of high prokaryotic sequences obtained across the three samples was  
23 229,296, with an average of 76,432 each. Regarding fungi, the amount of high-confidence  
24 sequences across the three samples was 85,259, with an average of 28,419 each. Rarefaction curves  
25 for both bacterial and fungal communities (see Supplementary Information) showed that sequencing

1 results covered bacterial and fungal diversity, since the rarefaction curves tended toward a  
2 saturation plateau. The alpha diversity, Chao, Shannon and Simpson indexes (see Supplementary  
3 Information) showed that the bacterial and fungal communities in the bioreactor were well  
4 characterised. Regarding the bacterial community, 97% of sequences were classified into 113 OTUs  
5 by using the 97% similarity threshold. At phylum level, OTUs were characterised by four phyla.  
6 The *Proteobacteria* phylum was the most abundant, with 88.1% representation among the total  
7 sequences. The other phyla were *Actinobacteria* (8.2%), *Bacterioides* (2.4%) and *Proteobacteria*  
8 (0.5%). At the genus level, most of the sequences were attributed to a taxonomical identity (Figure  
9 4). The most abundant genus was *Novosphingobium* (13.7%). Other representative genera were  
10 *Kaistia* (12.9%), *Comamonas* (11.9%), *Sphingobium* (7%) and *Microbacterium* (5.2%). The above-  
11 mentioned genera were distributed across the samples (from reactor RT, RT-9h and RT-28h, and  
12 reactor RQ, RQ-28h) in different ways: *Novosphingobium* was distributed with a percentage of  
13 33.2% (RT-9h), 5.2% (RT-28h) and 2.7% (RQ-28h); *Kaistia* was very abundant in the RQ reactor  
14 (37.4 % into RQ-28h), while it reached only 0.7% (RT-9h) and 0.6% (RT-28h) of the total in the  
15 RT reactor. On the other hand, the *Comamonas* genus was most abundant in the RT reactor (with  
16 5.6% in RT-9h and 28.5% in RT-28h) and less abundant in the RQ reactor (1.5% in RQ-28h). The  
17 genus *Sphingobium* was recorded at 15% in the RQ reactor, but less abundant in RT reactor (2.7%  
18 in RT-9h and 3.4% in RT-28h). The *Microbacterium* genus was abundant in the RT bioreactor  
19 (15.2% in RT-28h) and less abundant in the RQ bioreactor (less than 1% in RQ-28h). The  
20 remainder of the genera was represented with lower percentages than those reported above or not  
21 classified at the genus level. In the case of the fungal community, 93.1% of the total number of  
22 sequences represented 46 different OTUs. At phylum level, OTUs were divided into two phyla:  
23 *Ascomycota*, with an abundance of 65%, and *Basidiomycota*, with an abundance of 28.1%. The  
24 remaining 6.9% of sequences were uncharacterised. As shown in Figure 4, the *Basidiomycota*  
25 phylum *Apiotrichum* was the most-represented genus (27.9% of total sequences). Within the  
26 phylum *Ascomycota*, the genera *Ophiostoma* (16.2%), *Myxocephala* (15.2%), *Exophiala* (11.1%)

1 and *Aspergillus* (9.3%) were most common. The above-mentioned genera were distributed within  
2 the two bioreactors as follows: *Apiotrichum* genus was predominant in RT, with an abundance of  
3 24.4% in RT-9h, as compared to RQ where it accounted for just 15%. The *Ophiostoma* genus was  
4 most abundant in the RT bioreactor (23.4% in RT-28h) as compared to RQ where it accounted for  
5 just 14%. The *Myxocephala* genus accounted for 26% in the RT bioreactor (RT-9h), and for 4.4%  
6 in RQ. The *Exophiala* genus was more abundant in RQ (27.9%) compared to RT (2.3%). The  
7 *Aspergillus* genus accounted for 23% in RQ-28h and approximately 3% in RT (RT-9h and RT-28h).  
8 Since *Aspergillus tubingensis* was the fungal strain inoculated (immobilised onto the PUF), it might  
9 be reasonable to assume that the strain constituted the majority of the total *Aspergillus ssp.* in the  
10 RQ (23% of the total fungi in the RQ) and RT (about 3% of the total fungi in the RT in both HRT).  
11 Furthermore, biomolecular analysis showed that there was an important fungal consortium  
12 development at steady-state conditions in RQ. Starting from a single fungal strain inoculum, the  
13 *Aspergillus* relative abundance was reduced to a residual value in RT. It is worth mentioning that  
14 among the most abundant bacterial genera found in the reactors (*Kaistia* and *Sphingobium* in RQ;  
15 *Comamonas*, *Novosphingobium* and *Microbacterium* in RT), several species were identified that  
16 have not been previously reported in literature to be tannin degraders. Instead, such species are  
17 reported to be related to the biotransformation of phenols. Likewise, for the fungal genera  
18 *Exophiala*, *Aspergillus*, *Apiotrichum*, and *Ophiostoma* in RQ and *Apiotrichum*, *Myxocephala* and  
19 *Ophiostoma* in RT, several species were identified that were not previously reported as tannin  
20 degraders. In the experimental context, the presence of a diversified microbial community in  
21 addition to the inoculated *Aspergillus* was reasonably linked to the non-sterile nature of the  
22 commercial tannins used and the environmental contamination. The involvement of these  
23 communities in the tannin removal process under the experimental conditions adopted herein is  
24 reasonable. The relationship between organisms in a fungi and bacteria consortium is complex. A  
25 comparison between RQ and RT nevertheless provides some indications about the organisms more  
26 suitable to degrade TT or QT. Further studies are required to reach a better understanding about the

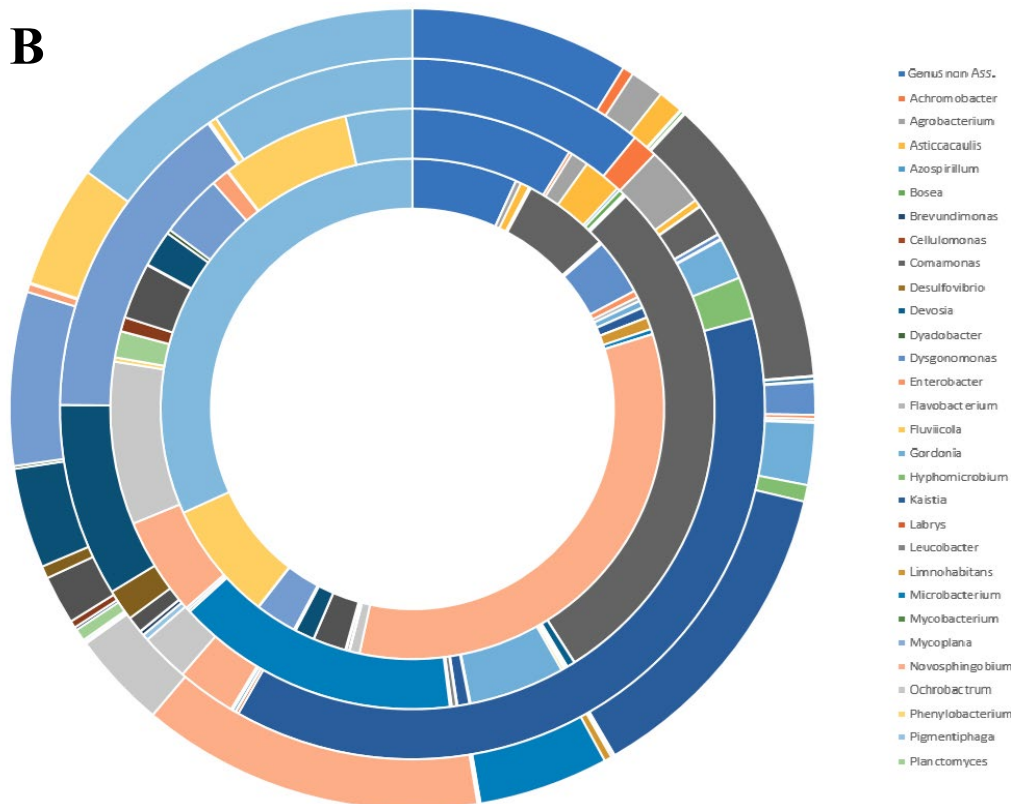
1 interactions between microorganisms and the influence of the operating conditions of the reactor.  
2 As an example, the different HRT in RT was, as expected, a reactor operating condition able to  
3 strongly influence the microbial populations. Thus, *Basidiomycota* phylum was present with an  
4 abundance of 47.3% in RT-28h while decreasing sharply to 15.3% in RT-9h. In fact, the fungi of  
5 *Basidiomycota* phylum are characterised, in general, by a slower growth rate than *Ascomycota*.  
6 Looking to the relative abundance, is it possible that the genera *Novosphingobium*, *Comamonas*,  
7 and *Microbacterium* could better tolerate/degrade TT with respect to QT and could be an invading  
8 species for most fungi. *Kaistia* and *Sphingobium*, on the other hand, could have symbiotic  
9 interactions with fungi and could better tolerate/degrade QT.

A



1

B



2

3 **Figure 4.** Genus percentage in both bioreactors for fungal (A) and bacterial (B) communities. From inside to  
4 outside: The total percentage; RQ-28h, RT-28h and RT-9h.

### 1 3.5 Comparison between RQ and RT

2 The volumetric organic loading rate (OLR) and the volumetric organic removal rate (ORR) in  
3 RQ and RT are compared in Figure 5. Under the conditions tested in this work, a maximum ORR of  
4 37.9 mg sCOD L<sup>-1</sup> h<sup>-1</sup> was obtained for TT while a maximum ORR of 9.1 mg sCOD L<sup>-1</sup> h<sup>-1</sup> was  
5 obtained for QT, which can be used for further sizing of larger scale reactors. Because of the  
6 difficulty to measure biomass concentration in the QT, specific rates could only be compared at an  
7 OLR of 28 mg sCOD L<sup>-1</sup> h<sup>-1</sup>. The specific ORR of RT was 61.2 mg sCOD g<sup>-1</sup> TSS h<sup>-1</sup> while that  
8 estimated for RQ was 2.7 mg sCOD g<sup>-1</sup> TSS h<sup>-1</sup>.

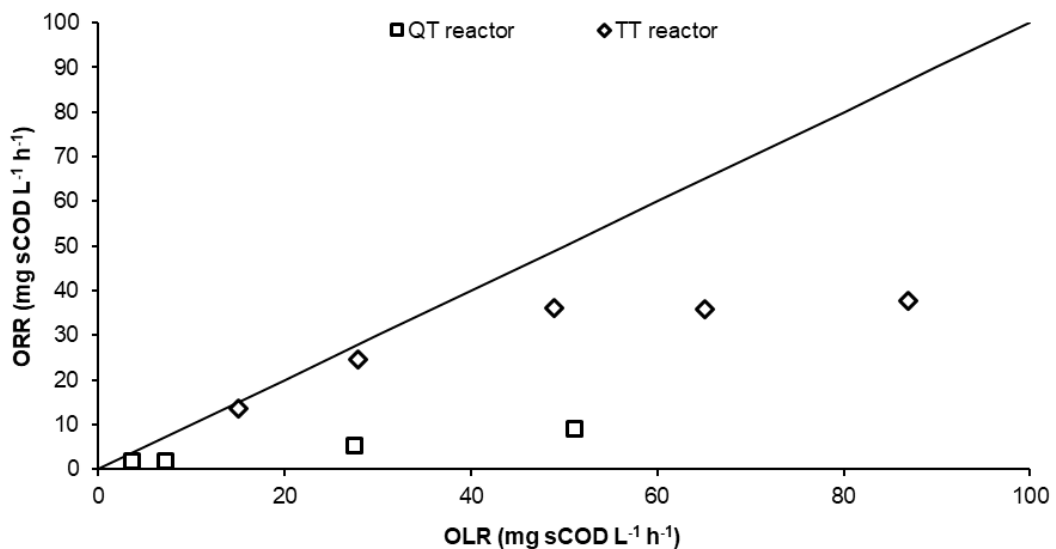
9 The different performance of RQ and RT bioreactors in terms of maximum ORR reached as  
10 well as the diversity changes in both reactors, can be explained because of the different competition  
11 for substrates between fungi and bacteria. It has been previously observed that bacterial growth can  
12 lead to the disruption of the fungal mycelium (Rene et al., 2010). Moreover, TT was mainly  
13 composed of hydrolysable tannin structures based on gallic acid and quinic acid (Garro Galvez et  
14 al., 1997) and, in agreement with previous studies, the minimum inhibitory concentration (MIC) of  
15 TT for most bacteria was found to be in the range of 0.16 to 16 g TT L<sup>-1</sup> (Aguilar-galvez et al.,  
16 2014). As a consequence, the selected fungal strain was outcompeted by bacterial strains able to  
17 degrade TT (Mingshu et al., 2006), with a MIC higher than the concentration used in the medium  
18 solution (1 g TT L<sup>-1</sup>). The operating conditions in RT (such as slightly acidic pH) allowed the  
19 development of bacterial-based suspended biomass with faster growth kinetics that lead to larger  
20 volumetric and specific rates compared to the RQ, which was mostly a fungal based reactor. In the  
21 RQ, the RE was satisfactory only when QT was fed at a concentration below 0.2 g QT L<sup>-1</sup>. A QT  
22 removal rate of 53% was obtained with 0.1 g L<sup>-1</sup> concentration in the inlet solution. Then, the RQ  
23 fed with QT outlasted the colonisation of bacteria and allowed a stable fungal biofilm thanks to the  
24 operating conditions, the QT concentration and consequently the condensed tannins load. This

1 finding was also verified during the simultaneous feeding of TT and QT since biofilm was  
2 preserved in the PUF cubes as well as the suspended biomass in the liquid phase.

3 Overall, results obtained in the present study represent an important progress in the  
4 application of biological processes in tannins removal from wastewater. To the authors' knowledge,  
5 this work is the first to report the continuous treatment of QT and TT, providing removal rates and  
6 operating conditions that could aid the design of industrial scale reactors. The reactor design and  
7 operating conditions demonstrated successful, stable tannin removal under non-sterile conditions,  
8 even without the addition of any co-substrate. In both RT and RQ, the development of a fungal and  
9 bacterial consortium was observed. Moreover, results allow designing a process for simultaneous  
10 treatment of different types of tannins as well as to expand the range of application of fungal-based  
11 technologies.

12

13



14

15 **Figure 5.** Overall performance rates of RQ and RT reactors.

16



## 1    **4    Conclusions**

2    The effects of two different types of tannins, condensed and hydrolysable, on the fungal biomass  
3    were investigated and separately tested in two reactors. Fungal and bacterial consortia developed in  
4    the long-term operation of bioreactors fed with QT and TT after initial inoculation with *Aspergillus*  
5    *tubingensis*. Reactor fed with QT outlasted the colonization of bacteria and allowed a stable fungal  
6    biofilm able to remove QT with RE up to 53%. Reactor fed with TT suffered the detachment of the  
7    inoculated fungal biofilm after start-up. Fungi were outcompeted by bacteria as demonstrated  
8    through biomolecular analysis leading to a stable culture of suspended biomass in the long run that  
9    reached 90% removal of hydrolysable tannins. Additionally, co-treatment of both tannins was  
10    demonstrated feasible. Fungi have been shown to be more resistant to the inhibitory effect of QT  
11    than bacteria, and the QT concentration therefore allowed for the maintenance of a stable fungal  
12    biofilm in non-sterile conditions. The study provided promising results application of the new  
13    technology based on fungal biofilms reactors.

## 15    **Acknowledgments**

16    The authors thank the Miur (Fir project RBFR13V3CH), the UE (Marie Curie Irses Carbala project  
17    295176) and the Tuscany region (Lightan POR FESR 2014-2020). Moreover, the authors thank  
18    Andrea Nardo for the support given with respirometric tests; Clara Reino for the support given with  
19    DNA extraction; Chimont International Spa for providing the tannins and the Mycoteca  
20    Universitatis Taurinensis (MUT) for providing the fungal strain.

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