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Quantification of Aspergillus fumigatus and enteric bacteria in European compost and biochar.

This is a pre print version of the following article:			
Original Citation:			
Availability:			
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1 Quantification of Aspergillus fumigatus and enteric bacteria in European compost and

2 **biochar**

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Abstract

29 Although most potential human pathogens (PHPs) can be inactivated during composting, the risk 30 that such substrates represent for human health remains largely unknown due to the shortage of 31 information on presence and abundance of PHPs in finished composts. This study focused on the 32 assessment of Salmonella spp., Listeria monocytogenes, Shiga toxin-producing Escherichia coli 33 (STEC) and the opportunistic fungal pathogen Aspergillus fumigatus in different compost 34 commodities. A total of fifteen European composts, made from different waste types and processes, were evaluated for the occurrence of the selected PHPs using molecular and traditional techniques. 35 36 The analyses were extended to five biochar because of their growing application in agriculture, 37 horticulture, floriculture and private gardening. 38 Enteric bacteria were detected by molecular methods in eight out of fifteen composts; however, 39 viable propagules were confirmed for L. monocytogenes only in two composts, and for STEC in 40 further three composts. No bacterial pathogens were found in biochar. Living A. fumigatus was present in eleven composts and two biochars. None of the eighteen isolates contained SNPs relevant 41 42 for resistance to azole fungicides. The role of compost and biochar as a source of PHPs in the environment and the risk for human health is discussed. 43

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Introduction

47 Composting is a self-heating microbiologically driven process that allows the recycling of a variety 48 of organic materials with different origin (Ryckeboer *et al.* 2003; Stentiford and Bertoldi, 2010). 49 Municipal and green wastes or a mixture of those is common substrates but compost is also a mean 50 for recycling human sewage and animal wastes, although with increasing human hazard (Jones and 51 Martin, 2003). The process includes a mesophilic phase (10-42°C) at the beginning of the process, a thermophilic phase when temperature reaches 55-65°C for different extent of time, a further mesophilic stage and a final maturing phase when the temperature declines and the material stabilizes. Most of the potential human pathogens are eliminated during the thermophilic phase with 55°C for 3 days (Anon, 1981) but different time-temperature combinations are applied depending on country composting standards (Jones and Martin, 2003).

58 Composts in Europe are produced by following appropriate standard procedures and before 59 commercialization have to satisfy microbiological standards (e.g. levels of *E. coli* below 1000 CFU 60 per gram of fresh mass and absence of *Salmonella spp*.). Microbiological standards are also 61 recommended in the Draft Final Report on End-of-Waste for Compost and Digestate 62 (<u>http://ipts.jrc.ec.europa.eu/publications/pub.cfm?id=6869</u>), and expected in the future EU 63 harmonised compost quality regulation.

When composting processes are conducted in an inefficient manner, a substrate susceptible to recolonisation may be generated, and consequently compost could become a substrate maintaining a number of enteric bacteria in the environment such as *Salmonella*, *Escherichia*, and *Listeria*, posing human health issues (Jones and Martin, 2003).

68 To date, many studies have been carried out on the occurrence and survival of bacterial pathogens 69 in human and animal wastes and biosolids, and were extensively reviewed (Wiley and Westerberg 70 1969; Jones and Martin 2003; Sidhu and Toze 2009). However, despite the reported risk of plant 71 contamination by enteric pathogens when using composts (Islam et al., 2004), there is a shortage of 72 information on the presence and abundance of these organisms in green and mixed composts 73 (Avery et al., 2012). Indeed, microbiologist's attention is mainly focused towards the control and 74 inactivation of enteric pathogens to maintain their level under mandatory limits (Heringa et al. 75 2010; Shepherd et al. 2011; Singh et al. 2011).

Although the presence and abundance of the opportunistic fungal pathogen *Aspergillus fumigatus*Fresenius cause of the so-called "aspergilloses" of which the most severe is represented by the

78 Invasive Aspergillosis (IA), have been reported in different types of compost, it has been rather 79 underestimated even though its role in the compost degradation process and its health implications 80 are widely recognized. This organism is a ubiquitous fungus normally inhabiting the soil and 81 decaying materials (Dagenais et al. 2009; Gisi, 2013) but it is well equipped to survive successfully 82 in a wide range of environments due to a number of features, recently discussed by Kwon-Chung 83 and Sugui (2013), first of all the wide growth temperature range. A. fumigatus is present in compost samples at concentrations of 10^{6} - 10^{7} CFU/gdw (Millner *et al.* 1994), the spores are released to the 84 air during compost activities such as turning reaching concentrations of 10^4 - 10^7 /m³ (Recer *et al.* 85 86 2001; Wheeler et al. 2001).

According to O' Gorman (2011) the menace for IA coming from the airborne inoculum of *A*. *fumigatus*, also originating from compost commodities, is highly underestimated.

Organic substrates, including them compost, have been claimed to be one of the environmental
sources for itraconazole resistant strains of *A. fumigatus* further contributing to hazard for human
health (Snelders *et al.* 2009; Verweij *et al.* 2009; Gisi 2013).

92 Biochars are obtained from plant and/or animal wastes transformed to carboniferous porous 93 material by pyrolysis processes (Beesley *et al.* 2011). Although biochar is considered safe for users, 94 thanks to the high temperatures used for its production, possible contaminations by human 95 pathogens may occur in later stages. Therefore, a proper storage and avoidance of cross 96 contamination must be taken into consideration.

97 In this study, a combination of relevant microbiological and molecular techniques have been 98 adopted to assay the occurrence of targeted *Salmonella spp.*, *L. monocytogenes*, Shiga toxin-99 producing *E. coli* (STEC) and the opportunistic fungal pathogen *A. fumigatus*. The vitality and 100 concentration of enteric bacteria and *A. fumigatus* have been estimated by plating. Real time PCR 101 kits have been used as quick detection method of genomic and/or metagenomic DNA directly 102 extracted from compost or obtained after culture enrichment. While the presence of bacterial 103 pathogens have been confirmed by selective plating and real time PCR, the identification of *A*. *fumigatus* was completed by macro-morphological assessment and by sequencing relevant gene regions including ITS and β-tubulin (Samsom *et al.* 2007). The 14α-sterol demethylase gene *cyp51A* and the gene promoter were also sequenced in order to inspect whether strains resistant to demethylation inhibitor (DMI) fungicides were present (Diaz Guerra *et al.* 2003; Chen *et al.* 2005; Verweij *et al.* 2009; Howard *et al.* 2011). All testing were extended to five biochars to understand if and to what extent these new soil amendments may embody a further environmental reservoir of the targeted PHPs.

The aim of this study was to estimate the presence and abundance of targeted PHPs in organic substrates such as compost and biochars, made from different waste types and processes. Because there is a growing health concern linked to the increased recovery of *A. fumigatus* isolates resistant to azole fungicides in clinical and environmental samples (Gisi 2013; Vermeulen *et al.* 2013), a further objective of the study was to verify the potential contribution of compost and biochar as environmental source for DMI resistant *A. fumigatus* strains as was recently hypothesised by some medical researchers (Snelders *et al.* 2009; Verweij *et al.* 2009).

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Materials and methods

Composts and biochars, collection and storage

121 Fifteen composts originated from six different European countries (Hungary, The Netherlands, 122 Spain, Italy, United Kingdom and Portugal) have been analyzed. Three types of composting 123 systems were selected: open outdoor composting; closed (in-vessel, turning) composting; combined closed (first phase) and open (second phase) composting. Different types of waste were considered: 124 125 only green waste (garden and park waste, ERC 20 02 01); green waste and municipal waste (kitchen and canteen waste, ERC 20 01 08); animal manure (manure, ERC 020106 and sludge, ERC 126 127 020301); agrifood waste (Olive mill pomace, ERC 02 03 01; Olive leaves, ERC 02 03 04). Specific 128 features of each substrate examined are listed in Table 1.

In their country of origin representative samples were taken from an approximately 1 m^3 big bag of compost, consisting of a composite sample obtained by pooling 5-6 individual subsamples taken from a 20-30 m³ pile of siewed and ready to market compost.

The five biochars came from four different countries and, except for BIOCHAR 1 which was obtained by the carbonization of animal bone, all others derived from plant wastes. Upon their arrival (April-May) composts/chars were maintained in big bags stored under an outdoor canopy and subsamples of 1kg (approx weight) were kept at 4°C for the following microbiological and molecular analyses.

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Detection and identification of enteric pathogens

139 Compost and biochar samples (25 grams) were collected in sterile Blender bags (VWR, Radnor, 140 PA, USA). L. monocytogenes, Salmonella spp., Shiga toxin-producing E. coli (STEC) were 141 detected by using real-time PCR kits (iQ-CheckTM,Bio-Rad, France) following manufacturers' 142 instructions with only minor modifications. In brief, for each pathogen, selective enrichment broth 143 at the appropriate incubation temperature, and DNA extraction protocol for environmental samples, 144 were followed. Simultaneously, 0.1 mL of samples were plated on Hektoen-Entero-Agar (Merck[®], 145 Germany), and coliform bacterial colonies (typically orange-red surrounded by a zone of 146 precipitate) were enumerated after incubation at 35 °C for 24 h.

147 Selective platings were carried out for all samples to verify the presence of the targeted pathogens. 148 The procedures were as follows: 25 g of each sample was transferred from the container into a 149 sterile Blender bag (VWR, Radnor, PA, USA) together with 225 mL of 0.1% sterile peptone water 150 (Sigma-Aldrich, St.Louis, USA) and homogenized for 180 s in a Masticator (IUL instruments, 151 Barcelona, Spain). To enumerate L. monocytogenes, Oxford selective agar (Sigma-Aldirich, St. 152 Louis, USA) added with Oxford *Listeria* selective supplement (Sigma-Aldirich, St. Louis, USA) 153 were used. Two 100 µL homogenates were taken, and decimal dilution series were spread onto two 154 Oxford agar plates and incubated for 48 h at 35 °C. Similarly, homogenates of STEC positive 155 samples were seeded in duplicate onto CHROMagarTM STEC (CHROMagar, Paris, France) plates 156 were incubated at 37 °C for 24 hours. *Salmonella* spp. positive samples were checked by using 157 Xylose-Lysine-Desoxycholate Agar (XLD) plates after 24 h of incubation at 35 °C. For each 158 selective media, typical colonies appearance were counted according to the user's manual, 159 randomly selected (at least three colonies), and picked for further confirmation by using appropriate 160 real-time PCR kit as above.

All bacterial concentrations have been expressed as Colony Forming Units per gram dry weight of
substrate (CFU/gdw) for each compost/biochar.

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Isolation and identification of Aspergillus fumigatus

Three subsamples of 0.5g (fresh weight) 10fold diluted in Ringer (Merck[®], Germany) solution (2 165 166 tablets per litre), the flasks were incubated in a rotary shaker at 100 rpm for one hour and the broth 167 diluted ten-fold. One ml of suspension was spread in three 90 mm Petri dishes for each dilution 168 series containing 15 ml of Potato Dextrose Agar (PDA) amended with 60mg/l of streptomycin, and 169 plates sealed and incubated at 42°C (species optimum). After four days plates have been assessed 170 for the presence of typical dark green/grey colonies exhibiting A. fumigatus morphology. At least one isolate was retained per each compost/biochar sample and purified through mono-hyphal 171 172 subculturing at least twice on PDA at 37°C. The first identification was carried out through 173 comparing the ability to growth at 10 and 50°C and assessing the colony and fungal structures 174 morphology with the reference CBS isolate 133.61 (Samson et al. 2007; Arendrup et al. 2010).

175 Cultures for DNA isolation were grown on cellophane disks for 7 days at 37°C, and mycelia 176 collected in 1.5 ml tubes. DNA of *A. fumigatus*-like colonies has been obtained using the EZNA[®] 177 FUNGAL DNA KIT starting from 100mg of fresh mycelia following manufactured instructions with the 178 exception that DNA was finally eluted in MilliQ autoclaved water instead of using kit elution 179 buffer. Isolate identification was confirmed by sequencing the rDNA ITS (White *et al.* 1990) and β -180 tubulin (Glass and Donaldson, 1995). Each single reaction mixture contained: 1µl of template DNA, 2µl of nucleotides mixture 2.5mM, 1µl of each primer prepared at a concentration of 10mM,
1.25µl of MgCl₂ 50mM, 2.5µl of 10XPCR buffer, 16 µl of MilliQ autoclaved water and 0.25 µl of
Taq Polymerase (Qiagen[®], Germany). PCR cycle for ITS and *ben*A gene included a denaturing
stage of 95°C for 2 min and 35 cycles as follow: 94°C for 30 sec, 55°C for 30 sec, 72°C for 1 min
with a final elongation step at 72°C for 7 min.

186 To determine the presence of specific SNPs for DMI resistance among the pathogen isolates, the cyp51A gene was amplified using high fidelity Taq and specific primer pairs (Chen et al. 2005). 187 188 Each reaction was composed of the following components: 1µl of template DNA, 2µl of nucleotide 189 mixture 2.5mM, 1µl of each primer prepared at a concentration of 10mM, 1.25µl of MgCl₂ 50mM, 190 2.5µl of 10XPCR buffer, 17.4 µl of MilliQ autoclaved water and 0.1 µl of Pfu Taq 191 (Invitrogen[®]Carlsbad, CA USA). For *cyp51A* gene identification, the cycles were: a denaturing stage at 94°C for 2 min followed by 94°C for 30 sec, 56°C for 30 sec, 68°C for 2 min with 35 192 193 cycles, without final elongation. The *cyp*51A promoter was amplified using the protocol reported by 194 Mellado et al. (2001). The promoter PCR included a denaturation step at 94°C for two minutes 195 followed by 35 cycles at 94°C for 30s, 60°C for 30sec, 68°C for 2min.

196 ITS, β -tubulin and *cyp51A* amplicons were sequenced through Macrogen Europe sequencing 197 service, and sequences manually edited using BioEdit v. 7.9 (Hall 1999). The coding region was 198 compared with the *cyp51A* sequences present in the database to verify the existence of reported 199 mutations linked to DMI resistance. Confirmed *A. fumigatus* isolates were included in 200 AGROINNOVA culture collection and long-term stored in Tryptic Soy Broth (Merck[®], Germany) 201 25% glycerol at -20°C. All potentially hazardous materials were destroyed through autoclaving for 202 25min at 121°C.

203 The concentration of *A. fumigatus* was expressed as Colony Forming Units per gram of dry weight
204 (CFU/gdw) for each compost/biochar.

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DNA isolation and real time PCR for A. fumigatus direct detection

207 Metagenomic DNA extraction was made with samples of 0.5 g of compost/char using NUCLEO SPIN 208 SOIL KIT (Macherey-Nagel GmbH & Co. KG) following manufacturers' instructions, with a final elution step in MilliO autoclaved water as for fungal DNA. Genesig[®] commercial kit adopted in 209 medical labs for the specific detection and quantification of A. fumigatus in clinical samples was 210 purchased by Primer DesignTM Ltd, UK. The kit uses a taq-Man probe developed for a gene of the 211 212 hypothetical protein AFUA 3G08890 to detect in 50 PCR cycles the presence/absence of A. fumigatus in clinical samples with a cut-off Ct value of 39. Following manufacturers' instruction 213 214 the kit was first used on 1:20 diluted DNA obtained from mycelia and then applied to metagenomic 215 DNA extracted from compost and biochars.

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Results and discussion

Compost and biochar characteristics

219 The main characteristics including percent organic carbon, C/N ratio, pH, provenances, input 220 materials and processes of 15 compost types and 5 chars screened in the study are summarized in 221 Table 1. Six composts each came from green waste (C2, C6, C7, C8, C11, C13) and municipal 222 waste (C3, C4, C9, C12, C14, C15), two from animal waste (C5, C10) and one from a combination 223 of olive pomace and sheep manure (C1). One char was animal based (CHAR1), the others plant 224 based. Most compost types were neutral (pH 6.5 to 7.5), some (C3, C6, C9, C10, C15) were slightly 225 basic (pH 7.5 to 8.1) and most chars strongly basic (pH > 9). Only a few compost types were rather 226 low in organic carbon (< 20%, C2, C5), all others ranged between 20 and 43%. The degradability of organic substrates can be estimated by the C/N ratio; if >25, degradation is assumed to slow down. 227 228 All compost types showed rather favourable C/N ratios (10 to 20), some (C8, C9, C13, C15) even 229 below 10 indicating the presence of rather high nitrogen concentrations. No relation was found 230 between chemical properties and provenances of compost types.

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Presence and abundance of enteric pathogens

233 Through real time PCR assays eighteen positive results were obtained for enteric bacteria: C6, C11, 234 C14 and CHAR3 for L. monocytogenes; C1, C2, C4, C5, C15, CHAR2 for Salmonella spp. and C1, 235 C2, C5, C10, C13, C14, C15, CHAR3 for STEC. Nevertheless, vital colonies were observed in only five cases out of twenty samples (Table 2): L. monocytogenes was found in green compost in one 236 Spanish (C6) and one Italian (C11) sample with a concentration of 2.3×10^3 and 2.8×10^4 237 238 respectively. Shiga toxin-producing E. coli were detected in three composts made from municipal 239 and animal manure from Italy, Spain and Hungary (C10, C12, C13) with concentrations of 1.88 to 2.46×10^3 CFU/gdw. 240

Coliform bacteria were detected in nine of twenty samples between a minimum of 4.1×10^2 241 CFU/gdw in two Italian composts of municipal origin and a maximum of 1.94×10^5 CFU/gdw in C2 242 243 green compost from Netherland. Values are within the range of those already reported in other 244 studies on composts (Gong et al. 2005). Plate counts for Salmonella spp. were always negative (<10 245 CFU/gdw). In biochars targeted bacteria were not found (<10 CFU/gdw) (Table 2).

246 Although selective platings allowed to detect vital enteric bacteria in finished composts the 247 overestimation of their abundance linked to the aerosol dispersion should be also contemplated 248 according to what reported by Cevallos-Cevallos et al. (2012).

249 The detection of enteric bacteria in animal derived composts was expected as their presence and 250 survival are well known in biosolids (Sidhu and Toze, 2009) which represent similar matrices in 251 terms of input material for animal waste compost. The finding of enteric bacteria in green composts 252 is also not uncommon (Jones and Martin, 2003) although few data are available on their level and it is still unclear which factor(s) correlate greatly with their presence and survival (Avery et al. 2012). 253 254 According to our results the detection of enteric pathogens in composts seem to be more linked to 255 handling, transport and/or external contamination (outdoor storage) rather than being a feature of 256

257 Detected concentrations are unlikely to cause contamination of vegetables growing on substrates containing PHP's; however, considering that only few cells could cause illness they should not be 258

the material or process itself (Pietronave et al. 2004).

underestimated. Favourable conditions, in terms of humidity, temperature and lack of antagonistic
competitors could allow enteric bacteria to re-grow and re-colonise the substrates (Santamaria and
Torazos, 2003; Sidhu and Toze, 2009).

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Presence and abundance of A. fumigatus

Viable propagules of *A. fumigatus* were found in eleven out of fifteen composts and in two biochars out of five (Table 2). Interestingly, the fungus was not found (or below detection limit) in composts made from animal manure (C5 and C10, both from Spain) or a combination of olive debris and sheep manure (C1 from Spain) as well as in one compost from municipal waste (C15 from Portugal) and three of five chars.

The highest concentration was detected in a British compost made from municipal waste (C14) with 268 6.15×10⁵ CFU/gdw. In all compost samples made from green wastes (C2, C6, C7, C8, C11, C13) 269 and one from municipal waste (C4), the A. fumigatus load was intermediate: between 0.24 and 3.62 270 $\times 10^3$ CFU/gdw and comparable to those of biochar 5 (2.32 $\times 10^3$ CFU/gdw). Low concentrations 271 were recorded in two composts from municipal waste (C3 and C9) and char 4, with estimated 272 concentrations lower than 10^2 CFU/gdw. In one case, A. fumigatus concentrations increased as a 273 274 consequence of longer storage and turning of the compost: in fact, the C12 sample represents the 275 same substrate as C3 but outdoor stored and periodically turned for a period of one year. Concentrations lower than 10^2 CFU/gdw were obtained with the same methodology also for a soil 276 277 sample used as standard substrate in greenhouse experiments (data not shown). The A. fumigatus 278 concentrations observed in our study are within the range of those already reported in other 279 investigations on composts (Millner et al. 1994; Anastasi et al. 2005).

The highest concentrations of the fungus were detected in compost made with green wastes, suggesting a close link between the presence of cellulose and lignocellulose substrates and *A. fumigatus* abundance. In fact, *A. fumigatus* is a strong producer of cellulolytic enzymes (Liu *et al.* 2013). While the link between the presence of the fungus with the type (mainly green) of compost appear evident, the presence of *A. fumigatus* in biochar is most probably due to superficial proliferation of the fungus likely as consequence of an airborne contamination.

287 On the other hand according to our results it does not seem to exist a connection between the 288 fungus level and the composting process because *A. fumigatus* was recovered from all types of 289 production methods.

Although real time PCR assays allowed confirming the identity of *A. fumigatus*, it was not possible to detect its presence based on DNA extracted directly from compost, probably because of reaction inhibitors and/or too low concentration of the fungus in the substrates.

Some protocols are available for the detection of *A. fumigatus* in water, air and clinical samples (McDevitt *et al.* 2004; Bansod *et al.* 2008; Vesper *et al.* 2008; Serrano *et al.* 2011) but no studies were carried out on its direct diagnosis in soils or composts. In agreement with O' Gorman (2011) and Gisi (2013) further studies are needed to investigate the presence and abundance of *A. fumigatus* in such types of environmental samples.

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Identification and characterisation of A. fumigatus *strains*.

300 Eighteen fungal strains were retained from the isolations made on PDA at 42°C. Sequence analysis 301 confirmed full identity of all strains to the species A. *fumigatus*. Taq Man real time PCR further 302 confirmed their identity. Real time PCR assays failed to detect A. fumigatus presence in compost 303 samples when DNA was extracted directly from the organic substrates, despite several attempts of 304 sample dilutions up to 500 fold, probably because some humic acid compounds disturbed the PCR 305 reaction. DNA sequences of isolates were deposited in GenBank for the three regions assessed 306 (accessions KF921462-KF921475; KJ584392-95 for ITS; KF921476-KF921489; KJ584396-99 for 307 beta-tubulin; and KJ584374-90 for *cyp51A*).

308 All sequences shared 100% identity with *A. fumigatus* gene accessions present in the NCBI 309 database. However, there were some minor differences among isolates in the ITS and in the beta310 tubulin sequences: in four isolates, a T to C change at position 105 of the ITS sequence, and in one 311 other isolate a G to A change at position 203 of the beta-tubulin was found. Based on the analyses 312 of cyp51A gene, none of the eighteen isolates obtained from the compost samples carried any of the 313 known mutations for DMI resistance. However, isolate A11 showed several polymorphisms, but 314 only one translated in an amino acid change (E427K) (Table 3) which was reported previously in 315 either resistant or susceptible A. fumigatus isolates (Howard et al., 2011). In addition, isolates A56 and A57 showed an amino acid change (N>K) at the 248 position (Table 3), but this mutation has 316 317 not been reported among the ones linked to DMI resistance. Furthermore, none of the isolates had a 318 tandem repeat of 34 bp in the gene promoter (S1).

None of the *A. fumigatus* isolates from compost examined in this study contained relevant mutations in the *cyp51A* gene, encoding DMI resistance. However, it cannot be ruled out that azole resistant isolates may be detected in environmental samples especially when a larger study is undertaken including more compost types and other habitats where *A. fumigatus* can grow and sporulate (Gisi 2013). Two isolates (A56 and A57) showed an unknown mutation at the 248 position of the *cyp51A* protein sequence; whether or not this mutation may induce a reduced sensitivity to DMI fungicides is currently under investigation.

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Conclusions

Data on recovery and quantification of PHPs in green and mixed composts are either fairly limited or outdated (Millner *et al.* 1977; Clark *et al.* 1983; Gong *et al.* 2005; De Clercq *et al.* 2007) and missing for biochars, even though extended literature is available on the study of microbial communities in composts and during their production process with different experimental and technical approaches (Ryckeboer *et al.* 2003; Insham *et al.* 2003; Anastasi *et al.* 2005; Danon *et al.* 2008; Bonito *et al.* 2010; Neher *et al.* 2013). To our knowledge this work represents the first study on the detection and quantification of four of the main PHPs in a reasonable wide number of compost samples and it is definitely the first considering biochars. This combined approach was
adopted to have a broader, even if specific, view of PHPs inhabiting finished organic products.

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Results of the analyses confirm the variable presence in compost of some enteric bacteria, but mainly the consistent presence of *A. fumigatus*. This agrees with what is generally expected because most bacterial pathogens are inactivated by composting while *A. fumigatus* is known to play an active role in the process (Jones and Martin, 2003; O'Gorman 2011).

Among detected living PHPs, the presence of *L. monocytogenes* and Shiga toxin-producing *E.coli* in compost, which could lead to crop contamination when contaminated compost is used in agriculture, together with the abundance of the opportunistic fungus *A. fumigatus* in these organic substrates may represent an health issue. It remains uncertain whether environmental exposure to enteric pathogens by handling contaminated composts would present a tangible risk for humans mainly through plant contamination.

To limit the health risk imposed by the potential presence of these pathogens, good agricultural practices and proper handling of the substrates respecting strict hygienic rules by workers may be good enough.

351 The detection of living A. fumigatus in variable concentrations in the majority of samples confirms compost as being one of the major sources for this organism in the environment (O' Gorman 2011). 352 353 Furthermore, the pathogen was detected for the first time in biochars posing questions on how and 354 where these product types should be used to limit the hazard of unintentional transmission of fungal spores. In this study, we provide evidence that longer storage and turning of compost can increase 355 356 the concentration of A. fumigatus in the substrate. These findings highlight the need to widen future 357 studies to the dissemination of this organism within and outside compost facilities, in order to 358 identify the conditions favouring its dispersal and sporulation and to detect critical hazard points 359 during the process.

360 It is still an open question, whether *A. fumigatus* DMI resistant originates from medical treatments 361 (human and veterinary) spreading to the environment or vice versa. It can be assumed that 362 resistance emerges in all situations where azole (DMI) fungicides are used intensively at high 363 concentrations over a long period of time (Gisi 2013).

364 The use of compost and biochars in agriculture, horticulture, floriculture as well as for other 365 environmental applications is gaining more and more attention for a number of reasons. The major 366 value of such substrates are linked to their environmentally friendly features such as their effects for 367 long term crop plant fertilization and quality, favourable influence on soil structure, aeration, and 368 moisture, for suppressiveness of soil borne plant pathogens, for activation of nutrient cycling, 369 mineralization and bioremediation processes in the soil environment (Ahmad et al. 2007; Beesley et 370 al. 2010; Beesley et al. 2011). In addition, they can contribute to carbon sequestration (biochars). 371 Our results are of relevance for the ongoing discussion on regulatory aspect of these and similar 372 types of organic substrates for limiting the level of PHPs to reasonable levels in order to minimize 373 health hazard.

374 Further studies should be done for PHPs and A. fumigatus, in compost and biochar facilities as well 375 as in other relevant habitats of these organisms in order to understand their main environmental 376 sources. The presence and abundance of such organisms in commercial organic substrates 377 especially within horticultural and floricultural sectors should be considered in future studies along 378 with the contamination risk of vegetables by enteric bacteria. The development of reliable 379 molecular methods for the specific detection and quantification of living A. fumigatus inoculum in 380 soil, compost, biochar and similar substrates would be equally important. In this way, the origin and 381 migration of PHP's and A. fumigatus between different ecological niches (habitats) in the medical 382 (human and veterinary) and environmental (including agricultural, horticultural, floricultural) 383 sectors can and should be investigated in a more rational approach.

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Summary

Fifteen compost and five biochar were sampled in different European Countries. Enteric bacteria were detected by molecular methods in eight out of fifteen composts; however, viable propagules were confirmed for *L. monocytogenes* only in two composts, and for STEC in further three composts. No bacterial pathogens were found in biochar. Living *A. fumigatus* was present in eleven composts and two biochars. None of the eighteen isolates contained SNPs relevant for resistance to azole fungicides. The role of compost and biochar as a source of PHPs in the environment and the risk for human health is discussed.

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Acknowledgments

The research leading to these results has received funding from the European Union Seventh Framework Programme (FP7/2007- 2013) under grant agreement n° 289785 (Collaborative project REFERTIL: Reducing mineral fertilisers and chemicals use in agriculture by recycling treated organic waste as compost and biochar products).

- 400 All experimental materials were handled under a class 2 laboratory hood.
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