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5 **Effects of wood amendments on the degradation of terbuthylazine and on soil**  
6 **microbial community activity in a clay loam soil**

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18 **Abstract** The herbicide terbuthylazine is widely used within the EU; however its frequent  
19 detection in surface and groundwater, together with its intrinsic toxicological properties, may  
20 pose a risk both for human and environmental health. Organic amendments have recently been  
21 proposed as a possible herbicide sorbent in soil, in order to limit herbicide movement from soil  
22 to water. The environmental fate of terbuthylazine depends not only in its mobility, but also its  
23 persistence. The latter is directly dependent on microbial degradation. For this reason the effects  
24 of pine and oak residues on terbuthylazine soil microbial community functioning and on the  
25 potential of this community for terbuthylazine degradation were studied. For this purpose,  
26 degradation kinetics, soil dehydrogenase activity and the number of live bacteria were assessed  
27 in a clay-loam soil treated with terbuthylazine and either amended with pine or oak wood or  
28 unamended (sterilized and non-sterilized). At day 65, 85% of the herbicide applied still persisted  
29 in the sterile soil, 73% in the pine-amended one and 63% in the oak-amended and unamended  
30 ones. Pine residues increased the sorption of terbuthylazine to soil and hampered microbial  
31 degradation owing to its high terbuthylazine sorption capacity and a decrease in the  
32 bioavailability of the herbicide. On the contrary, in the presence of oak residues the herbicide  
33 sorption did not increase significantly. The overall results confirm the active role of the soil  
34 microbial community in terbuthylazine degradation in amended and unamended soils and in a  
35 liquid enrichment culture performed using an aliquot of the same soil as the inoculum. In this  
36 clay loam soil, in the absence of amendments, the herbicide was found to be quite persistent ( $t_{1/2}$   
37 > 95 days), while in the enrichment culture the same natural soil bacterial community was able to  
38 halve terbuthylazine in 24 days. The high terbuthylazine persistence in this soil was presumably  
39 ascribable to its texture and in particular to the mineralogy of the clay fraction.

40

41 **Keywords** Terbuthylazine, Degradation, Soil texture, Organic amendments, Pine and oak  
42 residues, Microbial community activity.

## 43 **1 Introduction**

44

45 Terbutylazine is an *s*-triazine herbicide widely used in agriculture to control grass and broad-  
46 leaved weeds in a variety of crops. In Italy terbutylazine is used in maize and sorghum (Fait et  
47 al. 2010), and in Spain is used also in olive tree cultures (Cabrera et al. 2007, 2008).

48 The fate and behaviour of terbutylazine in soil have raised environmental concern  
49 because, together with its metabolite desethylterbutylazine (DET), it has been frequently found  
50 in surface water and groundwater at levels above  $0.1 \mu\text{g L}^{-1}$ , which is the limit established in the  
51 EU for individual pesticides in drinking water (Guzzella et al. 2006; Hildebrandt et al. 2006).

52 European Food Safety Authority (EFSA) has recently reported that terbutylazine poses  
53 high long-term risks for mammals, aquatic organisms, non-target plants, earthworms (EFSA  
54 2001) and can have genotoxic effects (Mladinic et al. 2012). The fact that it has been recently  
55 (16 August 2011) re-evaluated and its placing in the EU market approved until 2021 by  
56 Commission Implementing Regulation 820/2011 makes its environmental occurrence, together  
57 with its toxicological relevant metabolite, DET, a risk both for the environment and human  
58 health.

59 Terbutylazine degradation depends on both abiotic and especially biotic processes,  
60 which are responsible for its complete degradation. The more it is degraded in soil, the less the  
61 likelihood of it being leached to groundwater or run off to surface water. Biodegradation and  
62 mineralization of *s*-triazines have been shown to be carried out by bacterial consortia and by  
63 strains isolated from contaminated sites (Grenni et al 2009a; Barra Caracciolo et al. 2010). The  
64 formation of cyanuric acid as an intermediate and then its transformation to biuret was found to  
65 be the common step before mineralization, although the sequence of pathway steps varied among  
66 degraders (Santiago-Mora et al. 2005; Barra Caracciolo et al. 2010). Degradation rates in  
67 agricultural soils may depend on the history of terbutylazine treatment, which may increase the

68 soil self-remediation potential (Rhine et al. 2003) and on the specific soil characteristics (soil  
69 depth, pH, temperature, water content, presence of exogenous nitrogen, organic matter content  
70 and texture) which can directly or indirectly influence the degradation process (Di Corcia et al.  
71 1999; Barra Caracciolo et al. 2010; Kodešová et al. 2011).

72 In soil, one of the primary mechanisms of its transformation is a biotic oxidative N-  
73 deethylation with the formation of desethylterbutylazine, DET (Di Corcia et al. 1999).  
74 Monitoring data show that DET is frequently present in groundwater and its concentration is  
75 often higher than its parent compound; this phenomenon is due to the intrinsic characteristics of  
76 DET (e.g. water solubility and soil organic carbon partition coefficient) which determine its  
77 lower adsorption and higher mobility in soils (Bottoni et al. 1996; Guzzella et al. 2003; Barra  
78 Caracciolo et al. 2005a; EFSA 2011; FOOTPRINT, 2011).

79 Point-source contamination by pesticides has been identified as a major concern  
80 contributing significantly to the deterioration in the quality of natural water resources. Indeed,  
81 monitoring studies have clearly shown that pesticide point-source contamination produced by  
82 improper pesticide handling before or after their field application (e.g. spills, uncontrolled  
83 disposal, equipment washing water, etc.) has resulted in the frequent detection of high  
84 concentrations of pesticides in natural water resources (De Wilde, 2007; Fait et al. 2010;  
85 Kravvariti et al. 2010). The addition of exogenous organic matter of different origin, including  
86 wastes, may prevent the mobility of pesticides released in soil from point as well as from non-  
87 point sources of contamination and enhance their biodegradation (Rodríguez-Cruz et al. 2007a;  
88 Delgado-Moreno and Peña 2009). In recent years different low-cost sorbent systems (biobed,  
89 biomassbed, biofilter) have been developed to minimize point sources of pesticide pollution.  
90 These systems consist of a mixture of different organic biomaterials and soil which can retain  
91 and degrade pesticides (Kravvariti et al. 2010; Castillo et al. 2008). The addition of organic  
92 amendments to soil can affect the biodegradation of pesticides owing to the application of an

93 additional source of organic matter and sometimes microorganisms (Briceño et al. 2007; Kan et  
94 al. 2007) with the result of accelerating the degradation of pesticides (Kravvariti et al. 2010;  
95 López-Piñeiro et al. 2011). In other cases, the addition of an organic residue to soil can lead, by  
96 decreasing the bioavailability of pesticides owing to their increased sorption capacity, to a  
97 decrease in pesticide degradation (Moorman et al. 2001; Briceño et al. 2007; Grenni et al 2009a;  
98 Kravvariti et al. 2010).

99 Pine and oak wood residues have recently been shown to be effective low-cost sorbents  
100 of the herbicide linuron in a sandy-loam soil (Grenni et al 2009a). The greater adsorption of  
101 linuron to pine than oak was related to its higher lignin content, the hydrophobic wood  
102 component (Rodríguez-Cruz et al. 2007b). However, the influence of the addition to soil of these  
103 wood residues on the adsorption and degradation of the herbicide terbuthylazine has not been  
104 studied so far.

105 In the present work the degradation of terbuthylazine was evaluated in an agricultural  
106 clay loam soil, where the groundwater beneath is found to be chronically contaminated by this  
107 herbicide and its metabolite DET (Barra Caracciolo et al., 2010). In order to assess the  
108 applicability of pine and oak amendments for the immobilization of herbicide in this soil, an  
109 experimental set-up, consisting of soil microcosms treated with terbuthylazine and either  
110 amended with pine or oak residues or unamended, was performed. The ability of  
111 microbiologically active soils (amended or unamended) to degrade the herbicide was evaluated  
112 by comparing the half-lives ( $t_{1/2}$ ) in microcosm studies in the various scenarios (pine-amended,  
113 oak-amended, and unamended) to that in sterile soil. Moreover, the effects of these amendments  
114 on soil bacterial community activities, such as dehydrogenase and viability, were also assessed.

115 Finally, an enrichment culture was set up with terbuthylazine as the sole carbon source,  
116 using aliquots of the same soil as the inoculum. This experiment was performed in order to

117 evaluate the capability of this soil microbial community to degrade the herbicide and to grow on  
118 it in a liquid culture.

119

## 120 **2 Material and Methods**

121

### 122 2.1 Experimental site

123

124 The criteria used for the selection of the site (located near Assisi, Central Italy) were the  
125 presence of intensive agriculture with previous terbuthylazine application and a shallow alluvial  
126 aquifer (water table at 12 m depth, alkaline-bicarbonate geochemical facies) vulnerable to  
127 herbicide contamination (Daly et al. 2002). According to the Umbria Regional Environmental  
128 Agency's monitoring surveys (2000–2010), terbuthylazine and its metabolite,  
129 desethylterbuthylazine, are commonly found in this groundwater ( $> 0.1 \mu\text{g L}^{-1}$  parametric value).  
130 It is also common to find significant nitrate contamination at this site ( $> 100 \mu\text{g L}^{-1}$ ).

131

### 132 2.2 Soil and wood samples

133

134 Soil samples were collected from the surface layer (0-20 cm depth) and left to dry at room  
135 temperature, then sieved ( $< 2 \text{ mm}$ ) and analysed for their physiochemical characteristics. The  
136 soil was classified according to F.A.O. World Soil Classification as Calcaric Cambisol  
137 (Giovagnotti and Calandra 1994) and the soil texture was classified as clay loam according to  
138 USDA (22.9% sand, 43.2% silt and 33.9% clay). The organic carbon and nitrogen content were  
139 1.87% and 0.13%, respectively, and the pH 7.7. The clay minerals in the soil were  
140 montmorillonite (9.25%), illite (20.6%) and kaolinite (4.01%).

141 Pine and oak wood residues (< 1 mm) were selected as the organic soil amendments  
142 because of their different lignin contents of 24.4% and 18.2%, respectively (Rodríguez-Cruz et  
143 al. 2007b). They were obtained from a local company in Salamanca (Spain). The amended soils  
144 were prepared by uniformly mixing soil with oak or pine (5% w/w).

145 The organic carbon and the pH of wood residues, determined in a previous work  
146 (Rodríguez-Cruz et al. 2007b) are reported in Table 1. The wood amended occurrence increased  
147 the organic carbon content of the soil and did not affect the soil pH (Table 1).

148

### 149 2.3 Chemicals

150

151 Terbutylazine (*N*<sup>2</sup>-*tert*-butyl-6-chloro-*N*<sup>4</sup>-ethyl-1,3,5-triazine-2,4-diamine) and its main  
152 metabolites desethylterbutylazine (DET) and desethyldebutylterbutylazine (DEDT), were  
153 supplied by Dr. Ehrenstorfer GmbH (Augsburg, Germany) (> 98.0% purity). Terbutylazine is a  
154 colourless powder with a water solubility of 8.5 mg L<sup>-1</sup> at 20°C and log K<sub>ow</sub> of 3 (Tomlin et al.  
155 2003; Rodríguez-Cruz et al. 2007b).

156

### 157 2.4 Laboratory degradation experiments with unamended and amended soils

158

159 The herbicide degradation experiment was conducted in duplicate microcosms for each different  
160 treatment in accordance with SETAC guidelines (Lynch 1995) and some previous experiments  
161 (Grenni et al 2009a; Barra Caracciolo et al. 2005a; 2005b). Terbutylazine was added to  
162 unamended or amended soil (200 g) to obtain a final concentration of 1.5 mg kg<sup>-1</sup>. Initially some  
163 soil samples were sterilized twice (autoclaved 120 ± 2°C, 20 min) and then treated with  
164 terbutylazine (SST); other soil samples were only treated with terbutylazine (ST); others were  
165 treated with both terbutylazine and pine (SPT) or oak (SOT) sawdust (5% w/w); lastly,

166 microbiological control soils (S) were prepared with only water and with water and pine (SP) or  
167 oak (SO) sawdust. All soils were thoroughly stirred with a sterilized spatula and the water added  
168 was in all cases sterilized by filtration (0.22  $\mu\text{m}$ ). The final moisture content was adjusted to 60%  
169 of the maximum soil water holding capacity ( $\text{WHC}_{\text{max}}$ ).

170 The soils were maintained in beakers closed with a sterilized cotton plug wrapped in  
171 gauze to allow air exchange. The soil moisture was kept constant during the entire period of the  
172 experiments by periodically weighing and replacing any losses with sterile water. Samples were  
173 incubated at  $20 \pm 2^\circ\text{C}$  in the dark. Solutions and instruments were sterilized and all steps were  
174 performed in a sterile cabinet. The overall experimental set consisted of 14 microcosms (two for  
175 each of the 7 different treatments, S, ST, SST, SOT, SPT, SO, SP). For each chemical or  
176 microbiological analysis we collected 2 sub-samples from each of the two replicate microcosms.  
177 Consequently, each value reported is the average of a total of four data. Sampling was performed  
178 at different times (0, 6, 12, 20, 33, 49, and 64 days) for both the chemical and microbiological  
179 analyses.

180

## 181 2.5 Enrichment culture on terbuthylazine

182

183 An enrichment culture experiment was performed in order to evaluate the occurrence of natural  
184 bacterial populations able to degrade the herbicide and to grow on it as the sole carbon source.

185 Three soil samples (3.5 g each) were inoculated into 30 mL of a liquid medium MB 1 $\times$   
186 ( $\text{K}_2\text{HPO}_4$ , 1.6 g  $\text{L}^{-1}$ ;  $\text{KH}_2\text{PO}_4$ , 0.4 g  $\text{L}^{-1}$ ;  $\text{CaSO}_4 \times 2\text{H}_2\text{O}$ , 0.1 g  $\text{L}^{-1}$ ;  $\text{MgSO}_4 \times 7\text{H}_2\text{O}$ , 1.0 g  $\text{L}^{-1}$ ;  
187  $\text{FeSO}_4 \times 7\text{H}_2\text{O}$ , 0.02 g  $\text{L}^{-1}$ ;  $(\text{NH}_4)_2\text{SO}_4$ , 2g  $\text{L}^{-1}$ ) with the herbicide terbuthylazine at a  
188 concentration of 2 mg  $\text{L}^{-1}$ , as performed in similar works (Sánchez et al. 2005; Grenni et al.  
189 2009b). Two other flasks containing a previously sterilized liquid medium (MB1 $\times$ ) and the



190 herbicide at the same concentration ( $2 \text{ mg L}^{-1}$ ) were used as controls and monitored for chemical  
191 analysis until 85 days.

192 The experimental set consisted of flasks (50 mL capacity) which were incubated at  $20^{\circ}\text{C}$   
193 in the dark and gently shaken. Samples for chemical and microbiological analysis were collected  
194 at selected times until the degradation of at least 80% of the initial concentration was reached.  
195 The concentrations of the terbuthylazine and its metabolite DET were measured immediately  
196 after the treatment and at different times (0, 3, 8, 15, 23, 30, 37, 44, 51, and 58 days). The  
197 bacterial growth of the soil microbial pool was monitored during the incubation period by the  
198 epifluorescence direct count method.

199

## 200 2.6 Sorption studies

201

202 The sorption of terbuthylazine both to wood residues and to wood-amended soils was determined  
203 using a batch equilibrium method described in detail in Rodriguez-Cruz et al. (2007b). Briefly,  
204 triplicates 100 mg wood samples or triplicates 5 g unamended and wood-amended (5 %) soils  
205 were equilibrated with 10 mL of an aqueous solution of terbuthylazine at an initial concentration  
206 ( $C_i$ ) of  $7 \text{ mg L}^{-1}$ . Preliminary experiments pointed out that a contact for 24 h was long enough for  
207 the equilibrium to be reached. The pesticide amount adsorbed ( $C_s$ ) was considered to be the  
208 difference between that initially present in the solution ( $C_i$ ) and that remaining after equilibration  
209 ( $C_e$ ) with the wood or wood-amended soil. Sorption distribution coefficients,  $K_d$ , were  
210 calculated from the relationship between  $C_s$  and  $C_e$  ( $K_d = C_s/C_e$ ), and were considered a  
211 measure of pesticide adsorption capacity by the wood or wood-amended soils. All measurements  
212 were carried out in duplicate. The quantification of terbuthylazine was performed by HPLC-MS  
213 in a Waters chromatograph (Waters Assoc., Milford, MA) equipped with a model e2695  
214 multisolvent delivery and autosampler system attached to a ZQ mass spectrometer detector

215 (MS), and Empower software as the data acquisition and processing system. A Waters Symmetry  
216 C18 (75 mm x 4.6 mm I.D., 3.5  $\mu\text{m}$ ) column was used at ambient temperature. The mobile phase  
217 was 80:20 methanol/water in a 0.1% formic acid solution. The flow rate of the mobile phase was  
218 0.3  $\text{mL min}^{-1}$  and the sample injection volume was 20  $\mu\text{L}$ . Detection by HPLC/MS to quantify  
219 terbuthylazine was monitoring the positive molecular ion ( $m/z$ ) 230. The quantification of  
220 terbuthylazine was done with the external standard method using the calibration curves obtained  
221 by the injection of standard solutions at a concentration range between 0.05 and 1  $\mu\text{g mL}^{-1}$   
222 ( $r^2 > 0.99$ ).

223

## 224 2.7 Herbicide analysis

225

226 Soil sub-samples (1 g) were taken from each microcosm and shaken at 60 rpm with 6 mL of  
227 methanol for 24 h at 20°C for residue analysis. Samples were centrifuged and 4 mL of each  
228 supernatant were evaporated at 30°C under nitrogen stream (Concentrator EVA VLM-EC-2V-  
229 130, Germany) and re-dissolved in 0.5 mL of methanol for analysis.

230 Quantitative determination of terbuthylazine and its main metabolite DET and its further  
231 transformation product desethyldebutylterbuthylazine (DEDT) in soil and in enrichment culture  
232 samples was performed by GC-MS in a 7890A Agilent gas chromatograph coupled to a 5975C  
233 Agilent mass spectrometer (Agilent Technologies, Avondale, USA) with an Agilent 7683  
234 autosampler. Chromatographic separation was performed on a 30 m  $\times$  0.25 mm I.D, 0.25  $\mu\text{m}$   
235 film thickness HP-5MS capillary column. The carrier gas was helium at a rate of 1  $\text{mL min}^{-1}$ . A  
236 split/splitless injector was used in the split-less mode. A sample volume of 0.2  $\mu\text{L}$  was injected  
237 in the splitless mode with an injector temperature of 225°C. The following temperature program  
238 was used: the temperature was increased from 100°C to 150°C at 50°C  $\text{min}^{-1}$  and maintained for  
239 1 min, then at 5°C  $\text{min}^{-1}$  to 200°C and finally increased to 290°C at 30°C  $\text{min}^{-1}$  and maintained

240 for 1 min. The quadrupole mass spectrometer was operated in the electron impact ionization (EI)  
241 mode at 70 eV. The transfer line and the injector were set up at 250°C and the source and the  
242 quadrupole were at 230°C and 150°C, respectively. Measurements in the GC-MS were  
243 performed in the single-ion monitoring (SIM) mode. The more abundant ions were chosen for  
244 quantification (terbuthylazine m/z 214, DET m/z 186 and DEDT m/z 173). The quantification  
245 was carried out by double injection. Recoveries for terbuthylazine, DET and DEDT were 90%,  
246 80%, 72% respectively. Samples were extracted and analysed in duplicate. The quantification of  
247 terbuthylazine and its metabolites was performed by the external standard method using the  
248 calibration curves obtained by the injection of standard solutions at a concentration range  
249 between 0.1 and 1  $\mu\text{g mL}^{-1}$  ( $r^2 > 0.99$ ).

250

## 251 2.8 Soil dehydrogenase activity, total cell number and cell viability

252

253 At different times (0, 6, 12, 20, 33, 49, and 64 days) after herbicide application, the  
254 dehydrogenase activity, total cell number and cell viability were assessed. Soil dehydrogenase  
255 activity was determined following the method described by Tabatabai (1994). The method is  
256 based on extraction and colorimetric determination of the intensely coloured 2,3,5-triphenyl  
257 formazan (TPF) produced from the reduction of colourless 2,3,5-triphenyltetrazolium chloride  
258 (TTC) in soils after an 24 h incubation at 37°C in the dark. Results were expressed as  $\mu\text{g TPF g}^{-1}$   
259 dry soil. Measurements were performed in duplicate for each microcosm.

260 The total cell number (No. bacteria  $\text{g}^{-1}$  dry soil) was assessed (in duplicate for each  
261 microcosm) in 1 g of fixed soil with the epifluorescence direct count method, using 4',6-  
262 diamidino-2-phenylindole (DAPI) as the DNA fluorescent agent, as reported in detail in previous  
263 works (Barra Caracciolo et al. 2005a; 2005b).

264 Cell viability (% Live/Live+Dead) was measured in 1 g of fresh soil (in duplicate for  
265 each microcosm) using two fluorescent dyes, SYBR Green II and propidium iodide (Sigma-  
266 Aldrich, Germany) in order to distinguish between viable (green) and dead (red) cells under a  
267 fluorescence microscope (Leica DM 4000B Leica Microsystems GmbH, Wetzlar, Germany), as  
268 reported in a previous work (Grenni et al. 2009a).

269

## 270 2.9 Statistical analysis

271

272 Analysis of variance (two-way analysis of variance) was used to determine the significant  
273 differences ( $p < 0.05$ ) in the dehydrogenase activity, bacterial number and viability among the  
274 different soil treatments during the experimental period, using the Statistical software  
275 SIGMASTAT (version 3.0).

276

## 277 **3 Results**

278

### 279 3.1 Degradation of terbuthylazine in soil microcosm experiments

280

281 The decrease in the herbicide concentrations (expressed in percentages of residual  
282 terbuthylazine) over a period of 64 days in unamended and sterile soil (SST), unamended soil  
283 (ST) and soils amended with oak (SOT) or pine (SPT) is shown in Fig. 1. Degradation of  
284 terbuthylazine was fitted first-order kinetics:  $C_t = C_0 e^{-kt}$ , where  $C_t$  is the concentration at time  $t$ ,  
285  $C_0$  is the initial concentration at  $t=0$ , and  $k$  is the constant rate. In the ST and SOT treatments  
286 there was a phase of slow degradation at the beginning of the incubation. The theoretical half-life  
287 values ( $t_{1/2}$ ) calculated from the corresponding exponential equations, obtained from the  
288 regressions between concentrations and time, were:  $257 \pm 27$  d ( $r^2 = 0.90$ ,  $p < 0.01$ ) in SST < 161

289  $\pm 38$  d ( $r^2 = 0.75$ ,  $p < 0.05$ ) in SPT  $< 105 \pm 10$  d ( $r^2 = 0.94$ ,  $p < 0.01$ ) in ST  $< 95 \pm 7$  d ( $r^2 = 0.93$ ,  
290  $p < 0.01$ ) in SOT. These values suggest that the herbicide was quite persistent in the soil studied.  
291 At the end of the experiment (day 64) the lowest herbicide concentrations observed were in the  
292 unamended (ST) and oak-amended (SOT) soils and about 40% of the initial concentrations were  
293 degraded.

294 DET and its further metabolite DEDT were found in microbiologically active soils (ST,  
295 SOT, SPT) during the degradation process and the amounts of these metabolites generally  
296 increased over time, as shown in Fig. 2. Both metabolites were found in higher concentrations in  
297 SOT and ST from day 49 to day 64, in line with a higher terbuthylazine degradation observed in  
298 these treatments at the end of the incubation.

299

### 300 3.2 Microbiological analysis: soil dehydrogenase activity, total cell number and cell viability

301

302 The herbicide effects on soil dehydrogenase activity, total cell number and cell viability were  
303 studied in all herbicide-treated soils and compared with non-treated ones. Fig. 3 shows soil  
304 dehydrogenase activity ( $\mu\text{g TPF g}^{-1}$  dry soil) in relation to time in the terbuthylazine treated soils  
305 (A) and in the control ones (B).

306 A significant difference ( $p < 0.05$ ) in dehydrogenase activity was observed among the  
307 different soil treatments. Dehydrogenase activity was significantly higher in all the soils  
308 amended with pine or oak than in unamended ones. After 28 days in the SOT treatment a  
309 significant increase ( $p < 0.05$ ) of dehydrogenase activity was observed compared to the other  
310 treatments.

311 The initial total cell numbers (No. bacteria  $\text{g}^{-1}$  dry soil) obtained by DAPI counts were  
312 higher ( $p < 0.05$ ) in all the amended soils (SP:  $4.8 \times 10^7 \pm 3.2 \times 10^6$ ; SO:  $4.0 \times 10^7 \pm 2.4 \times 10^6$ ).

313 However, cell numbers were subsequently not significantly different in the various treatments  
314 (data not shown).

315 The cell viability values (% Live/Live+Dead) are reported in Fig. 4. In the presence of  
316 the amendments a transient decrease in viability at day 6 was observed; this was particularly  
317 evident in the SOT and SPT treatments.

318

### 319 3.3 Degradation of terbuthylazine in the enrichment culture

320

321 The soil microbial pool in the enrichment culture was able to degrade the terbuthylazine with a  
322  $t_{1/2}$  of  $24 \pm 2$  days ( $r^2 = 0.99$ ) (Fig. 5A on the left axis). In contrast, after 85 days in the sterile  
323 medium more than 98% of the initial concentration of the herbicide was still present. The total  
324 cell number (No. bacteria  $\text{mL}^{-1}$ ) of the microbial pool was assessed during the experimental time.  
325 A positive correlation ( $p < 0.05$ ) was found between the terbuthylazine concentration (% of TBA  
326 applied) and the cell number (Fig. 5A), indicating that the soil bacterial populations were able to  
327 grow using the herbicide as a carbon source.

328 The metabolite DET was immediately detected and it was exclusively found in the  
329 presence of the soil microbial pool (Fig. 5B). Its formation was correlated ( $p < 0.05$ ) to the  
330 terbuthylazine degradation.

331

### 332 3.4 Sorption of terbuthylazine by wood residues and soil

333

334 The sorption of terbuthylazine by the woods used and the unamended and wood-amended soils  
335 are reported in Table 1. The  $K_d$  value for the sorption of terbuthylazine by pine was much higher  
336 than for oak. Similarly, the sorption of terbuthylazine by the pine amended soil ( $11.7 \pm 2.63$ ) was

337 higher than the oak amended one ( $5.3 \pm 0.38$ ). However, the latter  $K_d$  value was not significantly  
338 different from the unamended soil ( $4.27 \pm 1.08$ ).

339

#### 340 **4 Discussion**

341 The overall results show that the microbial community had a significant role in the  
342 terbuthylazine degradation, as shown when comparing the degradation results for sterile soil and  
343 microbiologically active soil in both the soil and enrichment culture experiments. DET was  
344 found as the main metabolite in accordance with other studies (Navarro et al. 2003; Barra  
345 Caracciolo et al. 2005a; Delgado-Moreno and Peña 2007). The slight decrease in terbuthylazine  
346 concentration in the sterile conditions (both in soil and MB medium of the enrichment  
347 experiment) was presumably due chemical hydrolysis (Fig. 1A and Fig. 5A). DET and DEDT  
348 were not detected in sterile soil (data not shown) in line with the fact that their formation is  
349 reported to occur exclusively via biotic transformations (Di Corcia et al. 1999; Barra Caracciolo  
350 et al. 2005a; 2010).

351 The results of the microbiological analysis indicate that the presence of oak and pine  
352 amendments, rich in labile carbon fractions, stimulated soil dehydrogenase activity during the  
353 experimental period (Fig. 3). The positive influence of organic amendments on the  
354 dehydrogenase activity of the overall microbial community was found in several works  
355 (Moorman et al. 2001; Delgado-Moreno and Peña 2007; 2009) and in our previous experiment  
356 using the same pine and oak residues (Grenni et al. 2009a). Terbuthylazine did not negatively  
357 affect bacterial community functioning in terms of dehydrogenase activity (in the case of SOT  
358 the activity even increased), presumably because it was adapted to its presence (Fig. 3A).

359 The initial bacterial numbers were higher in all the amended soils than in the unamended  
360 ones and this was due to the fact that with pine and oak both organic matter and microorganisms  
361 were added to the soil (Briceño et al. 2007). However, since this difference was limited to the

362 start of the experiment, the allochthonous bacterial populations introduced by the residues were  
363 presumably not able to survive in the soil and they were both excluded competitively by the  
364 autochthonous populations and also affected negatively by the herbicide in the case of the treated  
365 soils. This hypothesis is confirmed by the cell viability values (% Live/Live+Dead) reported in  
366 Figure 4. In fact a transient decrease in viability was observed in all amended soils at day 6 and it  
367 was particularly evident in the SOT and SPT treatments.

368 The bacterial viability trend can be linked to the activation of bacterial populations  
369 involved in the herbicide degradation and this is particularly evident in the ST. In fact, in the  
370 control soil S (non-terbuthylazine treated soil) the overall cell viability tended to decrease during  
371 the experimental period. The  $t_{1/2}$  values of TBA were related to the  $K_d$  values found in the  
372 different treatments (Fig. 1A and Table 1); therefore the adsorption phenomena affected the  
373 amount of herbicide bioavailable for degrading populations.

374 The higher sorption of terbuthylazine by pine-amended than oak-amended soil is in line  
375 with its higher lignin content (24.4% in pine vs 18.2% in oak) and its greater organic carbon  
376 content (41.5% in pine vs 38.5% in oak). The  $K_d$  coefficients obtained in this work are higher  
377 than those found in a previous work (Rodríguez-Cruz et al. 2007b) for the sorption of linuron,  
378 alachlor and metalaxyl herbicides. The latter result is ascribable to the higher hydrophobicity of  
379 terbuthylazine ( $\log K_{ow} = 3.21$ ) compared to other non-ionic pesticides ( $\log K_{ow}$  range 1.75-  
380 3.09) (Rodríguez-Cruz et al. 2007b). The  $K_d$  value obtained in the SPT is comparable with that  
381 obtained by Cabrera et al. (2008) in a soil amended with alperujo. The initial decrease of TBA  
382 concentration in SPT can be explained by the degradation of the limited bioavailable fraction  
383 (20%) occurring in the amended soil; the pine sorption capability then presumably increased  
384 with the incubation time and no significant amount was further degraded during the experimental  
385 period. With the increase in incubation time ageing phenomena, which imply the formation of  
386 bound-residues or strong immobilization of pesticide residues in non-amended or amended soils



387 are commonly found (Gevao et al. 2000). The addition of organic amendments (urban sewage  
388 sludge, poultry compost and alperujo), increasing terbuthylazine sorption to soil, has been found  
389 to retard its degradation by other authors (Navarro et al. 2003; Cabrera et al. 2007; 2008;  
390 Dolaptsoglou et al. 2007; Sayara et al. 2010).

391 In the presence of oak amendment the herbicide sorption did not increase significantly  
392 and therefore did not substantially hamper the biodegradation, with the TBA decrease in  
393 concentration being just slightly delayed by about 12 days; however, at the end of the experiment  
394 the residual herbicide concentration was identical (60% of the initial concentration) to that in the  
395 unamended soil.

396 The enrichment culture experiment confirms the fact that the capability of the same  
397 herbicide degrading populations was hampered in different ways by adsorption phenomena. In  
398 fact, it demonstrates that the autochthonous soil microbial pool was able to degrade the  
399 terbuthylazine in the liquid culture ( $t_{1/2} = 24$  days, Figure 5A) significantly better than in its  
400 original soil ( $t_{1/2} = 105$  days in ST, Figure 1A) and to grow on the herbicide as the sole carbon  
401 source. This result not only confirms that terbuthylazine biodegradation can be carried out by  
402 bacterial consortia (de Souza et al. 1998; Grenni et al. 2009b) but it also demonstrates how the  
403 same microbial pool was prevented in the original clay loam soil from performing the  
404 degradation efficiently.

405 In fact in this soil, without any amendments, the herbicide terbuthylazine was quite  
406 persistent ( $t_{1/2}$  95-105 days). The TBA degradation rate is indeed reported highly variable, from a  
407 few weeks to more than 200 days (Di Corcia et al. 1999; Barra Caracciolo et al. 2005a;  
408 Kravvariti et al. 2010) and it depends on bacterial activity and on abiotic factors that directly or  
409 indirectly influence the degradation rate. Abiotic factors, such as temperature, soil moisture,  
410 organic carbon content and pH, have been known to significantly influence the degradation  
411 process of terbuthylazine; however, the soil intrinsic characteristics (such as texture and

412 mineralogy) have been taken into consideration only very recently (Vischetti et al. 2010). In this  
413 context, the relatively high half-life values (95-105 days) found in this clay loam soil could be  
414 ascribed to its fine texture and in particular to its clay mineral montmorillonite fraction which  
415 has a great capacity to adsorb organic matter (Arnarson 2000), including triazine herbicides,  
416 decreasing their bioavailability for degradation (Bailey et al. 1986). In fact in our previous  
417 microcosm studies, performed with terbuthylazine in the same laboratory conditions but on  
418 different soils (silty-loam and sandy-loam), we observed a  $t_{1/2}$  of 22 days and 30 days,  
419 respectively (Barra Caracciolo et al. 2001; 2005a).

420         The slow degradation rate found in the unamended soil was due firstly to the intrinsic  
421 characteristics of its clay, such as texture and mineralogy (e.g. the montmorillonite fraction),  
422 which made the herbicide less available for both abiotic and biotic degradation processes.

423         Moreover, although the soil studied had a fine texture, which is known to limit the  
424 diffusion and transport of contaminants, the groundwater beneath has been found to be  
425 contaminated by triazines. This fact can be ascribed to preferential flow pathways (Flury et al.  
426 1996; Kördel et al. 2008) occurring when large and discontinuous macropores operate and cause  
427 rapid movement of chemicals through the unsaturated zone. The transport of pesticides via  
428 macropores has been frequently found in fine textured soils and herbicides may contaminate  
429 groundwater especially if the degradation phenomena in surface soil do not significantly reduce  
430 their concentration (Guzzella et al. 2003, 2006), as in the case of our soil.

431         The evaluation of the environmental fate of terbuthylazine in clay soil requires the  
432 knowledge of how its persistence and movement can be affected by the possible interactions  
433 with its fine texture. Owing to the high adsorption of terbuthylazine to this soil, its leaching is  
434 not likely to occur through micropore flow, but through a preferential flow. The occurrence of  
435 terbuthylazine and DET in the groundwater beneath can therefore be explained by their transport  
436 by macropore flow.

437 Particular attention has therefore to be paid to clay soils because their structure can  
438 strongly affect both the transport and the degradation of soil contaminants and consequently  
439 makes it difficult to forecast their fate and control their mobility. The latter statement has to be  
440 taken in consideration for a better risk management of pesticide use in line with the requirements  
441 of the recent EU Directive 2009/128/EC and Regulation (EC) No. 1107/2009.

442

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450

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595 **Figure captions**

596 **Fig. 1. (A)** Percentage of residual terbuthylazine in unamended soils. (*SST*, sterile soil; *ST*, non-  
597 sterile soil) and soil amended with oak (*SOT*) or pine (*SPT*) vs time. *Vertical bars* represent  
598 standard errors. **(B)** Terbuthylazine structure (on the *left*) and images of oak and pine wood  
599 residues (on the *right*).

600  
601 **Fig. 2.** Metabolites detected over time in unamended (*ST*) and soil amended with oak (*SOT*) or  
602 pine (*SPT*) **(A)** Desethylterbuthylazine (DET); **(B)** Desethyldebutylterbuthylazine (DEDT). *Bars*  
603 represent standard errors.

604  
605 **Fig. 3.** Soil dehydrogenase activity ( $\mu\text{g TPF g}^{-1}$  dry soil) detected over time in the soils **(A)**  
606 treated with terbuthylazine (*SPT*, terbuthylazine + pine; *SOT*, terbuthylazine + oak; *ST*,  
607 terbuthylazine) and **(B)** in the control ones (*SP* amended with pine; *SO*, amended with oak; *S*,  
608 unamended). *Bars* represent standard errors.

609  
610 **Fig. 4.** Cell viability (% Live/Live+Dead) vs time **(A)** in the soils treated with terbuthylazine  
611 (*SPT*, terbuthylazine + pine; *SOT*, terbuthylazine + oak; *ST*, terbuthylazine) and **(B)** in the  
612 control ones (*SP* amended with pine; *SO*, amended with oak; *S*, unamended). *Bars* represent  
613 standard errors.

614  
615 **Fig. 5.** Terbuthylazine degradation, desethylterbuthylazine (DET) formation and total cell  
616 number in an enrichment culture. **(A)** Degradation (%) of terbuthylazine (TBA) vs time; **(B)**  
617 Formation of the metabolite desethylterbuthylazine (DET). *Bars* represent standard errors.

618

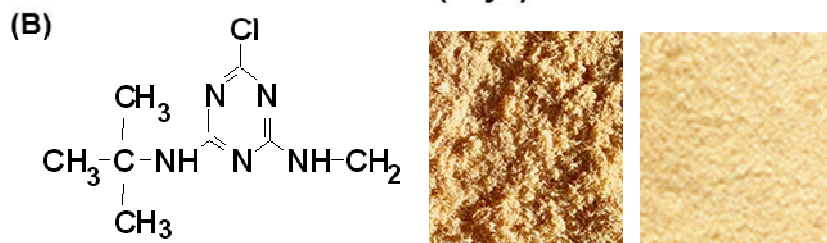
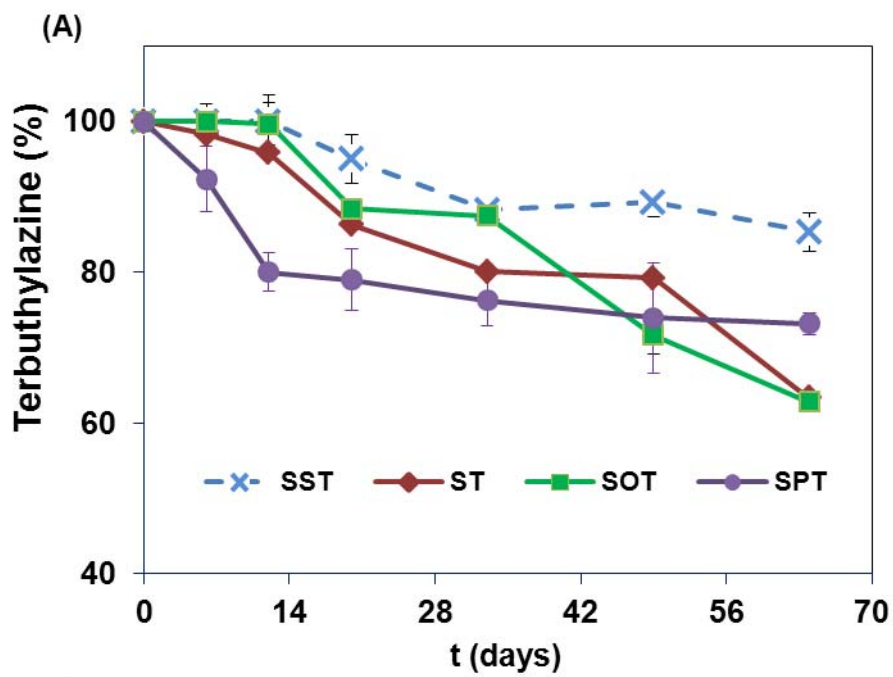


Fig. 1.

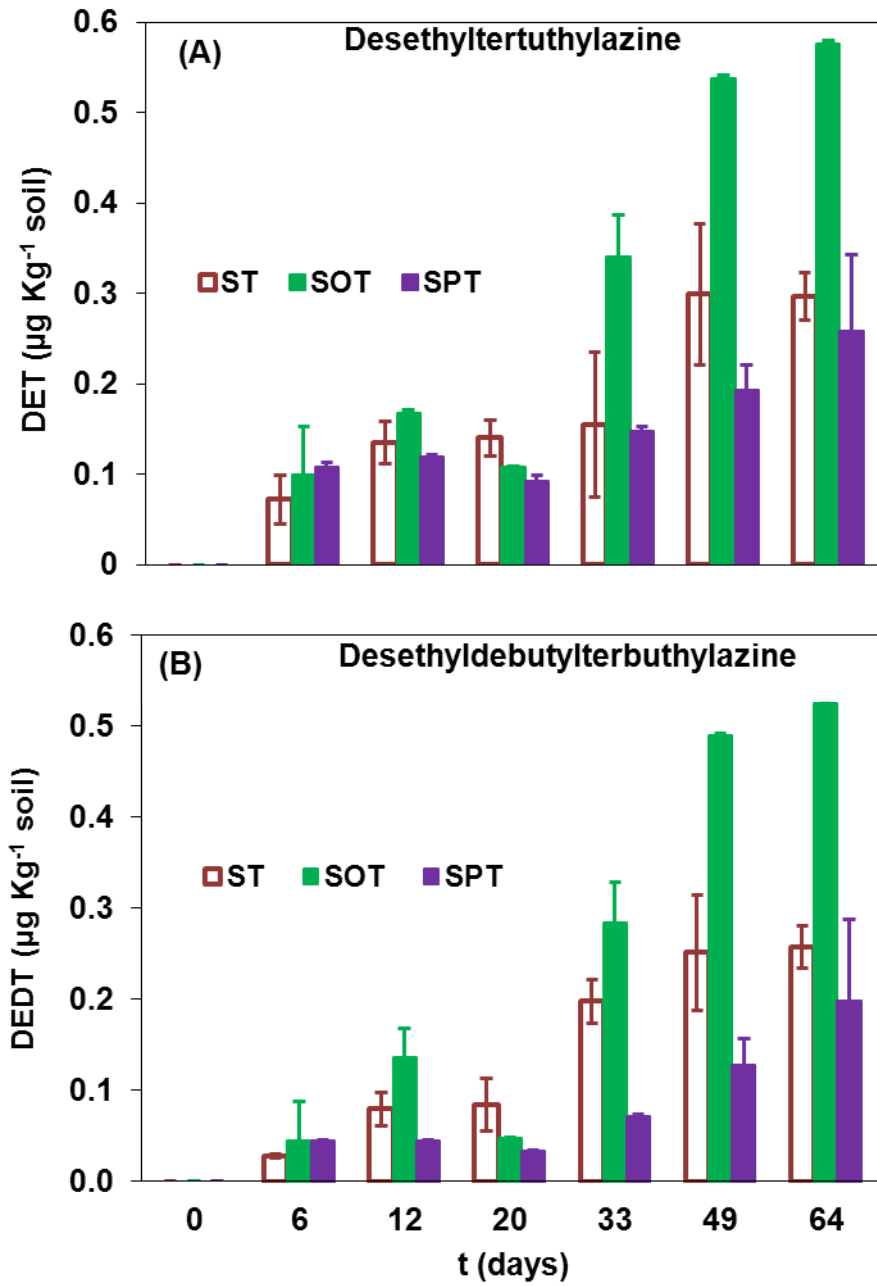


Fig. 2.

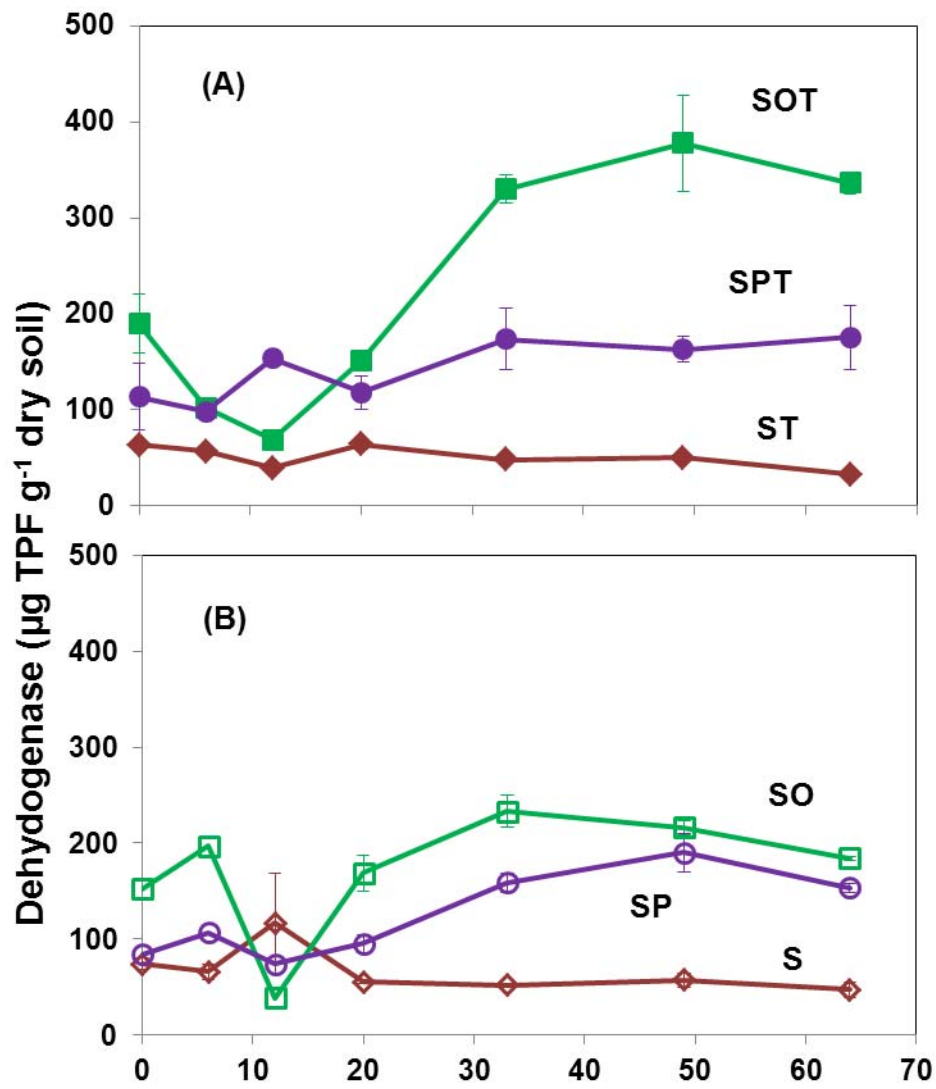


Fig. 3.

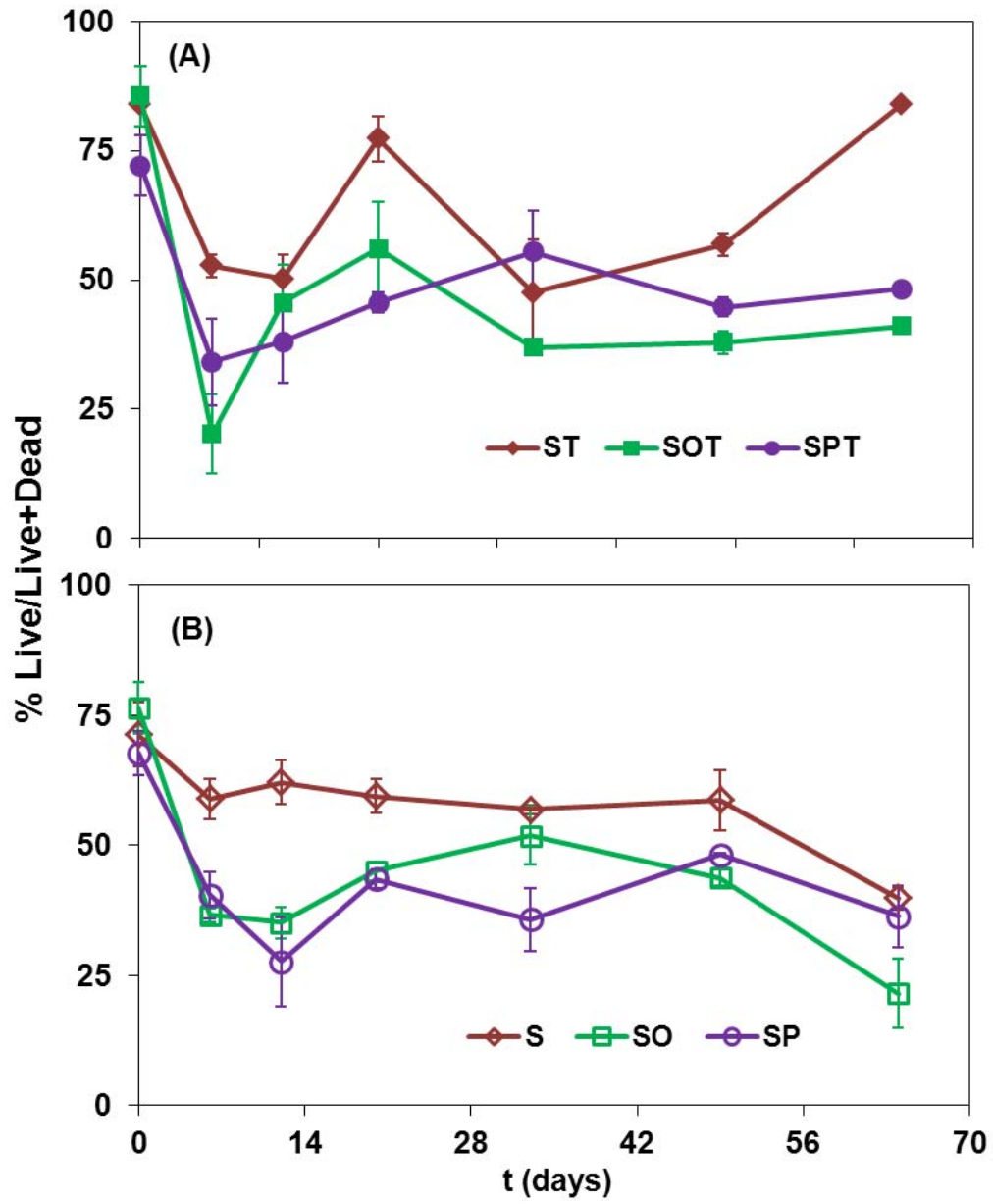


Fig. 4.

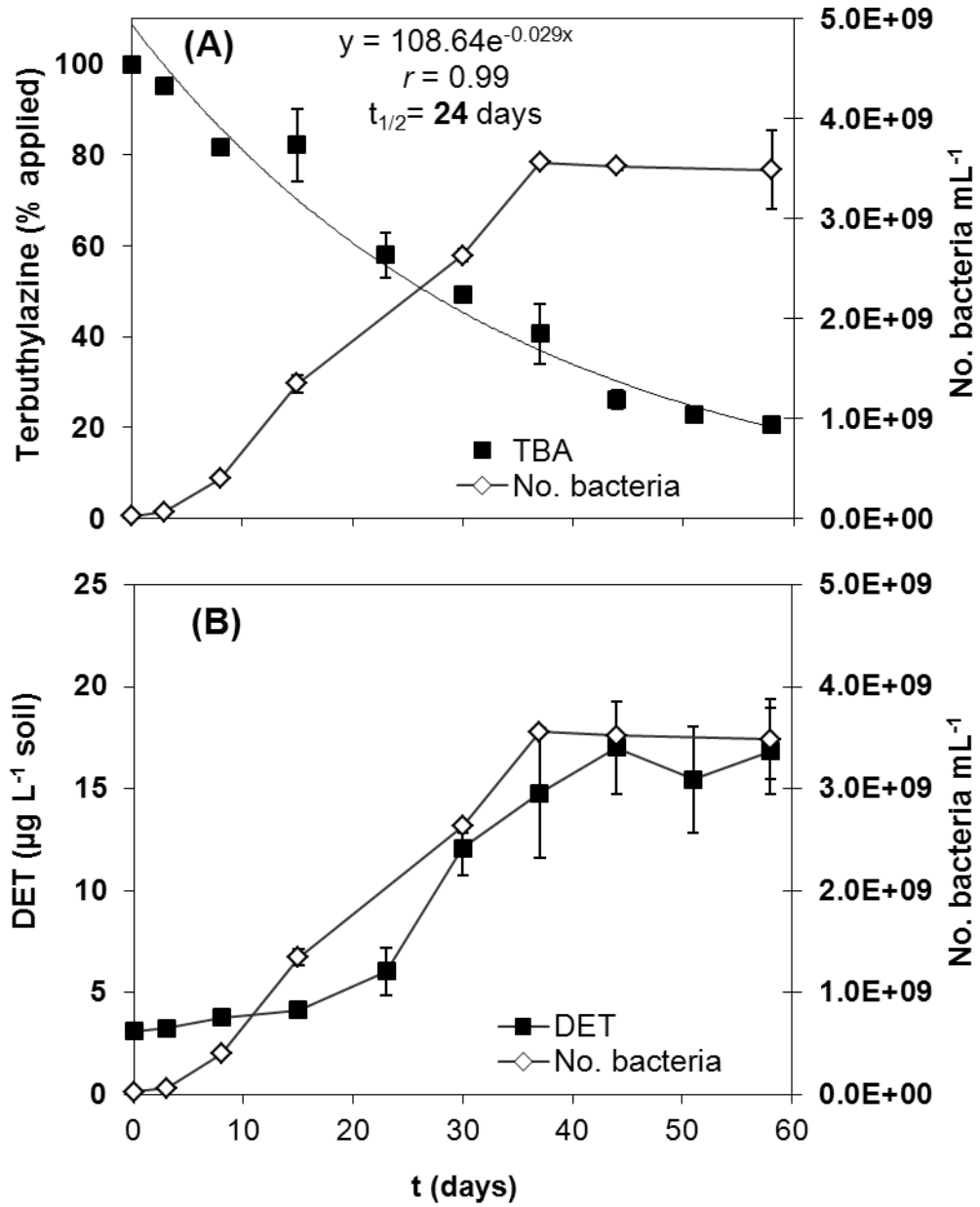


Fig. 5.

**Table 1** Organic carbon (OC %), pH and Sorption distribution coefficient ( $K_d$ ) of terbuthylazine by the wood residues (pine and oak), unamended (soil) and amended soils (soil + pine, soil + oak).

	OC %	pH	$K_d$ (mL g <sup>-1</sup> ) ± SD
Pine	41.5	5.0	1856 ±46.4
Oak	38.5	4.0	71.2 ±1.18
Soil	1.87	7.7	4.27 ±1.08
Soil + Pine	4.21	7.7	11.7 ±2.63
Soil + Oak	4.09	7.8	5.30 ±0.38

*SD* standard deviation