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1	Metabolic synergies in the biotransformation of organic and metallic toxic compounds by a
2	saprotrophic soil fungus
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21	Running head: Fungal biotransformation of hexachlorocyclohexane and vanadium
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23 Abstract

24	The saprotrophic fungus Penicillium griseofulvum was chosen as model organism to study responses to a
25	mixture of hexachlorocyclohexane (HCH) isomers (α -HCH, β -HCH, γ -HCH, δ -HCH) and of potentially
26	toxic metals (vanadium, lead) in solid and liquid media. The P. griseofulvum FBL 500 strain was isolated
27	from polluted soil containing high concentrations of HCH isomers and potentially toxic elements (Pb, V).
28	Experiments were performed in order to analyse the tolerance/resistance of this fungus to xenobiotics, and
29	to shed further light on fungal potential in inorganic and organic biotransformations. The aim was to
30	examine the ecological and bioremedial potential of this fungus verifying the presence of mechanisms that
31	allow it to transform HCH isomers and metals under different, extreme, test conditions. To our knowledge,
32	this work is the first to provide evidence on the biotransformation of HCH mixtures, in combination with
33	toxic metals, by a saprotrophic non-white-rot fungus and on the metabolic synergies involved.
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36	Keywords: soil saprotrophic fungi, hexachlorocyclohexane, vanadium, biotransformation, metabolic
37	phenotype, medium pH
20	

39 Introduction

40 Hazardous persistent organic pollutants, e.g. pesticides, pharmaceuticals, explosives, and potentially toxic 41 elements (PTEs), e.g. Pb, Cr, As, Sn, constantly enter ecosystems (waters, soils, and sediments) causing 42 severe environmental and health problems (Polti et al. 2014; Wan et al. 2015). Anthropogenic processes 43 associated with domestic, municipal, agricultural, industrial, and military activities represent the major 44 sources while PTEs can also be released from natural geological processes such as weathering and volcanic 45 eruption (Vargas-García et al. 2012; Griffith et al. 2015). Pollution due to human activities is often the 46 result of extensive histories of multiple land use which creates sites containing mixed pollutants. More than 47 40% of the United States National Priority List sites are co-contaminated by organic (volatile and semi-48 volatile organic compounds) and inorganic pollutants (metals including radionuclides) while metals and 49 mineral oil contribute jointly to around 60% of soil contamination and 53% of groundwater contamination 50 in Europe (Sandrin and Maier 2003; Panagos et al. 2013). The co-occurrence of organic and metal 51 pollutants is not only a threat to human and ecosystem health, but is also a challenge because the 52 technologies required for remediation of polluted sites are different for each group of pollutants (Sandrin 53 and Maier 2003; Zhu et al. 2012). As more than one third of contaminated sites are polluted by more than 54 one type of contaminant, it is imperative to develop cost-effective and sustainable techniques that can 55 transform organic compounds while also extracting PTEs or stabilizing them in non-toxic forms (Polti et al. 56 2014).

57 In recent years, several studies on microbial communities or on single microbes isolated from 58 historically contaminated sites have shown their ability to tolerate, adapt and grow in the presence of 59 organic compounds and PTEs (Alisi et al. 2009; Wasi et al. 2011). This suggests that bioremediation based 60 on microbial activities is feasible for the recovery of such sites by transformation or immobilization of both 61 organic compounds and PTEs (Zhu et al. 2012; Polti et al. 2014). Previous studies on microbial 62 biotransformations of multiple contaminants have mainly concentrated on bacteria (Alisi et al. 2009; Wasi 63 et al. 2011), but fungi can also represent ideal candidates for future challenges in complex multi-64 contaminated contexts. Fungi are ubiquitous chemoorganotrophic organisms, playing fundamental roles in 65 ecological and geological processes (Gadd 2010; Gadd et al. 2012). As decomposers, pathogens, and 66 symbionts (mycorrhizas, lichens), fungi provide fundamental ecological functions for ecosystems and 67 human well-being (Mace et al. 2012; Lange et al. 2012). Fungi can transform a huge variety of organic 68 substrates, including natural polymers such as cellulose, lignin, chitin and starch but also many 69 anthropogenic products like pesticides, explosives and other xenobiotics (Gadd 2013; Harms et al. 2011). 70 Due to their filamentous growth habit and ability to exude organic acids, protons and other metabolites, 71 fungi are important biological weathering agents of rocks and mineral-based substrates (Gadd 2004; 2007). 72 The potential of fungi to tolerate and transform both organic and inorganic pollutants has been highlighted 73 in many studies which have also reported some unusual abilities shown by fungi isolated from 74 contaminated soil (; Tigini et al. 2009; Ma et al. 2014; Mishra and Malik 2014). In fact, isolation of 75 indigenous fungi could provide the best candidate organisms for bioremediation of polluted soil since they already belong to an established soil microbial community, and are best adapted to the site conditions(Czaplicki et al. 2016).

78 In this research, we have examined tolerance to both organic and inorganic pollutants by a soil 79 saprotrophic fungus, Penicillium griseofulvum Dierckx isolated from a historically polluted soil. The study 80 area was the Italian National Site of Interest "Valle del Sacco" (Lazio, Italy) which is affected by high 81 concentrations of multiple pollutants such as hexachlorocyclohexane, vanadium and lead (Ceci et al. 2012; 82 Bernardini et al. 2016; Bernini et al. 2016). The aims of this research were to evaluate the 83 tolerance/resistance of P. griseofulvum FBL 500 to different combinations of vanadium, lead, and isomers 84 of HCH; to study the inorganic biotransformation in the presence of metals, and possible biomineralization 85 phenomena; and to analyse biodegradation of an isomeric mixture of HCH, in the absence and in the 86 presence of vanadium, and the impact of any synergistic effects on fungal metabolism.

87 Materials and methods

88 Organism, media, and growth conditions

A strain of *P. griseofulvum* Dierckx (FBL 500), obtained from the culture collection of the Fungal
Biodiversity Laboratory (FBL) (Sapienza, University of Rome), was used in all the biotransformation tests.
Czapek-Dox medium was used for all the experiments (; Ceci et al. 2015b, c). The strain is also preserved
in the public mycological collection of Mycotheca Universitatis Taurinensis (MUT) as MUT 5854. The
Czapek-Dox agar medium contained the following (g/l distilled water): NaNO₃, 3; K₂HPO₄ 1;
MgSO₄·7H₂O, 0.5; KCl, 0.5; FeSO₄·7H₂O, 0.01. All chemicals were purchased from Merck (Darmstadt,

95 Germany).

96 The experimental design was composed of three different experiments, in which (A) 97 tolerance/resistance responses and biotransformation of metals (V, Pb), (B) of HCH mixture of isomers, 98 and (C) of the synergic effects of combination of V and HCH isomers were respectively investigated (see 99 Table 1, Online Resource). The whole phenotypic response of this fungus to the different treatments was 100 examined using the Phenotype MicroArrayTM system (Pinzari et al. 2016). Sucrose and D-glucose (Difco, 101 Sparks, MD, USA) were used as substrates for fungal growth (Table 1, Online Resource). The glucose 102 concentration was 30 g/l in experiment A, in which P. griseofulvum FBL 500 was tested with metals (V and 103 Pb compounds): this concentration was used following the same cultural conditions as in Ceci et al. (2015a, 104 c) (Table 1, Online Resource). 5 g/l sucrose was used in experiment B to stimulate the biotransformation of 105 HCH isomers as used in previous tests of biotransformation of β-HCH by *P. griseofulvum* FBL 500 (Ceci et 106 al. 2015b) (Table 1, Online Resource). The same concentration was maintained in experiment C to study 107 synergic interactions between the fungus and different combinations of V and HCH. In experiment A, prior 108 to autoclaving, the medium pH was adjusted to 5.5 using concentrated HCl (Ceci et al. 2015a, c), while in 109 experiments B and C, the pH was kept at 7 in order to prevent acidic variations of the medium pH, which 110 could result in toxicity to P. griseofulvum FBL 500 during HCH biotransformation because of benzoate 111 formation (Guillén-Jiménez et al. 2012; Ceci et al. 2015b) (Table 1, Online Resource).

112 In experiments A and C, prior to inoculation, 84 mm diameter discs of sterile cellophane 113 membrane (Focus Packaging and Design Ltd, Louth, UK) sterilized by autoclaving in distilled water and 114 were placed aseptically on the surface of the agar in each Petri dish (Ceci et al. 2015c). Growth of P. 115 griseofulvum FBL 500 was evaluated by measuring diametric extension of the colony and by biomass yield 116 since extension of the colony alone does not take into account the density of fungal mycelium (Ceci et al. 117 2015c). After 12 days, fungal colonies were removed from the agar by peeling the biomass from the 118 dialysis membranes using a sterile razor blade. Mycelia were oven-dried at 100°C until reaching constant 119 weight for at least 2 days. Results were expressed in terms of a tolerance index (TI) as reported in Ceci et 120 al. (2015c). After the dialysis membrane and mycelium were removed, the surface pH of the agar was 121 measured at specific intervals across the diameter of the Petri dish using a conical tip FC 202D pH 122 electrode (Hanna Instruments, Woonsocket, RI, USA) and a pH portable meter, HI 99161 (Hanna 123 Instruments, Woonsocket, RI, USA).

125 Genetic identification of *P. griseofulvum* FBL 500

126 P. griseofulvum FBL 500 was isolated and previously identified through conventional taxonomic keys on 127 the basis of macro- and microscopic features. ITS sequence analysis was carried out in order to confirm the 128 previous taxonomical identification findings. P. griseofulvum FBL 500 was inoculated by transferring 129 mycelial portions with a flamed glass rod from the actively growing periphery of stock colonies grown in 130 solid Czapek-Dox medium. After 15 days, fungal colonies were removed from the agar by peeling the 131 biomass from the dialysis membranes using a sterile razor blade. Mycelia were freeze-dried, pulverized by 132 using liquid N_2 by using mortal and with the addition of polyvinylpolypyrrolidone to protect DNA. DNA 133 purification and extraction was carried out from single replicates following the standard cetyltrimethyl 134 ammonium bromide (CTAB) protocol (Doyle and Doyle 1987). Identification of P. griseofulvum FBL 500 135 was achieved after extraction of fungal DNA and ITS sequence analysis using ITS 1F (5'-136 CTTGGTCATTTAGAGGAAGTAA-3') and ITS 4 (5' -TCCTCCGCTTATTGATATGC-3') primers for 137 polymerase chain reaction (PCR) analysis (Bellemain et al. 2010). DNA was quantified by using Nanodrop. 138 The PCR amplification was performed in a final volume of 25 µl using: 2.5 µl of 10× buffer, 2.5 mM 139 MgCl₂, 0.2 µM of each primer, 0.2 mM of dNTPs, 1.5 u. of DNA Taq polymerase (Promega, Milan, IT) 140 and 5 ng DNA. PCR was conducted using 35 cycles of the following reaction conditions: initial 141 denaturation at 95°C for 4 min, 39 cycles of three-step cycling (denaturation at 92°C for 50 s, primer 142 annealing at 55°C for 50 s and extension at 72°C for 50 s) and final extension at 72°C for 10 min. The PCR 143 products were sent to Macrogen Europe (Amsterdam, Netherland) for purification and sequencing. 144 Sequences were edited using the software CHROMAS 2.33 (Technelysium Pty Ltd, Australia). The partial 145 sequence of 18S ribosomal RNA gene, the complete sequence of the internal transcribed spacer 1, the 146 complete sequence of the 5.8S ribosomal RNA gene, the complete sequence of the internal transcribed 147 spacer 2 along and the partial sequence of the 28S ribosomal RNA gene were pairwise compared with those 148 available in the public online databases International Nucleotide Sequence Databases using the BLAST 149 search program (Altschul et al. 1997) and UNITE database (Kõljalg et al. 2005; Abarenkov et al. 2010). 150 The genetic sequence for P. griseofulvum FBL 500 was deposited in GenBank with the accession number 151 KY560469.

152

153 Experiment A. Metal-amended plates and inoculation

154 Stock solutions of vanadium pentoxide, V_2O_5 (Riedel-deHaën, Seelze, Germany), ammonium 155 metavanadate, NH₄VO₃ (Merck, Darmstadt, Germany), and lead carbonate, PbCO₃ (GPR), were prepared 156 from oven-sterilized aliquots (48 h, 100°C). Growth experiments at different concentrations of ammonium 157 metavanadate (2.5 and 5 mM) were performed. In addition, combinations of insoluble 2.5 mM lead 158 carbonate with insoluble 1.25 mM vanadium(V) oxide or with slightly soluble 2.5 mM ammonium 159 metavanadate were examined to study the toxic effects of both substances on *P. griseofulvum* FBL 500. 160 These concentrations take account of the range of concentrations of these metals found in soils and volcanic 161 rocks at the same site. The fungus was isolated where vanadium concentrations were over the range 3–6 162 mM, while lead concentrations were <1 mM (data not shown). Observations of colonies and media were 163 performed using light- and stereo-microscopy to monitor growth, sporulation, pigment production and

164

secondary mineral precipitation.

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166 Experiment B. Batch experiments on HCH biodegradation by P. griseofulvum FBL 500

167 High purity mixture of hexachlorocyclohexane isomers ($\alpha:\beta:\gamma:\delta=1:1:1:1$) was acquired from Sigma-Aldrich 168 (Seelze, Germany). Ethyl acetate, acetone and n-hexane were all purchased from ROMIL Ltd (Cambridge, 169 UK) with chemical purity >99.9%. The internal standard γ -HCH-d₆ was obtained from CDN Isotopes 170 (Pointe-Claire, Quebec, Canada) and stored at 4°C until use. The tests were carried out at 25°C under 171 shaking conditions at 110 rpm, and with the addition of a mixture of α -HCH, β -HCH, γ -HCH, and δ -HCH 172 isomers (1:1:1:1) to a final concentration of 4 mg/l. Prior to HCH addition, the fungus was grown for 26 173 days, and the concentration of all isomers and formation of fungal metabolites were monitored at regular 174 intervals over 23 days. Uninoculated sterile flasks with HCH mixture and flasks with fungal culture but 175 without HCH mixture were used as chemical and biological controls. Growth of P. griseofulvum FBL 500 176 and tolerance to the HCH mixture were evaluated by biomass yield. Collected mycelial pellets were oven-177 dried at 100°C for at least 2 d, until reaching constant weight and fungal tolerance was evaluated using a 178 tolerance index (TI), based on dry weights as described above.

179

180 Experiment C. Batch experiments on HCH and vanadium biotransformation by *P. griseofulvum* FBL 181 500

182 Tests with at least three replicates were carried out at 25°C in the dark with different combinations of 183 vanadium and HCH isomeric mixture. HCH was directly added in each Petri dish to the final concentration, 184 while Czapek-Dox medium was at a temperature between 40 to 50°C to homogenize HCH in the agar and 185 to avoid HCH volatilization. Growth experiments with insoluble 2.5 mM vanadium(V) pentoxide, and with 186 4 mg/l and 50 mg/l HCH mixture were performed. In addition, a combination of insoluble 2.5 mM 187 vanadium(V) pentoxide and 4 mg/l HCH mixture was used to study any synergic effects on P. griseofulvum 188 FBL 500. The production of fungal metabolites was monitored in biomass and in agar for 20 d at regular 189 intervals. Data of fungal growth (diameter, dry weight) and pH were collected, and tolerance indices were 190 calculated as described above.

191

192 Chemical analysis

193 In experiment B, 5 ml samples were collected from the culture flasks: 1 ml aliquots were spiked with γ -194 HCH-d₆ as an internal standard and extracted using solid phase extraction (SPE) cartridges packed with 195 Graphitized Carbon Black (GCB — Carbograph, Rome, Italy). HCHs were retained on the solid phase and 196 eluted with 10 ml ethyl acetate through a vacuum manifold (Grayledge Pump & Industrial, LLC, Pelham, 197 NE). In experiment C, the agar and the membranes were collected and analysed to measure HCH concentration and metabolites. Specimens were posed in glass tubes with 15 mL ethyl acetate and sonicated
 for 30 min. After centrifugation at 2000 rpm, ethyl acetate solutions were recovered in vials and 10 ml were
 analyzed.

201 202

Residual HCH for each isomer in the culture medium was calculated using the formula reported in Salam and Das (2014) and in Ceci et al. (2015b).

203 In order to study the uptake of each HCH isomer and fungal transformation, mycelial biomass was 204 collected in all experiments and extracted with Dionex ASE200 Accelerated Solvent Extractor (Dionex, 205 Sunnyvale, USA) after γ -HCH-d₆ addition as an internal standard. Operating conditions are reported in 206 Ceci et al. (2015b). The concentration of isomers and formation of metabolites were analyzed by gas 207 chromatography-mass spectrometry (GC-MS). HCH isomers and the possible metabolites were analyzed 208 using a Hewlett-Packard 6890 gas chromatograph with a 5973A mass selective detector (Agilent 209 Technologies, Palo Alto, California, USA). GC-MS analyses of liquid media were performed to detect any 210 possible intermediate metabolites of dechlorination of HCH (e.g. pentachlorocyclohexene, 211 tetrachlorocyclohexene), or intermediates of HCH reductive dechlorination and hydroxylation as in other 212 studies with bacteria and fungi (Phillips et al. 2005; Guillén-Jiménez et al. 2012).

213

Analysis of fungal metabolic profile in the presence of HCH mixture and vanadium using Biolog FFmicroplates

216 The Phenotype MicroArrayTM system (Pinzari et al. 2016) was used to gather information on the whole 217 phenotype of the P. griseofulvum FBL 500 strain and on the effects of organic and inorganic toxic 218 compounds on its carbon metabolism. The method we used was based on the inoculation of a fungal spore 219 suspension in FF MicroPlates (BiologTM, Inc., Hayward, California, USA) (Bochner et al. 2001;Pinzari et al. 220 2016). A combined inoculum of the fungus with: a) 4 mg/l HCH mixture in toluene, or; b) 2.5 mM V_2O_5 , or 221 c) with the combination of 4 mg/l HCH mixture and 2.5 mM V₂O₅ were performed in FF MicroPlateTM 222 arrays in triplicate. The inoculation procedure for pure cultures of P. griseofulvum FBL 500 in the arrays 223 was based on the protocol used by Tanzer et al. (2003).

224 Conidia of the fungus were obtained by cultivation of the pure strain on 2% MEA plates in the 225 dark at 25°C for 7 d. Operating conditions are reported in Ceci et al. 2015b. The optical density of Biolog 226 plates was read using a microplate reader (Molecular device, Vmax) at 490 nm (OD₄₉₀), which was used to 227 measure the intensity of the purple colour resulting from the reduction of the tetrazolium redox dye (p-228 iodonitrotetrazolium), present in the wells of the FF plates, through the action of fungal succinate 229 dehydrogenase as a proxy for respiratory activity. Moreover, optical density at 750 nm (OD_{750}) was used to 230 assess fungal biomass and mycelial growth (Tanzer et al. 2003;Ceci et al. 2015b). Immediately after 231 inoculation, OD₄₉₀ and OD₇₅₀ were measured in order to zero the spectrophotometer specifically for each 232 Biolog plate. Plates were then read at intervals of 24, 48, 72, 96, 168, 192 and 240 h of incubation (Tanzer 233 et al. 2003; Ceci et al. 2015b). In order to evaluate possible redox effects of vanadium pentoxide on the 234 tetrazolium dye, two sets of microplates with V2O5 and V2O5 with HCH mixture were prepared without the fungus inoculum. Data obtained from the Phenotype MicroArrayTM assays were used to compare the three growth conditions to evaluate the overall differences in metabolism (co-metabolism, inhibition, synergic effects) by studying the utilization of different substrates in the absence or presence of the xenobiotic and PTEs (vanadium and HCH) and their combinations.

239

240 Statistical analysis

R elaboration and programming software, version 3.3.2 (The R Foundation for Statistical Computing, Vienna, Austria) and the statistical package XLStat (Addinsoft 2007-Pro, Paris, France) were used to perform statistical analyses (Fahmy and Aubry 2003). They were used to perform one-way ANOVA tests on means for dry weight, diametric growth, surface pH, and HCH concentrations (at least three replicate determinations were used). One-way ANOVA tests on means were performed for the OD₄₉₀ and OD₇₅₀ values for all different treatments at 168 h, when a plateau was reached in the metabolic curves.

247 The Phenotype MicroArray data were further analysed using the opm R package (Vaas et al. 2013). All the 248 OD values were combined in a dataset which comprised three replicates \times 96 substrates \times four treatments 249 (control, HCH mixture in toluene, vanadium, combination of vanadium and HCH mixture in toluene) × two 250 metabolic parameters (respiration and mycelial growth), giving rise to 2304 individual phenotypic curves. 251 Comparison of substrate utilization in the different treatments was carried out using the estimated curve 252 parameter of A — maximum height of the metabolic curve — calculated with the opm package and plotted 253 as confidence-interval plots and heatmaps (Vaas et al. 2012). The confidence intervals and the ANOVA 254 were performed on the optical density measurements for 168 h in FF microplates.

256 Results

257 Genetic identification of *P. griseofulvum* FBL 500

Sequence analysis using the internal transcribed spacer (ITS) regions (ITS1F and ITS4 primers) confirmed
 the identification of *P. griseofulvum* FBL 500 through conventional taxonomic keys. A 100% sequence
 identity over the BLAST alignment was obtained with the name of the reference sequence being *P. griseofulvum* SH207147.07FU and KJ467353 for UNITE and NCBI databases, respectively.

262

263 Experiment A. Fungal interactions with metals

264 The presence of vanadium and lead compounds did not inhibit colony expansion of P. griseofulvum FBL 265 500 under all test conditions. In fact, all the calculated tolerance indices were greater than 1 and colony 266 extension rates in the presence of metal (R_t) showed higher values than the control extension rate (R_c) , 267 which was statistically significant in all tests (P < 0.01) (Table 2, Online Resource). Multiple comparisons 268 using the Tukey test showed that fungal extension in all tests was significantly higher than the control (P <269 0.01). Tolerance indices (TI) were used to compare biomass yields of control and test conditions (Table 3, 270 Online Resource). Biomass yields were stimulated (TI > 100%) by the lead and vanadium compounds. The 271 highest TI value occurred with 2.5 mM NH_4VO_3 (TI = 231.49%). Multiple comparisons using the Tukey 272 test revealed that biomass production in all tests was significantly higher than the control (P < 0.01). Table 273 4 (Online Resource) shows the differences (ΔpH) between average surface pH values of uninoculated agar 274 and agar underneath fungal colonies of P. griseofulvum FBL 500, growing on Czapek-Dox medium in the 275 different treatment conditions. In uninoculated agar of all experimental conditions, the range of pH medium 276 was 5–5.3. In contrast, for test conditions, the medium pH decreased after growth for 12 days (pH =4-4.3) 277 with the exceptions of those with combinations of vanadium and lead compounds where the pH values 278 were similar to the uninoculated controls. In the latter, the buffering effect of carbonate might have 279 neutralized fungal acidification of the media (Table 4, Online Resource). Sporulation in tests was limited to 280 the central part of the colonies, occasionally green coloured, and the production of vellow-orange pigments 281 and exudates was observed to be higher in treatments than in controls. No secondary mineral precipitation 282 was observed over 12 days. In contrast, the complete dissolution of insoluble crystals of lead carbonate and 283 vanadium pentoxide by P. griseofulvum FBL 500 occurred. Notably, few crystals were observed 284 underneath growing fungal colonies in media amended with combinations of ammonium metavanadate and 285 lead carbonate after 3 months incubation of P. griseofulvum FBL 500 (Fig. 1). The crystals were red, 286 acicular and tapered, and morphologically resembled lead oxalate. However it was not possible to extract 287 them for further identification because of their limited quantity.

288

289 Experiment B. Fungal interactions with HCH mixture of isomers in liquid Czapek-Dox medium

290 Addition of 4 mg/l isomeric HCH mixture to liquid Czapek-Dox medium had no obvious adverse effects on

the growth of *P. griseofulvum* FBL 500. Biomass yield was not strongly reduced (TI > 50%) by the

292 presence of the HCH mixture, and an average TI value of 87.2% was obtained. Results from the time-

293 dependent studies of HCH concentration in liquid batch tests are shown in Fig. 2. No measurable changes 294 in isomer concentration were detected in the abiotic controls throughout the experiments. The HCH mixture 295 was added 26 d after fungal inoculation, and the isomer concentrations were monitored for 23 d. Up to the 296 third day, the isomer concentration increased and reached a maximum concentration for all the isomers. 297 This phase was followed by a reduction of α -HCH, β -HCH, and γ -HCH in the medium with residual α -298 HCH = 63.0%, residual β -HCH = 67.1%, and residual γ -HCH = 63.5% (Fig. 2). In contrast, the δ -HCH 299 concentration appeared to be stable and close to the initial concentration of $\sim 1 \text{ mg/l}$. According to ANOVA, 300 there was a significant difference (P < 0.01) between the means of the γ -HCH concentrations measured 301 during the second and third day. There was a significant difference (P < 0.05) between the means of α -HCH 302 and the β -HCH concentrations measured over the same days. At the end of the experiment, the 303 concentrations of α -HCH, β -HCH, γ -HCH and δ -HCH in the solution were respectively 0.56 \pm 0.02 mg/l, 304 0.76 ± 0.04 mg/l, 0.48 ± 0.02 mg/l and 0.92 ± 0.02 mg/l with a resulting substantial deficit with regard to 305 the initial value of 1 mg/l for each isomer.

306

307 Experiment C. Fungal interactions with HCH mixture of isomers, vanadium and combinations of V 308 and HCH in solid Czapek-Dox medium

309 The presence of 4 mg/l HCH mixture in Czapek-Dox agar medium had no apparent adverse effects on the 310 growth of P. griseofulvum FBL 500. The fungus showed a slight reduction of the extension rate in 311 comparison to the control (significant at P < 0.01) (Table 2, Online Resource). Biomass yields were not 312 reduced (TI > 70%) by the HCH mixture (Table 3, Online Resource). The TI value was 82.8%, and the 313 difference in biomass yields was statistically significant (P < 0.01). Table 4 (Online Resource) shows the 314 differences (ΔpH) between average surface pH values of uninoculated agar and agar underneath fungal 315 colonies of P. griseofulvum FBL 500, growing on Czapek-Dox medium in the different treatment 316 conditions. The ΔpH revealed a low medium acidification (Table 4, Online Resource). After growth of P. 317 griseofulvum FBL 500, the inoculated medium pH measured in the test condition (6.67 ± 0.19) was slightly 318 higher than the one of the control (6.31 ± 0.19) and statistically significant (P < 0.01).

50 mg/l HCH mixture had a toxic effect on growth of *P. griseofulvum* FBL 500 with the extension rate being significantly reduced (Table 2, Online Resource). Biomass yields were strongly reduced (TI < 25%) (Table 3, Online Resource). The TI value was 23.8%, and the difference in biomass yields was statistically significant (P < 0.01). After growth of *P. griseofulvum* FBL 500, the average pH value of inoculated medium pH measured in the test (6.73 ± 0.11) was slightly lower than the one of the control (6.31 ± 0.19) and statistically significant (P < 0.01).

The addition of 2.5 mM V_2O_5 to Czapek-Dox did not result in adverse effects on the fungal extension rate in comparison to the control, although the differences were not statistically significant (Table 2, Online Resource). Biomass yields were stimulated (TI > 100%) by the presence of vanadium pentoxide (Table 3, Online Resource). The TI value was 123.7%, although the difference in biomass yield was not statistically significant. After growth of *P. griseofulvum* FBL 500, the average pH value of inoculated medium pH in the test (5.98 \pm 0.41) was slightly lower than the one of the control (6.31 \pm 0.19) and statistically significant (*P* < 0.01); Δ pH was slightly negative (Table 4, Online Resource).

The combination of V and HCH mixture in Czapek-Dox agar showed a slight reduction of the extension rate in comparison to the control, although differences were not significantly different (Table 2, Online Resource). Biomass yields were not reduced (TI > 70%) by the HCH mixture (Table 3, Online Resource). The TI value was 88.1%, although the difference in biomass yields was not statistically significant. The average pH value in the test (6.24 ± 0.26) was slightly lower than the control (6.31 ± 0.19), although it was not statistically significant. ΔpH was slightly negative (Table 4, Online Resource).

338

339 Fungal metabolites from HCH biotransformation

340 Different metabolic intermediates were observed in experiments B and C (Fig. 3) and different isomers of 341 tetrachlorocyclohexene pentachlorocyclohexene (PCCH) (TCCH), trichlorobenzene (TCB), 342 dichlorobenzene (DCB) and chlorobenzene (CB) were detected. In experiment B, only PCCH was always 343 detected during all the monitoring period in the solid Czapek-Dox medium experiments, while no 344 metabolites were found in liquid Czapek-Dox medium. In experiment C, after 12 d fungal growth, PCCH 345 was found in all tests in the presence of HCH mixture, i.e. 4 or 50 mg/l HCH mixture and the combination 346 of HCH mixture and vanadium. TCB was found only in the combination of HCH and vanadium and in the 347 experiment with 50 mg/l HCH mixture. In the latter, all metabolites (PCCH, TCCH, TCB, DCB, CB) were 348 detected. Moreover, the benzoates, benzaldehyde and benzyl alcohol, were also detected during the 349 experiments.

350

Analysis of fungal metabolic profile in the presence of HCH mixture and vanadium using Biolog FFmicroplates

353 The Phenotype MicroArrayTM system (Biolog Inc., Hayward, CA, USA) was used to investigate the whole 354 phenotype and nutrient utilization by P. griseofulvum FBL 500 in both control and test conditions. P. 355 griseofulvum was able to grow in the presence of 71 substrates out of the 96 available in the FF plates 356 (Pinzari et al. 2016). A heatmap of A values — the maximum height of the growth curve — for all 357 treatments and all substrates at 750 nm is presented in Fig. 4. The clustergram above the heatmap shows 358 clusters of substrates that refer to the different intensities of fungal metabolism according to the A values 359 measured. The clustergram on the left side shows clusters of treatments in which vanadium and 360 combinations of vanadium and HCH are together, as well as the control and HCH mixture, resulting in a 361 different general response of fungal metabolism to the presence of the xenobiotics. The confidence intervals 362 and the ANOVA were performed on the optical density measurements at 750 nm for the estimation of 363 mycelial growth and at 490 nm for the estimation of respiration after incubation for 168 h in FF microplates 364 in the different treatment conditions - control (no xenobiotics, only fungus), 2.5 mM vanadium oxide, 4 365 mg/l HCH mixture and combination of both xenobiotics. Significant differences were detected according to 366 specific growth substrates. P. griseofulvum FBL 500 was able to use the same substrates in the treatments

367 as in control conditions, but the presence of xenobiotics influenced fungal metabolism, reducing or 368 increasing respiration rates and/or mycelial growth. The negative effect on respiration and fungal growth 369 was significant (P < 0.05) in the presence of vanadium or with a combination of vanadium and HCH for 370 some substrates. In particular, there was a statistically significant reduction of fungal metabolism for D-371 ribose (pentoses), rhamnose (hexoses), D-gluconic acid (sugar acids), N-acetyl-D-glucosamine 372 (hexosamines), maltose, maltotriose, D-melibiose, D-raffinose (oligosaccharides), all considered glucosides 373 with the exception of arbutin, D-salicin and sucrose, all nitrogen-containing compounds and all the 374 biochemical group "other" with the exception of L-asparagine, L-phenylalanine, L-pyroglutamic acid, L-375 threonine, ethanolamine, malic acid, and sebacic acid (Fig. 5). For all other substrates (36 - 50 %) used by 376 *P. griseofulvum* FBL 500, there was no significant difference (P > 0.05) in respiration and growth between 377 control and test conditions. It is worth mentioning that for several substrates, HCH and V (alone or in 378 combination) affected fungal metabolism as revealed by calculated confidence intervals and the absorbance 379 curves for respiration and growth. In particular, for L-sorbose, D-mannitol, L-asparagine, i-erythritol and L-380 threonine, vanadium influenced fungal metabolism, increasing respiration and growth compared to the 381 control (Fig. 5). In contrast, for D-mannitol, D-cellobiose, D-sorbitol and D-glucosamine, the presence of 382 HCH inhibited fungal metabolism (Fig. 5). For ethanolamine, L-phenylalanine and D-salicin, HCH 383 increased respiration and mycelial growth (Fig. 5). For sebacic acid, the presence of HCH and the 384 combination of HCH and V resulted in metabolic stimulation (Fig. 5).

385 Discussion

386 This work examines the potential of *P. griseofulvum* FBL 500, which was isolated from polluted sites, to 387 tolerate high concentrations of the potentially toxic metals, vanadium and lead, and hexachlorocyclohexane, 388 and mediate their biotransformation.

389 Hexachlorocyclohexane (HCH) is a persistent organic pollutant (POP) of global concern with 390 potentially toxic effects on humans and ecosystems. It is a halogenated xenobiotic which has been reported 391 to be carcinogenic and an endocrine disrupter for humans and other organisms (; Ceci et al. 2015b). Three 392 isomers of hexachlorocyclohexane, α -HCH, β -HCH and γ -HCH, were included as persistent organic 393 pollutants in the 2008 Stockholm Convention because of their worldwide spread and environmental 394 persistence (Vijgen et al. 2011).

395 Vanadium is considered to be the one of the most abundant elements and one of the most 396 important metals in modern technology (Rehder 2008; Ceci et al. 2015c). Vanadium is also essential for 397 certain organisms (e.g. some algae, bacteria, fungi and lichens) as a cofactor of enzymes and a constituent 398 of metabolites (haloperoxidases, nitrogenases and amavadin) (Crans et al. 2004). In recent decades, 399 hydrocarbon fuel combustion, industrial activities and mining have increased the vanadium concentration in 400 the environment, raising concern over its spread and toxicity for humans and ecosystems (Rehder 2008; 401 Ceci et al. 2015c).

- *P. griseofulvum* has been found to successfully tolerate and accumulate potentially toxic metals
 such as Cu and Cr (Shah et al. 1999; Shi et al. 2011; Abigail et al. 2015), to tolerate and mediate the
 biotransformation of Cu-based wood preservatives (Bridbžiuviene and Levinskaite 2007) and Ni and V
 porphyrins (Cordero et al. 2015). Moreover, this fungus was reported to tolerate high concentrations of
 pyrene and mediate its biotransformation (Ravelet et al. 2000).
- 407 In experiment A, vanadium and lead compounds (NH₄VO₃, V₂O₅, PbCO₃) were used. 408 Vanadium(V) oxide and ammonium metavanadate are important products of industrial metal recovery 409 (Teng et al. 2006). Vanadium(V) oxide is often found in leachates from mining and milling activities that 410 account for the most significant fluxes of vanadium in the environment and can also originate from fossil 411 fuel combustion as an insoluble by-product (Teng et al. 2006). Lead carbonate (cerussite), a common 412 insoluble lead mineral in soil, has been used in metal tolerance tests to evaluate possible combined effects 413 of the two metals on P. griseofulvum FBL 500 and to evaluate possible metal biotransformations mediated 414 by the fungus (Ceci et al. 2015c). Aspergillus niger has been tested with the same metal compounds in 415 similar research on vanadium geomycology (Ceci et al. 2015c). In this work, P. griseofulvum FBL 500 416 showed good growth in experiment A (Table 2 and 3, Online Resource). The presence of ammonium added 417 in the test medium could explain the higher growth rates and biomass yields, being used as an additional N 418 source. However, ammonium was not present with the combination of lead carbonate with vanadium(V) 419 oxide where the extension rate and biomass yield were also significantly higher than the control. Moreover, 420 in the presence of vanadium(V) oxide and sucrose in experiment C, the TI was particularly high. Higher 421 tolerance indices than the control may be related to metabolism-dependent or -independent mechanisms of

422 tolerance/resistance, implemented to cope with the stress due to metal toxicity (Gadd 1993; 2007; Gadd et 423 al. 2012). The presence of toxic metals can strongly influence the physiology and morphology of fungal 424 mycelia and the resulting interactions can include acidolysis, complexolysis, redoxolysis, metal 425 accumulation, production of high local concentrations of extracellular enzymes, and other metabolites (e.g. 426 organic acids, siderophores, polyphenolic compounds and pigments), mycelial growth strategies (e.g. 427 explorative growth) and hyphal aggregation (e.g. phalanx growth) (; Gadd et al., 2014; Ceci et al. 2015a, 428 c). The excretion of organic acids (e.g., oxalic, citric, gluconic, and lactic acid) can be strongly influenced 429 by growth conditions such as the presence of toxic metal minerals, nutrient availability, the C and N 430 sources, pH and buffering capacity of the medium (Gadd et al. 2012; Ceci et al. 2015a, c). P. griseofulvum 431 FBL 500 was able to acidify the medium beneath the colonies either from the initial value of pH 5.5 or 432 from an initial value of pH ~7 (Table 4, Online Resource). In experiment A, the final pH values in all test 433 conditions were all acidic (pH 4–5), with the only exception being the treatments with the combinations of 434 Pb and V compounds. These were close to the uninoculated control and this is probably related to the 435 buffering effect of carbonate (Table 4, Online Resource). Generally, fungi lower the pH of their medium 436 during growth. Mechanisms such as the excretion of protons via the plasma membrane ATPase, the uptake 437 of essential cation nutrients in exchange for protons, the release of organic acids and acidification due to 438 fungal respiration can all cause acidification (Ceci et al. 2015c). On the other hand, in the presence of 439 specific metal compounds such as carbonates or apatites, buffering effects can be evident (Ceci et al. 2015a, 440 c). In this work, the formation of new biominerals, which are assumed to be lead oxalate, is evidence of 441 metal biotransformation. Similar results were observed with A. niger in the presence of V and Pb 442 compounds, vanadinite and mimetite (Ceci et al. 2015a, c). The organic acids produced by P. griseofulvum 443 FBL 500 can provide ligands for metal-complex formation, electrons for metal redox reactions and metal 444 precipitation as new mycogenic minerals, that can result in mobilization or immobilization of lead and 445 vanadium (Ceci et al. 2015a, c). In experiment C, the medium pH values were close to the control (Table 4, 446 Online Resource). The fungal growth in such conditions did not significantly change the pH of media 447 (Table 4, Online Resource), and a pH range close to neutrality could avoid some possible toxic effects in 448 acidic conditions due to the production of benzoates during HCH biodegradation. The benzoates detected 449 during these experiments were previously observed with P. griseofulvum FBL 500 in the presence of β -450 HCH (Ceci et al. 2015b).

451 In experiments B and C, P. griseofulvum FBL 500 was able to grow and tolerate high 452 concentrations of a mixture of HCH isomers. The tolerance of this fungus was tested in previous work at a 453 combination of 1 mg/l β -HCH (Ceci et al. 2015b). The presence of a 4 mg/l HCH mixture did not inhibit 454 fungal growth, and similar TI values were found for liquid and solid Czapek-Dox media. Similar results 455 were observed in previous work (Ceci et al. 2015b) confirming that the presence of different HCH isomers 456 had no obvious synergic effects on growth. In liquid medium, the biotransformation of the isomers was 457 observed to be different and isomer-specific, with β -HCH and δ -HCH being the most stable and recalcitrant. 458 These findings also agree with some other biotransformation studies (Willett et al. 1998; Phillips et al.

459 2005). The biodegradation of different mixtures of α -HCH, β -HCH, γ -HCH and δ -HCH by ligninolytic 460 white-rot fungi has been previously reported (Quintero et al. 2007; 2008; Mohapatra and Pandey 2015). To 461 our knowledge, this work is the first evidence of the biodegradation of a HCH mixture by a saprotrophic 462 fungus, isolated from a historically co-contaminated site with metals and HCH. In Quintero et al. (2007), 463 biodegradation values exhibited by Bjerkandera adusta (Willd.) P. Karst. of 91.5%, 94.5%, 78.5% and 66.1% 464 were attained after 30 d for γ -, α -, δ - and β -HCH isomers, respectively. The δ - and γ -HCH isomers were 465 degraded to between 15.1 and 70.8% by six different white-rot fungi tested and the highest β-HCH 466 biodegradation (56.6%) occurred with B. adusta (Quintero et al. 2008). In our tests with liquid Czapek-Dox 467 medium, it is possible to hypothesize the existence of a lag phase during the first 3 d followed by a 468 reduction of α -HCH, β -HCH and γ -HCH isomers in the media (Fig. 2a–2c). This could be related to slow 469 HCH solubilization in the media and/or enzyme induction and an adaptation period of the fungus to 470 addition of the HCH mixture. These phases were also observed in P. griseofulvum FBL 500 in the presence 471 of β -HCH as an adaptation period associated with time and a specific compound threshold for induction of 472 catabolic enzymes (Ceci et al. 2015b. In a slurry batch reactor using B. adusta, Quintero et al. (2007) 473 observed a lag phase during the first 5 d followed by a 7-day period in which the concentration of the HCH 474 isomers α , γ , δ , and β decreased to 73.9%, 57.4%, 40.8% and 28%, respectively. Valentin et al. (2007), 475 using the same slurry reactor system, observed a lag period of 6 d, but they related this to stressful culture 476 conditions, e.g. high agitation rates, or a decrease in oxygen transfer. Ceci et al. 2015b observed a lag 477 period of 4 days, followed by a reduction period of β -HCH in the medium (residual β -HCH = 44.8%). 478 Detrimental conditions, such as benzoate formation, could speed up the decay phase, which implies that 479 degradation stops at shorter treatment periods (Guillén-Jiménez et al. 2012; Ceci et al. 2015b). In liquid 480 medium, no expected metabolites that have been reported in the literature for HCH biodegradation were 481 observed, while different metabolites derived from fungal dehalogenation of HCH were observed in 482 experiment C with HCH (Fig. 3). In liquid media, possible intermediates could not be detected probably 483 because of the fast kinetics of biodegradation involved in liquid medium, the low detection limits of the 484 transient products, or their different values of polarity and volatility (Guillén-Jiménez et al. 2012; Ceci et al. 485 2015b). Isomers of PCCH, TCCH, TCB (1,2,3-TCB, 1,3,5-TCB and 1,2,4-TCB), DCB (1,2-DCB, 1,3-DCB 486 and 1,4 DCB), and chlorobenzene were detected in solid Czapek-Dox medium incubated with P. 487 griseofulvum FBL 500 in experiment C (Fig. 3). A combination of vanadium and HCH did not significantly 488 inhibit fungal growth, showing similar TI values as those with the presence of HCH alone, while a 50 mg/l 489 concentration of the HCH mixture negatively reduced the fungal growth (TI < 50%). Despite this, in both 490 cases HCH biodegradation was not influenced negatively, as some metabolites were observed. 491 Intermediates of HCH reductive dechlorination and hydroxylation have been observed in other studies on 492 biotransformation of lindane by bacteria and saprotrophic fungi (Phillips et al. 2005; Salam and Das 2015). 493 PCCH is reported to be the first intermediate of HCH dehalogenation in aerobic degradation pathways 494 (Willett et al. 1998; Middeldorp et al. 2005; Camacho-Pérez et al. 2012). The presence of TCCH was 495 generally observed during anaerobic HCH degradation by bacteria, while a *Pseudomonas* sp. isolated from

496 soil was able to transform lindane aerobically producing PCCH and TCCH (Phillips et al. 2005). Microbial 497 degradation of lindane has been reported to produce volatile transformation products such as chlorinated 498 benzenes and phenols (Phillips et al. 2005; Salam et al. 2013). Previously, tetrachlorocyclohexane and 499 tetrachlorocyclohexanol were observed in the biodegradation of lindane by white rot-fungi (Phillips et al. 500 2005), while PCCH and benzoic acid derivatives were observed in aerobic degradation of lindane by 501 Fusarium verticillioides AT-100 (Guillén-Jiménez et al. 2012), y-PCCH and different metabolites were 502 observed in lindane biodegradation by Candida VITJzN04 (Salam and Das 2014), while the proposed 503 degradation route for lindane in *Rhodotorula* sp. is very similar to the one that can be hypothesized in this 504 investigation and γ -PCCH, 1,2,4-TCB, 1,2.DCB, CB and other intermediates were reported (Salam et al. 505 2013).

506 The Phenotype MicroArrayTM microplate system was used to study the whole phenotype of P. 507 griseofulvum FBL 500 in response to the different combinations of V and HCH. This fungus was able to 508 use ~ 74 % of the all available microplate substrates. This confirms a high metabolic versatility of P. 509 griseofulvum FBL 500 in different environmental habitats. The different treatments did not change the 510 number of substrates used, but instead the rate of utilization, with significant effects on fungal metabolism. 511 The presence of vanadium alone or in combination with HCH negatively influenced respiration and growth 512 in the presence of different compounds, according to their different chemical categories. The results 513 showed a strong similarity with the treatments with vanadium and with the combination of V and HCH. 514 This can be explained by V inhibition of specific enzymes which are essential for the activation of different 515 metabolic pathways of substrate utilization and also masks the effects of HCH. For instance, enzymatic V 516 inhibition could be due to chemical similarity of vanadate with phosphate, and the resulting competition for 517 phosphorylation of some enzymes, which, in turn, could reduce utilization of specific substrates and 518 metabolism. It has been demonstrated that vanadium enters Neurospora crassa as vanadate, which is a 519 potent inhibitor of the plasma membrane ATPase when cells are growing in an alkaline medium and are 520 depleted for phosphate (Ceci et al. 2015c). Functional and phenotypic studies on microbial communities 521 from metal polluted sites using Biolog microplates with high concentrations of Cu and Zn, showed negative 522 effects on community functions due to metal toxicity, but also pollution-induced community tolerance to 523 metals (Klimek and Niklińska 2007). In particular, the fungal community was shown to be less sensitive to 524 metal toxicity than the co-occurring bacterial community (Klimek and Niklińska 2007). Kong et al. 2006 525 observed that synergic effects of Cu and the antibiotic oxytetracycline on a microbial community 526 negatively influenced functional diversity, resulting in a significantly stronger negative effect for each 527 substrate group on the utilization potential of carboxylic acids and carbohydrates than those of 528 oxytetracycline or Cu alone (Kong et al. 2006). It is worth noting that in our study it has been found that for 529 nearly 50% of substrates used by P. griseofulvum FBL 500 there was no significant difference among the 530 metabolic curves for all the treatments. This may indicate that the mechanisms of tolerance/resistance 531 utilized by the fungus were adequate to cope with the stress imposed by the different combinations of 532 xenobiotics and PTEs. Apparently P. griseofulvum FBL 500 could transform metal and/or organic

533 pollutants into less toxic forms since in their presence it showed the same metabolic profile as the control. 534 In fact, for some substrates higher values of A, the maximum height of the metabolic curve, were observed 535 or higher values of OD for HCH, V and combinations of both. These evidences may represent a stimulatory 536 effect of HCH, V or combinations of both in the co-metabolism of specific organic compounds. In the 537 presence of carbon- and nitrogen-containing compounds, specific metabolic pathways, such as the pentose 538 phosphate pathway and glucuronate interconversion, and the presence of relatively non-specific enzymes, 539 such as laccases and cellulose dehydrogenase saprotrophic can be activated, and fungi, could play roles in 540 the biotransformation of HCH alone or with V (Cameron et al. 2000; Mander et al. 2006; Ceci et al. 2015b). 541 V can help in cancer treatment, because it may be actively involved in oxidative radicals' production by 542 Fenton reactions against the tumor cells (Rehder, 2008). These reactions could also enhance the enzymatic 543 catalyst reactions of HCH biotransformation by laccases or other fungal enzymes, involved also in the 544 biodegradation of other xenobiotics. Moreover, V can activate specific enzymes, such as V-dependent 545 haloperoxidases, that can further enhance the degradation of recalcitrant compounds (Ceci et al., 2015c)

546 The *P. griseofulvum* FBL 500 strain was isolated from an environment extensively polluted by 547 toxic organic and inorganic compounds over several years. Therefore the fungus occurred in an 548 environment where stress was a constant condition and stimulus for survival. The experiments exposed the 549 fungus to toxic compounds singly and in combination with the responses of the fungus being varied. On 550 one hand, the organism showed a generic response which was the same in both the presence of organic 551 compounds and metals. It is likely that fungus initiated a survival or buffering metabolism generically 552 directed to cope with immediate extreme environmental conditions, apparently without distinguishing 553 between metal or organic stressors. It is worth mentioning that the different treatments did not affect the 554 number of carbon sources used by the fungus, but instead modified their rate of utilization suggesting a 555 significant systemic effect on fungal metabolism, and not just on a single pathway. P. griseofulvum FBL 556 500 was able to acidify the medium and the formation of new biominerals was evidence of the occurrence 557 of metal biotransformation. This suggests that the fungus possesses particular mechanisms of tolerance to 558 metal toxicity. When exposed to HCH, however, the biotransformation of the xenobiotic by the fungus did 559 not significantly change the pH of the media, which remained close to neutrality. This could protect the 560 fungus from toxic effects that occur in acidic conditions due to the production of benzoates from HCH 561 biodegradation. In fungi, pH plays important roles in other ecological aspects. For example, in postharvest 562 pathogens, fungal pH modulation of the host environment regulates a multitude of enzymes implied in 563 fungal pathogenicity (Alkan et al. 2012). Moreover, for Paecilomyces marquandii (Massee) S. Hughes, the 564 pH of the medium affects the production of reactive oxygen species during biodegradation of the pesticide 565 alachlor. Neutral pH was favourable both for alachlor biodegradation and for oxidative stress reduction 566 (Słaba et al. 2015). The effectiveness of mycelial protection by anti-oxidative enzymes from oxidative 567 stress was observed to be dependent on the environmental pH, exposure time and fungal growth phase 568 (Słaba et al. 2015). Interestingly, the synthesis of laccase and other extracellular peroxidases by some 569 filamentous fungi was higher in the presence of intracellular oxidative stress (Chanda et al. 2015). Besides 570 a reaction to generic stress, *P. griseofulvum* FBL 500 was also capable of activating pathways to use 571 different substrates, tolerating a high concentration of a combination of HCH and V with a supposed co-572 metabolic mechanism. In fact, *P. griseofulvum* FBL 500 was able to tolerate high concentrations of a 573 mixture of HCH isomers, an ability described so far only in a few fungal species.

574 In this research, we have provided evidence of fungal tolerance to metals and HCH, and its ability 575 in metal biomineralization, metal mobilization and biotransformation of HCH mixtures of isomers, even in 576 the presence of vanadium at high concentrations. This could therefore be relevant to bioremediation 577 treatments (bioaugmentation, biostimulation) of co-contaminated sites through native fungal species

578 selected for extreme environmental conditions.

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591 **References**

- Abarenkov K, Henrik Nilsson R, Larsson K-H, Alexander IJ, Eberhardt U, Erland S, Høiland K, Kjøller R,
 Larsson E, Pennanen T, Sen R, Taylor AFS, Tedersoo L, Ursing BM, Vrålstad T, Liimatainen K,
 Peintner U, Kõljalg U (2010) The UNITE database for molecular identification of fungi recent
 updates and future perspectives. New Phytol 186:281–285. doi: 10.1111/j.14698137.2009.03160.x
- Abigail MEA, Samuel MS, Chidambaram R (2015) Hexavalent chromium biosorption studies using *Penicillium griseofulvum* MSR1 a novel isolate from tannery effluent site: Box–Behnken
 optimization, equilibrium, kinetics and thermodynamic studies. J Taiwan Inst Chem Eng 49:156–
 164. doi: 10.1016/j.jtice.2014.11.026
- Alisi C, Musella R, Tasso F, Ubaldi C, Manzo S, Cremisini C, Sprocati AR (2009) Bioremediation of
 diesel oil in a co-contaminated soil by bioaugmentation with a microbial formula tailored with
 native strains selected for heavy metals resistance. Sci Total Environ 407:3024–3032. doi:
 10.1016/j.scitotenv.2009.01.011
- Alkan N, Espeso EA, Prusky D (2012) Virulence regulation of phytopathogenic fungi by pH. Antioxid
 Redox Signal 19:1012–1025. doi: 10.1089/ars.2012.5062
- Altschul SF, Madden TL, Schäffer AA, Zhang J, Zhang Z, Miller W, Lipman DJ (1997) Gapped BLAST
 and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res
 25:3389–3402. doi: 10.1093/nar/25.17.3389
- Bellemain E, Carlsen T, Brochmann C, Coissac E, Taberlet P, Kauserud H (2010) ITS as an environmental
 DNA barcode for fungi: an in silico approach reveals potential PCR biases. BMC Microbiol. doi:
 10.1186/1471-2180-10-189
- Bernardini A, Salvatori E, Di Re S, Fusaro L, Nervo G, Manes F (2016) Natural and commercial *Salix*clones differ in their ecophysiological response to Zn stress. Photosynthetica 54:56–64. doi:
 10.1007/s11099-015-0155-9
- Bernini R, Pelosi C, Carastro I, Venanzi R, Di Filippo A, Piovesan G, Ronchi B, Danieli PP (2016)
 Dendrochemical investigation on hexachlorocyclohexane isomers (HCHs) in poplars by an
 integrated study of micro-Fourier transform infrared spectroscopy and gas chromatography. Trees
 30:1455–1463. doi: 10.1007/s00468-015-1343-8
- 620Bochner BR, Gadzinski P, Panomitros E (2001) Phenotype MicroArrays for high-throughput phenotypic621testing and assay of gene function. Genome Res 11:1246–1255. doi: 10.1101/gr.186501

- Bridbžiuviene D, Levinskaite L (2007) Fungal tolerance towards copper-based wood preservatives.
 Biologija 53:54–61
- 624 Camacho-Pérez B, Ríos-Leal E, Rinderknecht-Seijas N, Poggi-Varaldo HM (2012) Enzymes involved in
 625 the biodegradation of hexachlorocyclohexane: A mini review. J Environ Manage 95,
 626 Supplement:S306–S318. doi: 10.1016/j.jenvman.2011.06.047
- 627 Cameron MD, Timofeevski S, Aust SD (2000) Enzymology of *Phanerochaete chrysosporium* with respect
 628 to the degradation of recalcitrant compounds and xenobiotics. Appl Microbiol Biotechnol 54:751–
 629 758. doi: 10.1007/s002530000459
- 630 Ceci A, Kierans M, Hillier S, Persiani AM, Gadd GM (2015a) Fungal bioweathering of mimetite and a
 631 general geomycological model for lead apatite mineral biotransformations. Appl Environ
 632 Microbiol 81:4955–4964. doi: 10.1128/AEM.00726-15
- 633 Ceci A, Maggi O, Pinzari F, Persiani AM (2012) Growth responses to and accumulation of vanadium in
 634 agricultural soil fungi. Appl Soil Ecol 58:1–11. doi: 10.1016/j.apsoil.2012.02.022
- 635 Ceci A, Pierro L, Riccardi C, Pinzari F, Maggi O, Persiani AM, Gadd GM, Petrangeli Papini M (2015b)
 636 Biotransformation of β-hexachlorocyclohexane by the saprotrophic soil fungus *Penicillium*637 *griseofulvum*. Chemosphere 137:101–107. doi: 10.1016/j.chemosphere.2015.05.074
- 638 Ceci A, Rhee YJ, Kierans M, Hillier S, Pendlowski H, Gray N, Persiani AM, Gadd GM (2015c)
 639 Transformation of vanadinite [Pb₅(VO₄)₃Cl] by fungi. Environ Microbiol 17:2018–2034. doi: 10.1111/1462-2920.12612
- 641 Chanda A, Gummadidala PM, Gomaa OM (2015) Mycoremediation with mycotoxin producers: a critical
 642 perspective. Appl Microbiol Biotechnol 100:17–29. doi: 10.1007/s00253-015-7032-0
- 643 Cordero PRF, Bennett RM, Bautista GS, Aguilar JPP, Dedeles GR (2015) Degradation of nickel
 644 protoporphyrin disodium and vanadium oxide octaethylporphyrin by Philippine microbial
 645 consortia. Bioremediation J 19:93–103. doi: 10.1080/10889868.2013.827616
- 646 Crans DC, Smee JJ, Gaidamauskas E, Yang L (2004) The chemistry and biochemistry of vanadium and the
 647 biological activities exerted by vanadium compounds. Chem Rev 104:849–902
- 648 Czaplicki LM, Cooper E, Ferguson PL, Stapleton HM, Vilgalys R, Gunsch CK (2016) A new perspective
 649 on sustainable soil remediation—case study suggests novel fungal genera could facilitate in situ
 650 biodegradation of hazardous contaminants. Remediat J 26:59–72. doi: 10.1002/rem.21458

- Doyle JJ, Doyle LJ (1987) A rapid DNA isolation procedure for small quantities of fresh leaf tissue.
 Phytochem Bull 19:11–15
- 653 Fahmy T, Aubry P (2003) XLSTAT-pro, version 7.0. Paris, FR

- 655 Gadd GM (1993) Interactions of fungi with toxic metals. New Phytol 124:25–60. doi: 10.1111/j.1469-656 8137.1993.tb03796.x
- Gadd GM (2007) Geomycology: biogeochemical transformations of rocks, minerals, metals and
 radionuclides by fungi, bioweathering and bioremediation. Mycol Res 111:3–49. doi:
 10.1016/j.mycres.2006.12.001
- 660 Gadd GM (2004) Mycotransformation of organic and inorganic substrates. Mycologist 18:60–70. doi:
 661 10.1017/S0269-915X(04)00202-2
- 662 Gadd GM (2010) Metals, minerals and microbes: geomicrobiology and bioremediation. Microbiology
 663 156:609-643. doi: 10.1099/mic.0.037143-0
- Gadd GM (2013) Geomycology: fungi as agents of biogeochemical change. Biol Environ Proc R Ir Acad
 113:1–15. doi: 10.3318/BIOE.2013.16
- Gadd GM, Bahri-Esfahani J, Li Q, Rhee YJ, Wei Z, Fomina M, Liang X (2014) Oxalate production by
 fungi: significance in geomycology, biodeterioration and bioremediation. Fungal Biol Rev 28:36–
 55. doi: 10.1016/j.fbr.2014.05.001
- 669 Gadd GM, Rhee YJ, Stephenson K, Wei Z (2012) Geomycology: metals, actinides and biominerals.
 670 Environ Microbiol Rep 4:270–296
- 671 Griffith CM, Baig N, Seiber JN (2015) Contamination from industrial toxicants. In: Cheung PCK, Mehta
 672 BM (eds) Handbook of Food Chemistry. Springer, Berlin, Heidelberg, DE, pp 719–751
- Guillén-Jiménez F de M, Cristiani-Urbina E, Cancino-Díaz JC, Flores-Moreno JL, Barragán-Huerta BE
 (2012) Lindane biodegradation by the *Fusarium verticillioides* AT-100 strain, isolated from *Agave tequilana* leaves: kinetic study and identification of metabolites. Int Biodeterior Biodegrad 74:36–
 47. doi: 10.1016/j.ibiod.2012.04.020
- Harms H, Schlosser D, Wick LY (2011) Untapped potential: exploiting fungi in bioremediation of
 hazardous chemicals. Nat Rev Microbiol 9:177–192. doi: 10.1038/nrmicro2519

- Klimek B, Niklińska M (2007) Zinc and copper toxicity to soil bacteria and fungi from zinc polluted and
 unpolluted soils: a comparative study with different types of Biolog plates. Bull Environ Contam
 Toxicol 78:112–117. doi: 10.1007/s00128-007-9045-6
- Kõljalg U, Larsson K-H, Abarenkov K, Nilsson RH, Alexander IJ, Eberhardt U, Erland S, Høiland K,
 Kjøller R, Larsson E, Pennanen T, Sen R, Taylor AFS, Tedersoo L, Vrålstad T (2005) UNITE: a
 database providing web-based methods for the molecular identification of ectomycorrhizal fungi.
 New Phytol 166:1063–1068. doi: 10.1111/j.1469-8137.2005.01376.x
- Kong W-D, Zhu Y-G, Fu B-J, Marschner P, He J-Z (2006) The veterinary antibiotic oxytetracycline and Cu
 influence functional diversity of the soil microbial community. Environ Pollut 143:129–137. doi:
 10.1016/j.envpol.2005.11.003
- 689
- Lange L, Bech L, Busk PK, Grell MN, Huang Y, Lange M, Linde T, Pilgaard B, Roth D, Tong X (2012)
 The importance of fungi and of mycology for a global development of the bioeconomy. IMA
 Fungus 3:87–92. doi: 10.5598/imafungus.2012.03.01.09
- Ma X, Ling Wu L, Fam H (2014) Heavy metal ions affecting the removal of polycyclic aromatic
 hydrocarbons by fungi with heavy-metal resistance. Appl Microbiol Biotechnol 98:9817–9827.
 doi: 10.1007/s00253-014-5905-2
- Mace GM, Norris K, Fitter AH (2012) Biodiversity and ecosystem services: a multilayered relationship.
 Trends Ecol Evol 27:19–26. doi: 10.1016/j.tree.2011.08.006
- Mander GJ, Wang H, Bodie E, Wagner J, Vienken K, Vinuesa C, Foster C, Leeder AC, Allen G, Hamill V,
 Janssen GG, Dunn-Coleman N, Karos M, Lemaire HG, Subkowski T, Bollschweiler C, Turner G,
 Nüsslein B, Fischer R (2006) Use of laccase as a novel, versatile reporter system in filamentous
 fungi. Appl Environ Microbiol 72:5020–5026. doi: 10.1128/AEM.00060-06
- Middeldorp PM, van Doesburg W, Schraa G, Stams AM (2005) Reductive dechlorination of
 hexachlorocyclohexane (HCH) isomers in soil under anaerobic conditions. Biodegradation
 16:283–290. doi: 10.1007/s10532-004-1573-8
- Mishra A, Malik A (2014) Novel fungal consortium for bioremediation of metals and dyes from mixed
 waste stream. Bioresour Technol 171:217–226. doi: 10.1016/j.biortech.2014.08.047
- Mohapatra S, Pandey M (2015) Biodegradation of hexachlorocyclohexane (HCH) isomers by white rot
 fungus, *Pleurotus florida*. J Bioremediation Biodegrad 6:280. doi: 10.4172/2155-6199.1000280

- Panagos P, Van Liedekerke M, Yigini Y, Montanarella L (2013) Contaminated sites in Europe: review of
 the current situation based on data collected through a European network. J Environ Public Health
 4:11.
- Phillips T, Seech A, Lee H, Trevors J (2005) Biodegradation of hexachlorocyclohexane (HCH) by
 microorganisms. Biodegradation 16:363–392. doi: 10.1007/s10532-004-2413-6
- Pinzari F, Ceci A, Abu-Samra N, Canfora L, Maggi O, Persiani A (2016) Phenotype MicroArrayTM system
 in the study of fungal functional diversity and catabolic versatility. Res Microbiol 167:710–722.
 doi: 10.1016/j.resmic.2016.05.008
- Polti MA, Aparicio JD, Benimeli CS, Amoroso MJ (2014) Simultaneous bioremediation of Cr(VI) and
 lindane in soil by actinobacteria. Int Biodeterior Biodegrad 88:48–55. doi:
 10.1016/j.ibiod.2013.12.004
- Quintero JC, Lú-Chau TA, Moreira MT, Feijoo G, Lema JM (2007) Bioremediation of HCH present in soil
 by the white-rot fungus *Bjerkandera adusta* in a slurry batch bioreactor. Int Biodeterior Biodegrad
 60:319–326. doi: 10.1016/j.ibiod.2007.05.005
- Quintero JC, Moreira MT, Feijoo G, Lema JM (2008) Screening of white rot fungal species for their
 capacity to degrade lindane and other isomers of hexachlorocyclohexane (HCH). Cienc Investig
 Agrar 35:159–167.
- Ravelet C, Krivobok S, Sage L, Steiman R (2000) Biodegradation of pyrene by sediment fungi.
 Chemosphere 40:557–563. doi: 10.1016/S0045-6535(99)00320-3
- 728 Rehder D (2008) Bioinorganic vanadium chemistry. John Wiley & Sons Ltd, West Sussex, U.K
- Salam JA, Das N (2014) Lindane degradation by *Candida* VITJzN04, a newly isolated yeast strain from
 contaminated soil: kinetic study, enzyme analysis and biodegradation pathway. World J Microbiol
 Biotechnol 30:1301–1313. doi: 10.1007/s11274-013-1551-6
- Salam JA, Das N (2015) Degradation of lindane by a novel embedded bio-nano hybrid system in aqueous
 environment. Appl Microbiol Biotechnol 99:2351–2360. doi: 10.1007/s00253-014-6112-x
- Salam JA, Lakshmi V, Das D, Das N (2013) Biodegradation of lindane using a novel yeast strain,
 Rhodotorula sp. VITJzN03 isolated from agricultural soil. World J Microbiol Biotechnol 29:475–
 487. doi: 10.1007/s11274-012-1201-4
- Sandrin TR, Maier RM (2003) Impact of metals on the biodegradation of organic pollutants. Environ
 Health Perspect 111:1093–1101.

- Shah MP, Vora SB, Dave SR (1999) Evaluation of potential use of immobilized *Penicillium griseofulvum*in bioremoval of copper. Process Metallurgy 9: 227–235. doi.org/10.1016/S1572-4409(99)801126
- 742Shi Z, Bai S, Tian L, Jiang H, Zhang J (2011) Molecular detection of *Penicillium griseofulvum* as the743coastal pollution indicator. Curr Microbiol 62:396–401. doi: 10.1007/s00284-010-9720-4
- Słaba M, Różalska S, Bernat P, Szewczyk R, Piątek MA, Długoński J (2015) Efficient alachlor degradation
 by the filamentous fungus *Paecilomyces marquandii* with simultaneous oxidative stress reduction.
 Bioresour Technol 197:404–409. doi: 10.1016/j.biortech.2015.08.045
- Tanzer M, Arst H Jr, Skalchunes A, Coffin M, Darveaux B, Heiniger R, Shuster J (2003) Global nutritional
 profiling for mutant and chemical mode-of-action analysis in filamentous fungi. Funct Integr
 Genomics 3:160–170. doi: 10.1007/s10142-003-0089-3
- Teng Y, Ni S, Zhang C, Wang J, Lin X, Huang Y (2006) Environmental geochemistry and ecological risk
 of vanadium pollution in Panzhihua mining and smelting area, Sichuan, China. Chin J Geochem
 25:379–385. doi: 10.1007/s11631-006-0378-3
- Tigini V, Prigione V, Di Toro S, Fava F, Varese GC (2009) Isolation and characterisation of
 polychlorinated biphenyl (PCB) degrading fungi from a historically contaminated soil. Microb
 Cell Factories 8:5–5. doi: 10.1186/1475-2859-8-5
- Vaas LAI, Sikorski J, Hofner B, Fiebig A, Buddruhs N, Klenk H-P, Göker M (2013) opm: an R package
 for analysing OmniLog® phenotype microarray data. Bioinformatics 29:1823–1824. doi:
 10.1093/bioinformatics/btt291
- Vaas LAI, Sikorski J, Michael V, Göker M, Klenk H-P (2012) Visualization and curve-parameter
 estimation strategies for efficient exploration of phenotype microarray kinetics. PLoS ONE
 761 7:e34846. doi: 10.1371/journal.pone.0034846
- Valentín L, Lu-Chau TA, López C, Feijoo G, Moreira MT, Lema JM (2007) Biodegradation of
 dibenzothiophene, fluoranthene, pyrene and chrysene in a soil slurry reactor by the white-rot
 fungus *Bjerkandera* sp. BOS55. Process Biochem 42:641–648. doi:
 10.1016/j.procbio.2006.11.011

- Vargas-García M del C, López MJ, Suárez-Estrella F, Moreno J (2012) Compost as a source of microbial
 isolates for the bioremediation of heavy metals: in vitro selection. Sci Total Environ 431:62–67.
 doi: 10.1016/j.scitotenv.2012.05.026
- Vijgen J, Abhilash PC, Li Y, Lal R, Forter M, Torres J, Singh N, Yunus M, Tian C, Schäffer A, Weber R
 (2011) Hexachlorocyclohexane (HCH) as new Stockholm Convention POPs—a global perspective
 on the management of Lindane and its waste isomers. Environ Sci Pollut Res 18:152–162. doi:
 10.1007/s11356-010-0417-9
- Wan J, Meng D, Long T, Ying R, Ye M, Zhang S, Li Q, Zhou Y, Lin Y (2015) Simultaneous removal of
 lindane, lead and cadmium from soils by rhamnolipids combined with citric acid. PLoS ONE
 10:e0129978. doi: 10.1371/journal.pone.0129978
- Wasi S, Tabrez S, Ahmad M (2011) Suitability of immobilized *Pseudomonas fluorescens* SM1 strain for
 remediation of phenols, heavy metals, and pesticides from water. Water Air Soil Pollut 220:89–99.
 doi: 10.1007/s11270-010-0737-x
- Willett KL, Ulrich EM, Hites RA (1998) Differential toxicity and environmental fates of
 hexachlorocyclohexane isomers. Environ Sci Technol 32:2197–2207. doi: 10.1021/es9708530
- Zhu Z, Yang X, Wang K, Huang H, Zhang X, Fang H, Li T, Alva AK, He Z (2012) Bioremediation of CdDDT co-contaminated soil using the Cd-hyperaccumulator *Sedum alfredii* and DDT-degrading
 microbes. J Hazard Mater 235–236:144–151. doi: 10.1016/j.jhazmat.2012.07.033

787 Figures

Fig. 1 Biomineralization by *P. griseofulvum*. Biomineral precipitation observed underneath growing fungal
 colonies in Czapek-Dox agar amended with combinations of 2.5 mM ammonium metavanadate and 2.5
 mM lead carbonate after 3 months incubation of *P. griseofulvum* at 25°C in the dark. Scale bar = 0.4 mm.
 Typical image is shown from several examinations.

Fig. 2 Gas chromatographic analysis of HCH mixture in liquid Czapek-Dox medium after growth of *P*.
 griseofulvum. The HCH mixture was added 26 days after fungal inoculation, and the isomer concentrations

794 were monitored for 23 days. **a** α-HCH concentration. **b** β-HCH concentration. **c** γ-HCH concentration. **d** δ-

HCH concentration. The bars are the standard errors of HCH concentrations of three replicates.

Fig. 3 Proposed biodegradation pathway for HCH in *P. griseofulvum*. The initial reaction is the dehalogenation of HCH to pentachlorocyclohexene, the second is the formation of tetrachlorocyclohexene (TCCH) (3,4,5,6-TCCH), the third is the formation of trichlorobenzene (TCB) (1,2,3-TCB, 1,3,5-TCB and 1,2,4-TCB were detected), the fourth is the formation of dichlorobenzene (DCB) (1,2-DCB, 1,3-DCB and 1,4 DCB were detected), and the fifth is the formation of chlorobenzene.

801 Fig. 4 The parameter "A" (=asymptote) namely the maximum cumulative growth of the fungus on each 802 substrate/in each well, clustered according to the different treatments and visualised as a heat map. The heat 803 map was obtained using the function heat_map of the opm package. The x-axis corresponds to the 804 substrates clustered according to the similarity of their values over all treatments; the y-axis corresponds to 805 the plates clustered according to the similarity of their values over all substrates. The central rectangle is a 806 substrate \times plate matrix in which the colours represent the classes of values. Deep violet to blue indicate 807 low optical density values and light brown indicates high values. The four treatments were cont=control, 808 HCH= hexachlorocyclohexane mixture, V= vanadium and HCH+V= combination of vanadium and HCH 809 for measurements at 750 nm.

Fig. 5 Growth curves of *P. griseofulvum* measured at 750 nm (**a**) and 490 nm (**b**). Growth curves for the four treatments (cont=control; HCH= hexachlorocyclohexane mixture; V= vanadium; HCH+V= combination of vanadium and HCH) for some of the most representative carbon sources among the 95 substrates present in FF arrays were elaborated using the opm package for R.