

Ecological importance of soil bacterivores for ecosystem functions

Jean Trap, Michael Bonkowski, Claude Plassard, Cécile Villenave, Eric

Blanchart

► To cite this version:

Jean Trap, Michael Bonkowski, Claude Plassard, Cécile Villenave, Eric Blanchart. Ecological importance of soil bacterivores for ecosystem functions. Plant and Soil, Springer Verlag, 2015, pp.1-24. <10.1007/s11104-015-2671-6>. <hal-01214705v1>

HAL Id: hal-01214705 https://hal.archives-ouvertes.fr/hal-01214705v1

Submitted on 12 Oct 2015 (v1), last revised 7 Jan 2016 (v2)

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

1	Number of words (main text): 8583
2	Number of words (abstract): 157
3	Number of figures: 7
4	Number of tables: 1
5	Number of appendix: 1
6	
7	Title
8	Ecological importance of soil bacterivores on ecosystem functions
9	
10	Authors
11	Jean Trap ¹ , Michael Bonkowski ² , Claude Plassard ³ , Cécile Villenave ⁴ , Eric Blanchart ¹
12	
13	Affiliations
14	
15	¹ Institut de Recherche pour le Développement – UMR Eco&Sols, 2 Place Viala, 34060,
16	Montpellier, France
17	² Dept. of Terrestrial Ecology, Institut of Zoology, University of Cologne, D-50674 Köln,
18	Germany
19	³ Institut National de Recherche Agronomique – UMR Eco&Sols, 2 Place Viala, 34060,
20	Montpellier, France
21	⁴ ELISOL environnement, 10 avenue du Midi, 30111 Congenies, France
22	

24 Abstract

25 Background

Bacterivores, mostly represented by protists and nematodes, are a key component of soil
biodiversity involved in soil fertility and plant productivity. In the current context of global
change and soil biodiversity erosion, it becomes urgent to suitably recognize and quantify
their ecological importance in the maintenance of ecosystem functions.

30 Scope

Using meta-analysis tools, we aimed at providing a quantitative synthesis of the ecological importance of soil bacterivores on ecosystem functions. We also intended to produce an overview of the ecological factors that are expected to drive the magnitude of bacterivore effects on ecosystem functions.

35 Conclusions

Bacterivores in soil contributed significantly to numerous key ecosystem functions. We
propose a new theoretical framework based on ecological stoichiometry stressing the role of
C:N:P ratios in soil, microbial and plant biomass as important parameters driving bacterivoreeffects on soil N and P availability for plants, immobilization of N and P in the bacterial
biomass, and plant responses in nutrition and growth.

41

42 Keywords

Bacterivores; Protists, Nematodes; Microbial loop; Plant growth; Plant nutrition; Ecological
stoichiometry; Meta-analysis

46 **1. Introduction**

47

Soils are some of the most biologically diverse environments, encompassing about 25% of 48 global biodiversity in terms of species (Coleman and Wall 2015; Decaëns 2010). One gram of 49 soil can host 6000 different bacterial genomes, several hundred meters of fungal hyphae and a 50 wide range of protists, nematodes, enchytraeids or mites (Jones et al. 2009; Lavelle and Spain 51 2001; Wall et al. 2010). Through their activities, soil organisms ensure key soil functions 52 (Barrios 2007; Brussaard et al. 2007; Wagg et al. 2014) and contribute to many ecosystem 53 services (de Vries et al. 2013; Kibblewhite et al. 2008). 54 55 Unfortunately, soils are also among the most threatened environments in terms of biodiversity loss (Kibblewhite 2012). Many pressures (soil erosion, land use change, overexploitation, 56 pollution, biological invasion, etc.) have been identified as directly disturbing soil organism 57 abundance, distribution and activity (Bossio et al. 2005; Dupouey et al. 2002; Foley et al. 58 2005; Gardi et al. 2013; Pimentel 2006; Schlaghamersky et al. 2014). A thorough 59 understanding of the many roles of soil biodiversity is required in order to predict how these 60 threats could damage ecosystem services accomplished by soil organisms. 61 A key component of soil biodiversity involved in soil fertility and plant productivity are 62 63 bacterivores (Bardgett et al. 1999; Bonkowski 2004; Bonkowski et al. 2009; Brussaard 1997). These bacterivores are mostly represented by protists and nematodes (Ronn et al. 2012). 64 Bacterivorous soil protists are generally unicellular heterotrophic eukaryotes, that were 65 66 traditionally grouped into ciliates, flagellates, naked amoebae and testate amoebae

67 (Darbyshire 1994). However, molecular studies reveal an enormous taxonomic diversity in

soils (Geisen et al. 2014; Geisen et al. 2015), and only the ciliated protists are monophyletic,

69 while flagellated, and naked, and testate amoeboid morphotypes repeatedly originated during

70 protist evolution (Adl et al. 2005; Adl et al. 2012).

71	Despite their small size (average size 2-50 μ m), protists can largely contribute to total soil
72	faunal biomass (Bonkowski 2004). Nematodes are small multicellular eukaryotes (0.03-1 mm
73	size range) and form a highly diverse monophyletic group including about one million of
74	species (Lambshead 1993), making this phylum one of the most diverse animal taxa on Earth
75	(Lambshead et al. 2004). Both bacterial-feeding protists and nematodes are widely distributed
76	in terrestrial ecosystems and successful colonizers of soil niches (Cowling 1994; Geisen et al.
77	2014). It has been estimated than tens of millions of protists and hundreds of thousands of
78	nematode individuals co-exist in only one forest square meter (Cowling 1994; Foissner 1999;
79	Yeates 2007; Yeates 2003). Still, they remain poorly studied in comparison to other soil taxa
80	(Coleman and Wall 2015; Couteaux and Darbyshire 1998). Only a fraction of protist and
81	nematode species have been described yet, and their biology and ecology remain poorly
82	known (Brussaard et al. 2007). It thus becomes urgent, in the current context of global change
83	that is expected to alter soil bacterivore communities, to suitably recognize the species and to
84	quantify their ecological importance in the maintenance of ecosystems.
85	Here, as a complement to detailed reviews published in the past (Bonkowski 2004;
86	Bonkowski and Clarholm 2012; Bonkowski et al. 2009; Chen et al. 2007), we aimed at
87	providing a quantitative synthesis of the ecological importance of bacterivores in soil on
88	ecosystem functions using meta-analysis tools. We also intended to produce an overview of
89	the ecological (biotic and abiotic) factors that are expected to drive the magnitude of their
90	effects on ecosystem functions.
91	
92	2. Meta-analysis - Data sources, extraction and analysis

We investigated the literature published in peer-reviewed journals before December 2014
through computer searches in available databases (Google Scholar and ISI Web of

Knowledge), with no restriction on the year of publication, that investigate bacterivore-96 97 induced effects. We used the following key-words (alone or in combination): soil, biodiversity, bacteria, bacterivore, bacterial-feeders/grazers, protozoa, protists, nematodes, 98 predators, mycorrhiza, earthworms, nodules, nutrient, nitrogen, phosphorus, microbial loop. 99 We also searched studies in the reference lists of relevant articles and reviews. We selected 100 studies that (i) tested the effects of bacterivores on ecosystem functions by successfully 101 102 manipulating soil bacterivores in controlled experimental conditions, (ii) measured at least 103 one function of interest, and (iii) reported means of treatment and control. We excluded field studies that monitored soil bacterivore population changes after perturbation or during 104 105 seasonal cycles, or eliminated soil bacterivores by adding biocides in the soil, freezing or sieving the soil without measuring the success of the elimination. We also excluded studies 106 comparing low versus high soil bacterivore densities and modelling studies. 107 108 We analyzed the effects of bacterivores on 18 microbial and plant functions: soil microbial biomass (measured by fumigation or substrate-induced respiration), soil bacterial number 109 (measured by plate counting), soil microbial basal respiration, microbial metabolic quotient 110 (microbial respiration divided by microbial biomass carbon), soil phosphatase activity, soil 111 net N mineralization, soil net P mineralization, plant growth (shoot and root biomass, 112 shoot:root ratio) and plant nutrition (total N and P amounts expressed in mg-N or mg-P plant⁻¹ 113 and concentrations in shoot and root expressed in mg-N or mg-P g⁻¹). For each study, we 114 recorded the mean values of these functions in presence/absence of bacterivores as well as 115 experimental information (soil parameters, species, protocols, etc.). These data were extracted 116 directly from tables, the text or figures using PlotDigitizer 2.6.4 software. 117 For each observation, we calculated the individual effect size (ES_i) from the natural log of the 118 response ratio using Equation 1 (Hedges et al. 1999). 119

Version postprint

- 121 (1) $ES_i = Ln (T_i / C_i)$; where "T" was the treatment mean (presence of bacterivores) and 122 "C" the control mean (absence of bacterivores) for the observation i.
- 123

124 This metric reflects a relative change in a function due to the presence of bacterivores.

125 Positive and negative ES_i indicated respectively a positive and a negative effect of

bacterivores on the function of interest. We then calculated the mean effect size (ES_m) using

the equation 2 and estimated the 95% confidence interval around the ES_m using bootstrapping
(999 iterations).

129

130

(2) $ES_m = (\Sigma ES_i) / n$; with "n" the total number of observations

131

 ES_m was considered significant (P < 0.05) if its 95% confidence intervals did not overlap with 132 zero. For a more intuitive understanding of the effects of bacterivores on functions, we also 133 calculated the percentage of change from control (%CC) from the ES_m. We also tested the 134 effect of the medium used by the experimenters in their studies. To do so, we split the 135 database into 4 groups according to the medium: "agar" (or agarose), "humus", "sand" (mixed 136 with organic matter or with nutrient solution) and "soil" (alone or amended with sand, organic 137 138 matter and/or nutrient solution). We then calculated ES_m for each group and tested for significance among groups using the Kruskal-Wallis test at the P<0.05 level. 139 We also determined whether the presence of mycorrhizal fungi altered bacterivore-effects on 140 plant nutrition and growth. We collected data from studies comparing the effects of 141 bacterivores in the rhizosphere of plants infected or not by mycorrhizal fungi (7 studies, 14 142 observations). We did not include studies where the control (non mycorrhizal plant) was 143 lacking. For each observation, we calculated the ES_i for mycorrhizal and non-mycorrhizal 144

plants and tested for significant difference in ES_m using the Kruskal-Wallis test at the *P*<0.05 level.

Linear meta-regressions were performed to tested the relationships between the bacterivore-147 induced effect size on total N (or P) amount in shoot (or root) (responses variables) and the 148 149 bacterivore-induced effect sizes on shoot (or root) N (or P) concentration and shoot (root) biomass (explanatory variables). 95% confidence intervals were calculated for each 150 151 regression. The normal distribution of residuals was tested using Shapiro test. Finally, we searched for publication bias using funnel plots. We tested the significance (*P*-value < 0.05) 152 of regression between the bacterivore-induced ES versus the inverse sample size of the study 153 154 for each function (Peters et al. 2006). A significant regression indicated that the funnel plot was asymmetric, corresponding to possible publication bias, which was the case only for 155 respiration (*P*-value: 0.02). We also tested the relationships between experiment duration, 156 which ranged from 3 to 561 days, and ES_m values using regression. Except for the 157 concentration of N in roots, we did not find any effect of experiment duration on ES_m. 158

159

Version postprint

160 3. Quantitative synthesis of soil bacterial-bacterivore literature

161

162 We collected 41 experimental studies published since 1977 investigating the effects of soil bacterivores on ecosystem functions (Appendix 1). Among them, 38 and 17 studies 163 investigated effects of bacterivores on N and P concentrations (either in soil or plant tissues), 164 corresponding to ~93% and ~41% of reported studies, respectively (Figure 1). Around 66% of 165 experimental studies co-inoculated bacteria and bacterivores in the rhizosphere of plants, 166 using either herbaceous (73%) or woody (27%) species. According to our literature survey, in 167 a period of 37 years, only 1.1 papers per year on average focused on the ecological functions 168 accomplished by these organisms (without taking into account reviews and books). We 169

170 believe that the number of studies (and observations) is large enough to conduct a meta-

analysis but remains critically limited regarding the significant roles of these organisms onecosystem functioning (Bonkowski 2004; Chen et al. 2007).

This apparent lack of interest in soil bacterivores can be explained by the difficulties involved 173 174 in observing, characterizing (extraction, enumeration and description) and manipulating these small-sized organisms in comparison to larger soil fauna (e.g. arthropods or earthworms) 175 176 (Cowling 1994; Ekelund and Ronn 1994; Griffiths and Ritz 1988). This is particularly true for protists because the majority of taxa cannot be extracted from soils, and our taxonomic 177 knowledge is mostly restricted to few cultivable species (Ekelund and Ronn 1994; Foissner 178 179 1999). Furthermore, this practical difficulty is enhanced by the heterogeneous and opaque nature of soils at the micro-spatial scale in contrast to aquatic environments. Soil 180 protozoology and nematology thus require specific taxonomic skills and time-consuming 181 182 microscopy in addition to expensive equipment (high-quality light-, or electron-microscopy). These methodological drawbacks together with the difficulties working under strictly sterile 183 conditions have certainly constituted the main obstacles in the establishment of soil micro-184 food web experiments. 185

We also noted that contrasting experimental protocols have been used in these studies. For 186 187 instance, experimenters used various types of medium (agar, sand, humus or soil) or experimental devices (Petri dishes, glass tubes, rhizoboxes or pots) for the growth of 188 organisms (Table 1). The amount and composition (C:N:P ratio) of organic amendments and 189 nutrient solutions were also highly different among studies as well as the choice of model 190 species and the duration of the experiment (from 2 to 561 days). This high diversity in 191 protocols made the identification of ecological factors that are likely to drive bacterivore-192 effects on ecosystem functions more difficult. On the other hand, it also allowed us to 193

estimate an overall bacterivore-effect representative of a wide range of environmentalconditions.

196

197 **4.** Bacterivore-effects on the soil microbial community and nutrient availability

198 *4.1. Microbial biomass*

199

200 Because protists and nematodes feed on bacteria, the presence of these organisms usually induced a reduction in soil microbial biomass or bacterial abundance (Anderson et al. 1978; 201 Coleman et al. 1977; Darbyshire et al. 1994; Elliott et al. 1979; Zwart and Darbyshire 1992). 202 203 However, a number of studies showed that the presence of bacterivores can lead to higher bacterial abundance (and biomass) due to a strong reduction of senescent cells (Elliott et al. 204 1980; Ingham et al. 1985; Sundin et al. 1990). The global effect of grazing on the soil 205 206 microbial biomass and bacterial abundance estimated by the meta-analysis was -16% and -17% of control, respectively (Figure 2). 207

208 The effect of bacterivores on bacterial abundance has been shown to vary according to bacterivore species (Ingham et al. 1985; Postma-Blaauw et al. 2005), soil texture (Elliott et al. 209 1980), incubation time (Elliott et al. 1980), supply and availability of labile carbon (Anderson 210 211 et al. 1978; Elliott et al. 1980) or nitrogen (Baath et al. 1981) and identity of bacterial taxa (Glücksman et al. 2010; Griffiths et al. 1999; Ronn et al. 2002; Xiao et al. 2010), showing that 212 differences in bacterivore-effects on bacterial abundance are multifactorial. Certainly the long 213 co-evolution between bacteria and protists had a major influence on the evolution of prey-214 capture mechanisms (Parry 2004) and bacterial defense strategies (Jousset 2011; Matz and 215 Kjelleberg 2005). In consequence, predicting bacterivore-effect on soil bacterial community 216 in a particular environment (or experimental set up) remains difficult. 217

The whole composition of the soil microbial community drastically changed in the presence 221 of protists (Bonkowski et al. 2011; Ekelund et al. 2009; Griffiths et al. 1999; Koller et al. 222 223 2013c; Ronn et al. 2002; Rosenberg et al. 2009) and bacterial-feeding nematodes (Blanc et al. 2006; Djigal et al. 2010; Djigal et al. 2004; Postma-Blaauw et al. 2005). Unfortunately, most 224 225 published studies have a coarse taxonomic resolution limiting our ability to assess accurately grazing effects on microbial composition. Usually, the presence of bacterivores changed the 226 relative abundance of dominant bacterial populations (Djigal et al. 2004), often increasing the 227 228 proportions of bacteria that are grazing protected, either by physical means, such as grampositive bacteria (Griffiths et al. 1999; Ronn et al. 2002; Weekers et al. 1993) and those that 229 make filaments such as actinomycetes (Rosenberg et al. 2009), or by chemical means (Jousset 230 et al. 2009; Jousset et al. 2010; Jousset et al. 2008; Mazzola et al. 2009). The ability of 231 bacterivores to alter the composition of the microbial biomass can feed back on microbial 232 function, such as nitrification (Djigal et al. 2010; Jousset et al. 2006) or phosphatase activity 233 (Djigal et al. 2004; Gould et al. 1979). For instance, using a DGGE technique based on PCR 234 amplification of the amoA gene, Xiao et al. (2010) found a significant shift in the community 235 236 composition of ammonia-oxidizing bacteria (AOB) from *Nitrosospira* sp. to *Nitrosomonas* sp. in presence of nematodes bacterivores. 237

Selective grazing of cells has been proposed as the main mechanism causing a shift in
microbial composition in the presence of protists (Bonkowski et al. 2009). Bacteria evolved
different means to become grazing-resistant, which can be grouped in physical protections
such as bacterial shape and size (Bjornlund et al. 2012), cell wall resistance, and the ability of
cells to form micro-colonies, filaments or biofilms (Bonkowski et al. 2009; Jousset 2011); and
chemical protections, such as the production of pigments (Weekers et al. 1993) or of specific

toxins (Jousset et al. 2006; Jousset et al. 2009; Mazzola et al. 2009). In particular, the latter 244 245 have been identified as main drivers of changes in bacterial community composition, since bacterivores have been shown to consume preferentially the competitors of the grazing-246 resistant taxa (Jousset et al. 2008). The differences in growth rates of bacterivorous protist 247 (Ekelund 1996; Weekers et al. 1993) and nematode populations (Anderson and Coleman 248 1981; Blanc et al. 2006; Venette and Ferris 1998) according to bacterial species suggest that 249 250 also the digestibility and nutritional values of preys constitute an important trait involved in bacteria-bacterivore interactions (Pussard et al. 1994). 251

On the other hand, bacterivores have developed different forms of feeding (direct 252 253 interception, grasping, filter-, diffusion- or raptorial feeding); this appears as an important trait involved in selective grazing aptitude (Parry 2004; Ronn et al. 2012; Weisse 2002). 254 While ciliates move out of toxic biofilms, amoebae graze directly within biofilms and 255 256 grasping and raptorial feeders rather rely on bacteria they can dislocate from biofilm surfaces, while diffusion and filter feeders rely on the free-swimming bacteria (Ronn et al. 2012). In 257 addition, certain bacterivore nematode taxa evolved a grinder in the terminal bulb which 258 serves for crushing trapped bacteria (Fürst von Lieven 2003), thus providing a physical means 259 260 to consume bacteria with thick cell walls that are largely undigestible for other bacterivores, 261 like gram-positive bacteria.

Because bacterial-feeding nematodes can ingest around 20 cells during each suction of their 262 continuously pumping pharynx (Ferris et al. (1997), it appears unlikely that nematodes 263 actively select for specific prey taxa such as protists. Jousset et al. (2009), using mixed 264 populations of mildly toxic wild-type and non-toxic gacS-deficient mutants of Pseudomonas 265 fluorescens CHA0 in batch and rhizosphere systems, showed that the diet composition of the 266 nematode *Caenorhabditis elegans* did not vary with changes in the frequency of the two 267 bacterial strains, confirming its lack in selective compared to Acanthamoeba castellanii that 268

clearly preferred the non-toxic bacterial mutants. Accordingly, bacterial-feeding nematodes 269 270 are more likely to alter bacterial community composition through passive mechanisms. The high variations in the head shape of Cephalobidae species recorded by De Ley (1992) 271 supports this hypothesis. He investigated in detail how the shape of the labial probolae and the 272 development of cephalic probolae at the nematode head allow passive food specialization, 273 thereby optimizing niche partitioning among co-occurring bacterivore nematode species. 274 275 Lastly, bacterivores have been suggested to modify the amount and composition of rhizodeposits (Sundin et al. 1990), which can in turn feed back on the composition of the 276 bacterial community (Benizri et al. 2002; Puglisi et al. 2013). 277

278

279 4.3 Microbial activity and nutrient availability

280

281 The presence of bacterivores generally enhanced the overall microbial activity and turnover (Alphei et al. 1996; Coleman et al. 1978; Djigal et al. 2004; Kuikman et al. 1990). The meta-282 analysis revealed that the presence of bacterivores significantly increased soil microbial basal 283 respiration and microbial turnover, measured as microbial metabolic quotient (respired-carbon 284 relative to microbial biomass carbon) by +29% and 35% of control, respectively (Figure 2), 285 286 indicating that the microbial carbon-use efficiency (CUE) strongly decreases in presence of bacterivores (Manzoni et al. 2012; Sinsabaugh et al. 2013). The main mechanisms are that (i) 287 bacterivores can ingest senescent bacteria and contribute to maintain younger bacteria cells 288 with higher metabolic activity (see discussion in Bonkowski (2004)) and (ii) they release 289 undigested food particles, labile carbon and nutrients making the medium more favorable for 290 bacterial re-growth (Griffiths 1994a; Pussard et al. 1994). 291 The effects of bacterivores on microbial communities have strong consequences for soil 292

nutrient availability (Clarholm 1985a; Koller et al. 2013b; Kuikman and Vanveen 1989; Xiao

et al. 2010). We found that the presence of bacterivores almost doubled soil N (N_{min}) 294 295 mineralization (x 1.8 in absence of plants) (Figure 2). Two main pathways of bacterivoreeffects on soil nutrient availability do exist simultaneously: excretion of nutrients (direct way) 296 according to the consumer-driven nutrient recycling (CNR) theory (Elser and Urabe 1999) 297 and stimulation of the microbial activity and turnover as quoted earlier (indirect way). 298 Using stable isotopes, Crotty et al. (2013) found that soil protists only have a production 299 300 efficiency of 37%. Ferris (1997) found higher production efficiencies (ranging from 58% to 86%) for eight bacterivore nematodes species. Because bacterivores have to maintain 301 stoichiometric homeostasis relative to their food, and loose carbon due to respiration, a great 302 303 part of the ingested N will be ultimately released by bacterivores (Anderson et al. 1983; Darbyshire et al. 1994; Ferris et al. 1997; Ferris et al. 1998; Griffiths 1994b). Borkott (1989) 304 found mass C:N ratios of 3.5:1, 4.7:1 and 5.6:1 for *Naegleria gruberi* (amoeba), *Colpoda* 305 306 steinii (ciliate) and Rhabditis sp. (bacterivore nematode), respectively. Using recent values of global soil microbial stoichiometry (mass C:N ratios ranging from 5.5:1 to 7.4:1) (Cleveland 307 and Liptzin 2007; Xu et al. 2013) and production efficiencies from Crotty et al. (2013) and 308 Ferris et al. (1997), we calculated that theoretical ranges of N excretion would be 23-32%, 43-309 49% and 9-20% of assimilated-N for amoebae, ciliates and nematodes, respectively. The C:N 310 311 ratios of both bacteria and bacterivores will thus directly influence the amount of assimilated-N excreted by the bacterivores (Elser and Urabe 1999). 312 Furthermore, the presence of bacterivores will increase microbial respiration and turnover 313 (Coleman et al. 1978; Levrat et al. 1992), leading to lower microbial CUE and a decreased 314 microbial C:N ratio. In addition, the C:N ratio of bacteria determines the respiration rates in 315 316 presence of bacterivores, with C-losses increasing at decreasing C:N ratios (De Telegdy-Kovats 1932) leading to higher N release by bacterivores at narrow C:N ratios of the 317

318 prey. However, the soil microbial biomass C:N:P has been shown to be relatively stable

319	compared to its resource (Cleveland and Liptzin 2007; Xu et al. 2013); and microbes keep
320	strict homeostasis in respect to nutrient availability in soils (Griffiths et al. 2012). To maintain
321	their stoichiometry, microbes can regulate their nitrogen-use efficiencies by releasing
322	elements in excess depending on their CUE (Mooshammer et al. 2014a). The excess N may
323	then become available for roots in presence of bacterivores (Kuzyakov and Xu 2013).
324	However, small changes of microbial biomass stoichiometry can occur after shifts in the
325	microbial community structure (Fanin et al. 2013). For instance, fast-growing bacteria
326	(copiotrophs) have been suggested to exhibit lower biomass C:N:P ratios (higher nutrient
327	requirements) than slow-growing ones (oligotrophs) (Elser et al. 2003; Hodge et al. 2000).
328	Indeed, in line of the growth rate hypothesis (GRH) (Elser et al. 2003), N:P ratios in organism
329	decrease with increasing growth rates due to elevated demands for P for the synthesis of P-
330	rich ribosomes. Therefore, shifts in the dominance of fast- or slow-growing bacteria induced
331	in presence of bacterivores can transfer to shifts in microbial biomass stoichiometry, with
332	important consequences on the content of N and P available for plant growth.
333	Ferris et al. (1998) calculated that bacterial-feeding nematodes excreted NH ₄ -N with rates
334	ranging between 0.0012 and 0.0058 μ g-N ind. ⁻¹ day ⁻¹ according to species, but in addition,
335	considerable amounts of organic N (Norg) can be released by nematodes (Anderson et al.
336	1983). Wright (1975) found that that 3-14%, 15-34% and 35-45% of the total N (mean 13.7
337	μ moles N g ⁻¹ nematode fresh wt hour ⁻¹) liberated in Ringer solution by the nematode
338	Panagrelus redivivus were composed of urea, amino acids and proteins, respectively, while
339	ammonium contributed 23-35% of total N-release. Large releases of organic N (in total and in
340	proportion of total N) by nematodes may occur when nematodes exhibit high nitrogen-use
341	efficiency (NUE) in order to balance their stoichiometry to bacterial biomass C:N:P ratios,
342	explaining why Anderson et al. (1983) observed declines in excretion of Norg with decreasing
343	resource availability (i.e. when nematode populations became older). Knowing that dissolved

Norg (especially low molecular weight N compounds) can be re-used by soil bacteria or taken 344 345 up by plants (Gallet-Budynek et al. 2009; Nasholm et al. 2009; Neff et al. 2003), bacterivoreeffects on soil N availability likely have been under-estimated because experimenters 346 generally focused only on NH₄. 347 Interestingly, we found a great variability in ES_i for N_{min} as shown by the 95% confidence 348 interval. In parallel, we found that when the substrate used in experiments was soil, the ES_m 349 350 was very high in comparison to humus, while in sand it exhibited intermediate values (Figure 3.A). Humus had the highest mass C:N ratios (mean ~39:1) while soils had the lowest ones 351 (mean ~15:1) and sand exhibited intermediate values (mean ~20:1). We therefore suggest that 352 353 bacterial growth in substrates with high C:N ratios (humus), and in absence of living roots, will be mostly limited by N, leading to rapid immobilization of N released by bacterivores. In 354 contrast, in substrate with low C:N ratios (soil), bacterial growth will be limited by C and 355 356 bacterial N immobilization should decrease (Ferris et al. 1998). We also found that protists exhibited significant higher effects than nematodes on N_{min} but the 357 highest values were found when both bacterivore groups were present in the microcosms 358 (Figure 4.A). Accordingly, the choice in bacterivore species appears as an important 359 experimental parameter explaining the magnitude of bacterivore-effects on N_{min}. The broad 360 361 classification of bacterivores is by far not accurate enough to predict how bacterivores impact soil N cycling. For instance, Ferris et al. (1998) found high variation of different nematodes 362 species to net N mineralization. Predicting bacterivore-effects on nutrient availability 363 therefore remains difficult, especially when a mixture of bacterivore species is used. 364 Applying the same stoichiometric reasoning as for N (Borkott 1989; Cleveland and Liptzin 365 2007; Crotty et al. 2013; Ferris et al. 1997; Xu et al. 2013), theoretical ranges of excess-P 366 excretion would be 27-48%, 73-81% and 66-76% of assimilated-P for amoebae, ciliates and 367 nematodes, respectively. These estimates are higher than for N (except for nematodes) while, 368

in the meta-analysis, we found lower bacterivore-effects on P mineralization (P_{min}) (22% of

370 control) in comparison to N (Figure 2). Why did these theoretical differences in N and P

371 excretion rates of bacterivores not match with N_{min} and P_{min} data provided by the meta-

372 analysis?

The theoretical liberations of N and P by bacterivores did not take into account biotic 373 (microbial immobilization, microbial CUE, resource stoichiometry, etc.) and abiotic processes 374 375 (adsorption of P, leaching of N, etc.) occurring after nutrient excretion by bacterivores, thus differentiating gross from net effects of bacterivores on nutrient availability. In humus (where 376 the C:P ratio is high), the bacterivore-effect on P_{min} was negative, while in soil (where the C:P 377 378 ratio is low) we found a positive effect of bacterivores on P_{min} (Figure 3.B). Hence, in a high C:P ratio environment, bacterial growth may be mostly limited by P and rapid immobilization 379 of excreted P from bacterivores may have occurred. However, in contrast to N, P is poorly 380 381 mobile in the soil solution and can be rapidly adsorbed by soil colloids (Hinsinger et al. 2011), reinforcing rapid immobilization of P in the microbial biomass and hampering the 382 overall positive net effect of bacterivores on Pmin. 383

384

385 5. Bacterivore-effects on plant performance

386 *5.1. Plant growth and root architecture*

387

Positive effect of bacterivores on plant growth have been observed for both herbaceous
(Alphei et al. 1996; Djigal et al. 2004; Krome et al. 2009a) and woody (Bonkowski et al.
2001b; Irshad et al. 2011; Jentschke et al. 1995) plant species (see Table 1 for details). Our
meta-analysis revealed a significant effect of bacterivores on shoot (+27% of control) and root
biomass (+21% of control) without affecting the shoot:root ratio (Figure 2). Interestingly, the
95% confidence interval did not exhibit huge variation though many ecological factors have

been suggested in the literature to affect bacterivore-effects on plant growth. Both, protists and nematodes induced root growth of similar magnitude, but when both were present, their effects were significantly larger (Figure 4.C). We did not find such an additive effect for shoot biomass (Figure 4.B; *P*-value = 0.109).

Besides biomass, the presence of protists in the rhizosphere of plants has been shown to alter 398 profoundly root architecture by promoting lateral root production (Jentschke et al. 1995; 399 400 Kreuzer et al. 2006) (Table 1). Based on 11 studies, we found that bacterivores increased the number of root tips by 96% of control. However, this positive effect was greatly variable 401 among studies (Table 1). Shifts in root architecture were also observed for bacterial-feeding 402 403 nematodes but to a lesser extent than for protists (Cheng et al. 2011; Irshad et al. 2012; Irshad et al. 2011). For instance, while protists increased the number of root tips on average by a 404 factor of 2.5, nematodes only increased the number of tips by a factor of 1.2 (Table 1). The 405 406 presence of mycorrhizal fungi usually reduced the bacterivore effect on root length, specific root length, root area and number of root tips. 407

408

409 *5.2. Plant nutrition*

410

411 The positive effects of bacterivores on plant biomass could be mostly related to increased plant nutrient acquisition when protozoa and nematodes were inoculated in the soil 412 (Bonkowski et al. 2009). Generally, we found substantial bacterivore-effects on the total 413 amount of N in shoots and roots, *i.e.* +59% and +28% of control (Figure 2). Although the 414 increase of plant N concentration in shoots by bacterivores was large (+25% of control) it was 415 highly variable and not different from 0 (non-significant effect on this variable). By 416 conducting linear meta-regressions, we found that $\sim 34\%$ (P<0.001) and $\sim 65\%$ (P<0.001) of 417 the total variance of ES_m on total N amount in shoot (response variable) was explained by 418

bacterivore-effect on shoot biomass and shoot N concentration (explanatory variables), 419 420 respectively, without significant interaction between the two explanatory variables (Figure 5.A). These results imply that in some experimental conditions, bacterivores enhanced the 421 total shoot N amount by increasing the shoot N concentration while in others, bacterivores 422 increased the shoot N amount by increasing shoot biomass, explaining why we found high 423 variance in ES_m for shoot N concentration. Exceptions are the studies by Kuikman et al. 424 425 (1991) and Alphei et al. (1996) where increased plant biomass production in presence of protists or nematodes was neither paralleled by an increased total uptake of plant N, nor by 426 increased concentrations of N. 427

Interestingly, we found higher shoot N concentrations in experiments with bacterial-feeding nematodes than with protists (Figure 4.C). These results suggested that the increase in total shoot N amount was mostly caused by higher N concentrations in plant tissue when nematodes were inoculated. Together with the lower effects of nematodes on root architecture, the data suggest different mechanisms of plant responses to bacterivore nematodes or protists, and might explain why the combined effects of both bacterivore groups were generally larger than the effects of either group alone.

In respect to P, we found that bacterivores increased the total amount of P in shoots and roots 435 436 by 38% and 55% of control, respectively (Figure 2). The data suggest differences in the internal plant cycling of newly-acquired N and P: the bacterivore-induced gain of N in plant 437 tissues was mostly allocated to shoots (probably for the maintenance of high carbon fixation 438 rates via photosynthesis) while the gain of P was more allocated to roots. Krome et al. (2009a) 439 and Koller et al. (2013c) showed that the effects of protozoa on plant performance were more 440 441 related to enhanced plant carbon fixation rather than nitrogen uptake. Compared to N, the effects of bacterivores on shoot P concentration (+30% of control) were less variable and 442 different from 0, suggesting that surplus P made available by bacterivores was commonly 443

concentrated in plant tissues. Although the presence of bacterivores led to an important 444 445 increase in root P concentration (+23% of control), this effect was highly variable. The metaregression showed that ~16% (P<0.001) and ~75% (P<0.001) of the total variance of ES_m on 446 total P amount in roots (response variable) was explained by bacterivore-effects on root 447 biomass and root P concentration (explanatory variables), respectively without significant 448 interaction between these two explanatory variables (Figure 5.B). Some experimental 449 450 conditions, which remain to be known, may thus favor bacterivore-effects on nutrient concentration in plant tissues (without affecting plant biomass) and others may promote 451 bacterivore-effects on plant biomass (and increase total N and P amount in plants). 452

454 5.3. Plant reproduction & defense

456 It is not surprising that positive effects of bacterivores on nutrient uptake and biomass of plants will lead to increased plant reproduction. For instance, the presence of protozoa 457 increased the number of ears (+60%), seeds (24%) and the individual seed weight (+32%) of 458 barley plants, and the crop was even more tolerant to aphid herbivores (Bonkowski et al. 459 2001a). More recently, Krome et al. (2009a) showed that A. castellanii enhanced the seed 460 461 production of Arabidopsis thaliana (Brassicaceae) more than 3- and 7-fold compared to treatments with bacteria and sterile grown plants, respectively. These results imply that gains 462 in plant fitness might be much more profound than increases of plant biomass and/or nutrient 463 contents in tissues indicate. Unfortunately, only these two papers investigated bacterivores 464 effects on plant reproduction. More studies are certainly needed to investigate bacterivore 465 466 effects on plant fitness and on interactions with herbivores and plant pathogens. 467

468 **6.** Relationships between bacterivores and root-infecting symbionts

Version postprint

469 6.1. Bacterivores & Mycorrhizal fungi

The effects of bacterivores on plant functions have been mostly investigated in the
rhizosphere of non-mycorrhizal plants (84% of reported studies) (Figure 1). This is a severe
shortcoming knowing that, in natural conditions, more than 80% of herbaceous and woody
plants are infected by soil fungi, mostly forming arbuscular mycorrhizal (Smith and Read
2008) and ectomycorrhizal symbioses (Marmeisse et al. 2004).

The formation of mycorrhizal roots is considered as the most widespread response of plants to 476 low soil nutrient availability (Plassard et al. 2011). By prospecting a large volume of soil 477 478 away from plant roots, and by releasing carboxylates (Finlay 2008) and enzymes (Courty et al. 2006; Louche et al. 2010), mycorrhizal fungi have been often shown to significantly 479 enhance plant N and P acquisition (Plassard and Dell 2010; Tibbett and Sanders 2002; 480 481 Wallenda and Read 1999). The presence of mycorrhizal fungi can alter the positive effects of bacterivores on plant performance through two main pathways: at a first glance, the presence 482 of mycorrhizal fungi is a priori likely to reduce the positive effect of bacterivores. The growth 483 of bacteria and bacterivores clearly depend on plant carbon allocation to roots. It has been 484 estimated that ca. 7-30% of net carbon fixation will be directly allocated to the root-infecting 485 486 fungal symbionts by the host plant (Jones et al. 2009; Leake et al. 2004). It is likely that the amount of carbon released by root exudation will decrease in the presence of a mycorrhizal 487 partner (Meier et al. 2013; Olsson et al. 1996) with indirect detrimental effects on bacterivore 488 populations. Secondly, mycorrhizal fungi are known to profoundly change root biomass and 489 architecture (Brown et al. 2013). Usually, the presence of the mycorrhizal symbiont reduced 490 the stimulating effect of bacterivores on lateral root production (Table 1). Bacterivore effects 491 on plant nutrition through more complex root architecture can thus be altered by mycorrhizal 492 fungi. 493

However, this *a priori* hypothesis could not be confirmed by the meta-analysis. We did not 494 495 find any negative effect of mycorrhizal fungi on bacterivore effects on N and P concentrations or total amount in plant tissues. In contrast, the ES_m of bacterivores on shoot and root biomass 496 were significantly reduced when the plants were in association with mycorrhizal fungi (Figure 497 6). This negative effect of mycorrhizal fungi on ES_m was very low for shoot biomass (-5%) 498 but substantial for root biomass (-104%), showing that even when mycorrhizal fungi reduced 499 500 the positive effect of bacterivores on root growth, the positive effects of bacterivores on plant nutrition were still maintained. The few existing studies support this hypothesis for both 501 502 plants infected by arbuscular (Koller et al. 2013b; Koller et al. 2013c) and ectomycorrhizal 503 fungi (Bonkowski et al. 2001b; Irshad et al. 2012). These results also support the hypothesis formulated by Bonkowski et al. (2001b) assuming that the mycorrhizal and the bacteria-504 bacterivore mutualisms complement each other and plant resources are allocated to optimize 505 506 simultaneous exploitation of both mutualistic relationships. Because studies are lacking, it was not possible to distinguish the effect of each mycorrhizal fungi type (arbuscular-, or ecto-507 mycorrhiza) on bacterivore-effect sizes. Knowing that both fungal symbionts are 508 phylogenetically and physiologically distinct, this constitutes a serious shortcoming. More 509 studies are required, especially those focusing on different types of mycorrhizal fungi in 510 511 interaction with bacterivores. It is well known that the ability of plants to take up organic N as source of N is greater in 512 presence of mycorrhizal fungi (Cappellazzo et al. 2008; Nasholm et al. 2009; Neff et al. 2003; 513 Plassard et al. 2000; Wallenda and Read 1999). Because significant amounts of dissolved 514

organic N (amino acids) are released by bacterial-feeding nematodes (Anderson et al. 1983;

516 Sundin et al. 1990; Wright 1975), we expect that the presence of mycorrhizal fungi will favor

517 the acquisition of some organic N- (and maybe P-) forms released by bacterivores, before

518 nutrients can be re-used by rhizosphere bacteria. Mycorrhizal plants could thus gain a

519 competitive advantage over microorganisms for soil N released by bacterivores. We did not 520 find experimental study testing this hypothesis, but we believe that this organic N circuit 521 should not be overlooked, and may constitute an important mechanism by which bacterivores 522 improve mycorrhizal plant N nutrition.

523

524 6.2. Bacterivores & N₂-fixing plants

525

Because N usually limits plant productivity, biological N fixation is one of the most important 526 functions of terrestrial ecosystems. Yet, little attention has been given to the potential role of 527 528 bacterivores on symbiotic N fixation and legume growth (Appendix 1). Ramirez and Alexander (1980) showed that protists decreased the abundance of root-nodule bacteria in the 529 530 rhizosphere of *Phaseolus vulgaris*. Lennox and Alexander (1981) found higher numbers of 531 root nodules and growth of *P. vulgaris* when they used a biocide to suppress protists, but we can not judge the side-effects of the biocide, since no protist-specific biocides exist. High 532 symbiotic N fixation efficiency usually requires high levels of available P because high rates 533 of energy transfer take place in the symbiotic nodule (Drevon and Hartwig 1997; Vitousek et 534 al. 2002). The meta-analysis showed that bacterial bacterivores increased soil P availability on 535 536 average by 22% of control. We can hypothesize that protists and nematodes, by increasing soil P availability and subsequent plant P acquisition, might promote higher N fixation 537 efficiency once the nodules are formed. The N:P ratio of nodules would then be an important 538 proxy to predict how bacterivores in the vicinity of nodules may affect plant N uptake. 539 540

541 7. A 'stoichiometric' perspective on bacteria-bacterivore-plant interactions

542 7.1. Stoichiometric controls on bacterivore-effects on soil nutrient availability

544	Comparing the data of bacterivore effects on N- and P-availability point to an important role
545	of stoichiometry if we want to predict how bacterivores impact ecosystem functions.
546	(1) Bacterivores have to maintain stoichiometric homeostasis, but especially protists have
547	been shown to exhibit low production efficiency (Crotty et al. 2013). Accordingly
548	bacterivores release a great part of ingested N and P in mineral form (Elser and Urabe 1999).
549	The amounts of N- and P-release are expected to vary according to bacteria and bacterivores
550	C:N:P ratios. For example, Cole et al. (1978) showed that amoebae were highly efficient at
551	recycling microbially immobilized P to soil in comparison to bacterivore nematodes.
552	(2) Bacterivores often increase microbial turnover and further decrease microbial CUE.
553	Microbes are strongly homeostatic in terms of their biomass C:N:P at the community scale
554	(Cleveland and Liptzin 2007) and can regulate their nutrient-use efficiency by releasing
555	elements in excess depending on their CUE (Mooshammer et al. 2014a) leading to a strict
556	homeostasis with the available N- and P-levels in soil (Griffiths et al. 2012).
557	(3) Bacterivores can alter the soil microbial community structure. According to GRH (Elser et
558	al. 2003), C:N:P ratios in organism vary with growth rates. Therefore, small changes in
559	microbial stoichiometry can occur after shifts in the microbial community structure (Fanin et
560	al. 2013) caused by bacterivores.
561	(4) Soils with high C:N or C:P ratios may be nutrient deficient for bacterial growth, favoring
562	rapid immobilization of newly-mineralized N or P from bacterivores (Ferris et al. 1998). Soils
563	with low C:N or C:P ratios may induce energetic starvation for bacterial growth, reducing
564	bacterial N and P immobilization. The net effects of bacterivores on soil N and P
565	mineralization therefore depend on energy available in terms of labile plant C, either provided
566	by plant roots, or derived during initial phases of litter decomposition (Bonkowski and
567	Clarholm 2012).
568	

Clarholm (1985b) and Bonkowski (2004) proposed theoretical frameworks describing how 571 bacterivores impact plant functions. The 'soil microbial loop' hypothesis (Clarholm 1985a) 572 described nutritional mechanisms, quoted earlier, by which bacterivores may increase plant 573 574 nutrition and growth. Because bacterivores can enhance root growth without increasing plant 575 nutrient concentrations (Alphei et al. 1996; Jentschke et al. 1995; Kuikman et al. 1991), Bonkowski (2004) proposed a 'hormonal' hypothesis based on non-nutritional mechanisms 576 where protists increase lateral root branching by altering microbe-root communication, with 577 578 strong consequences on plant internal auxin metabolism (Krome et al. 2009b). Both nutritional and non-nutritional mechanisms will occur concurrently in the rhizosphere of 579 plants and are not exclusive of each other. However, it is unlikely that they contribute to the 580 581 same magnitude to bacterivore-effects under different experimental conditions. For instance, we showed that the increase in total N amount in shoots induced by bacterivores was either 582 explained by higher shoot N concentration or higher shoot biomass. We found similar 583 partitioning in explanatory variables for bacterivore-effects on total P amount in roots. We 584 believed that ecological stoichiometry can help to understand these patterns. 585 586 Ecological stoichiometry describes the importance of the relative proportions between elements for the growth of organisms (Elser et al. 2000; Gusewell 2004; Sardans et al. 2012; 587 Zechmeister-Boltenstern et al. 2015). Plant C:nutrient ratios are assumed to decrease with 588 increasing plant relative growth rate (Agren 2004; Elser et al. 2000). This was conceptually 589 presented by Ågren (2008) as a linear relationship between these two plant traits (Figure 7.A). 590 From this graph, we defined the "critical ratio" (R_{cri}) which designates the C:nutrient ratio 591 592 above which plant growth is null (nutrient limitation).

For a certain plant species, the major cause of natural high C:nutrient ratios in plant tissues is 593 assumed to be nutrient availability in soil (Ågren 2008). Nutritional mechanisms by 594 bacterivore activity rely on a relatively high release of nutrients from consumed microbial 595 biomass. Such conditions are experimentally met when detritus of narrow C:nutrient ratio is 596 added to soil, supporting high levels of microbial growth and a subsequent substantial release 597 of nutrients by bacterivores (Bonkowski et al. 2000). When the readily available organic C 598 599 from the amendments is depleted, microbial biomass turnover will release nutrients for plant uptake, and nutrient release will be further enhanced in presence of bacterivores. 600 Under more natural soil conditions the nutritional mechanisms should prevail when nutrients 601 602 are poorly available for plants, *i.e.* when competition for N and P between microbes and plants is strong. For example, Koller et al. (2013a) added litter of high C:N ratio (i.e. low 603 quality, LQ) or low C:N ratio (high quality, HQ) to soil inoculated with bacteria or with 604 bacteria and the amoeba A. castellanii, and planted with Plantago lanceolata. By ¹³C-labeling 605 of the plants, they showed that plants in presence of high C:N ratio substrates allocated 12% 606 more recently fixed photosynthates to roots in presence of protists compared to controls. 607 Subsequently, ¹³CO₂ respiration losses from the soil more than doubled, because the excess 608 release of root C could not be fixed in microbial biomass due to nutrient deficiency. Here the 609 610 presence of bacterivores caused a 17% decrease in plant C:N ratio which was likely due to two parallel processes, an increased nitrogen uptake and an excess energy expenditure 611 towards rhizosphere C-allocation. 612 The non-nutritional mechanisms suggest that bacterivores stimulate higher root production 613 and relatively more C-fixation before or without an increase in plant nutrient concentration. 614 The formation of lateral roots and the resulting root branching requires substantial plant 615 investment of photosynthates and nutrients: N for proteins, P for ribosomes and energy 616

617 synthesis. Manipulating nutrient availability to plants through the addition of HQ litter to soil,

Koller et al. (2013a) found a substantially higher increase of root biomass (x1.8) (and 618 619 increased root surface area) compared to shoot biomass (x1.3) in treatments with bacterivores compared to control. In contrast to the LQ treatment, the plant C:N ratio increased by 14%, 620 *i.e.* plant biomass increased while the nutrient concentration decreased. This relatively wider 621 plant C:N ratio might have been a result of a more efficient photosynthesis. At the same time, 622 623 due to greater nutrient availability in low C:N substrates, microbial biomass in the plant rhizosphere incorporated substantially more ¹³C from recent photosynthates. This may be 624 observed when nutrients are highly available for plants, *i.e.* when competition for N and P 625 between microbes and plants is low. 626

627 The relative dominance of nutritional and non-nutritional mechanisms induced by presence of bacterivores can be plotted along a soil C:N and C:P co-gradient likely to drive (i) plant-628 microbe competition towards N and P and (ii) plant stoichiometry status at the beginning or 629 630 during co-inoculation experiments (Figure 7.B). The term *dominance* refers to the relative importance of mechanisms to initiate and drive bacterivore-effects on plants. We 631 distinguished four contrasting cases. In case 1, plants exhibit C:N and C:P ratios above their 632 respective critical ratios, and are strongly limited by N and P (the relative growth rate is 633 therefore null). This case may happen when plants grow in soils with high C:N and C:P ratios. 634 635 Under such soil conditions, microbes are strongly limited by N and P and expected to exhibit high NUE and PUE (Mooshammer et al. 2014b), and consequently the competition for N and 636 P will be shifted in favour of microbes (Kuzyakov and Xu 2013). The experiment of Koller et 637 al. (2013a) cited above, demonstrates that plants under nutrient limitation will allocate 638 relatively more photosynthates belowground, thus providing energy for potential root 639 symbionts or priming effects on soil organic matter. Under these circumstances the 640 consumption of microbes by bacterivores will lead to increased microbial turnover and a 641 decreased microbial CUE. Plants will primarily benefit from the released N and P by 642

bacterivores. In cases 2a and 2b, plants have C:N or C:P ratios just below their respective 643 644 critical ratios; they are thus strongly limited by N or P according to soil C:N:P ratios. Similar mechanisms as described above may occur for other plant limiting nutrient concentrations, 645 but according to plant species stoichiometric behavior, an excess uptake of the non-limiting 646 nutrient may also occur (Ågren 2008). In cases 3a and 3b, plants grow in soils with 647 intermediate C:N or C:P ratios, respectively. Microbes become limited by carbon and their 648 649 NUE and PUE may decrease, while plants are limited by N or P but their C:nutrient ratios are below the critical ratios. The presence of bacterivores, and the subsequent changes in nutrient 650 availability and root architecture, may result in both nutrient storage and biomass production, 651 652 according to species stoichiometric behavior (co-dominance of mechanisms). In case 4, microbes are strongly limited by carbon (NUE and PUE are expected to be low) and plants 653 exhibit low C:N and C:P ratios. This may happen in substrates with low C:N and C:P ratios 654 655 (or soils with low availability of labile C). Plant nutrient concentrations are high enough to cause an increase of plant biomass (dilution of plant nutrient concentration can occur), but 656 also lateral root production (i.e. root surface area) for more efficient nutrient uptake in 657 response to bacterivores (dominance of non-nutritional mechanisms). This case is expected to 658 happen when plants grow in a relatively nutrient-rich environment (Jentschke et al. 1995), or 659 660 when plants experience nutrient imbalances which is critical, especially at early life stages (Fenner and Lee 1989; Hanley and Fenner 1997). 661

662

663 8. Conclusion & Perspectives

664

In this meta-analysis, we showed that bacterivores in soil contributed significantly to
numerous key ecosystem functions. We suggest a theoretical framework based on ecological
stoichiometry stressing the role of C:N:P ratios in soil, microbial and plant biomass as

important parameters driving bacterivore-effects on soil N and P availability for plants, 668 669 immobilization of N and P in the bacterial biomass, and plant responses in nutrition and growth. Unfortunately, it has not yet been possible to test this 'stoichiometric' hypothesis 670 using meta-analysis tools because of insufficient data on soil and plant C:N:P ratios. More co-671 inoculation experiments focusing on bacterivore-effects on plant nutrient acquisition in 672 673 various environmental conditions are therefore needed. In addition, we propose three outlooks 674 to contribute to our understanding of bacterivore functions in terrestrial ecosystems. From population to community-level experiments – Studies investigating the effects of 675 bacterivores on ecosystem functions usually used one model species. In this meta-analysis, we 676 677 found that 57% of studies focusing on protists effects selected the A. castellanii species as a model, corresponding to 28% of all studies reported in the meta-analysis. This species was 678 frequently used because it has a ubiquitous worldwide distribution (Geisen et al. 2014) and 679 680 has strong effects on the activity and composition of the soil microbial community (Griffiths et al. 1999; Kreuzer et al. 2006; Rosenberg et al. 2009) and on root architecture (Bonkowski 681 and Clarholm 2012). Concerning bacterial-feeding nematodes, the genera Mesodiplogaster, 682 Rhabditis, Acrobeloides (and several other Cephalobidae genera), were frequently used. On 683 average, we found that 38% of studies used a mixture of different protists or nematodes 684 685 species. Only 6 studies (15%) investigated soil protists and nematodes together and only 2 of them did it in the rhizosphere of a plant (Figure 1). 686

It is recognized that the effects of bacterivores on microbial community composition (Djigal
et al. 2004; Ronn et al. 2002), soil N mineralization (Ferris et al. 1998) or plant performance
(Bonkowski and Clarholm 2012; Cheng et al. 2011; Irshad et al. 2011) are species-specific.
Also, biological interactions among bacterivores are diverse and often antagonistic (Ronn et
al. 2012). Recent studies indicate that intraguild-predation between 'bacterivores', especially
antagonistic interactions between protists and nematodes are quite common and not

unidirectional. Bacterivore nematodes are known to prey on protists (Anderson et al. 1978; 693 694 Bonkowski et al. 2000; Neidig et al. 2010; Ronn et al. 2012) and vice-versa (Bjornlund and Ronn 2008; Neidig et al. 2010). The population density of one bacterivore taxon can thus 695 increase on the cost of other another bacterivore (Anderson and Coleman 1981). 696 Unfortunately, the effect of within-trophic group diversity of the soil bacterivore community 697 698 on ecosystem function has rarely been investigated (Postma-Blaauw et al. 2005; Saleem et al. 699 2012). As a consequence, we are still far from predicting which bacterivore community 700 attribute(s) drive(s) the overall effect of the bacterivore community on ecosystem functions. A number of community attributes might potentially drive the bacterivore community effects, 701 702 such as (i) species richness, (ii) keystone groups with specific ecology (*e.g.* amoebae), (iii) keystone species (e.g. Acanthamoeba castellanii), (iv) particular trait dissimilarity (head 703 704 morphology, feeding types, body size, demographic parameters; etc.) or (v) multi-trait 705 dissimilarity (Rao's dissimilarity coefficient). Identifying the most relevant attributes of soil bacterivore communities driving ecosystem functions should help us to get closer to 706 predicting the actual effects of bacterial bacterivores in the fields. 707 Kinetic experiments to monitor bacterivore-induced effects on plant performance – Most 708 studies investigated the effect of bacterivores on plant performance after a specific period of 709 710 growth. Kinetic experiments are scarce because monitoring bacterivore-effects on plant growth or nutrition requires a high number of replicates, which is difficult from a technical 711 point of view. However, this type of studies might provide crucial information on the 712 sequence of plant and microbial responses in the presence of bacterivores, and the dynamic 713 and persistence of bacterivore-effects on ecosystem functions (Krome et al. 2009a). For 714 715 instance, Ingham et al. (1985) observed a temporary effect of nematodes (*Pelodera* sp. and Acrobeloides sp.) on plant (Bouteloua gracilis) growth. Kinetic experiments conducted under 716 contrasting soil N and P availability would be optimal to test the *stoichiometric* hypothesis. 717

Emphasize on P and ecological factors – P is essential for plant growth and may be limiting 718 719 in many environments. In contrast to N, P is characterized by its relative immobility in soil and the very low concentration of its available form in the soil solution (Hinsinger 2001; 720 721 Hinsinger et al. 2011). Here, we found that N was 2.3-times more studied than P in both soil and plant tissues. This is a serious deficiency given that plant P acquisition efficiency is often 722 based on biological interactions between plant roots and rhizosphere organisms. Using P 723 724 radio-isotopes is probably the best way to better understand how bacterivores affect P cycling 725 at the soil-plant interface. Also, the literature indicates that a large number of other ecological factors are likely to impact bacterivore-effects on ecosystem functions: bacterivore species 726 727 (Cheng et al. 2011; Ferris et al. 1997), soil properties (Ekelund and Ronn 1994), plant species, or cultivars (Somasundaram et al. 2008), the presence of symbiotic mutualists (Herdler et al. 728 2008; Irshad et al. 2012) or other free fauna (Bonkowski and Schaefer 1997; Tao et al. 2009), 729 730 time after inoculation (Krome et al. 2009a), etc. Unfortunately, the number of observations for each factor was too low or difficult to aggregate to perform a meta-analysis. For instance, 731 it was impossible to test the effect of initial bacterivore biomass (or density), soil properties 732 (pH, texture, carbon content, water content; etc.), plant species or cultivar, or the presence of 733 rhizosphere organisms (e.g. mycorrhizal fungi or rhizobia) on bacterivore-induced effects on 734 735 ecosystem functions. Conducting more detailed studies on bacteria/root/bacterivore interactions should deeply improve our understanding of soil food web roles in ecosystems 736 functions, and ultimately help us to predict how the erosion of soil biodiversity will affect 737 738 ecosystem services in the future.

739

Version postprint

740 Acknowledgements

741

742 We thank Claire Marsden for providing helpful comments on an earlier draft of this review.743

744 References

Adl SM, Simpson AG, Farmer MA, Andersen RA, Anderson OR, Barta JR, Bowser SS, 746 747 Brugerolle G, Fensome RA, Fredericq S (2005) The new higher level classification of eukaryotes with emphasis on the taxonomy of protists. Journal of Eukaryotic 748 Microbiology 52: 399-451. 749 750 Adl SM, Simpson AG, Lane CE, Lukeš J, Bass D, Bowser SS, Brown MW, Burki F, 751 Dunthorn M, Hampl V (2012) The revised classification of eukaryotes. Journal of Eukaryotic Microbiology 59: 429-514. 752 753 Agren GI (2004) The C:N:P stoichiomestry of autotrophs - theory and observations. Ecol Lett 7: 185-191. 754 Ågren GI (2008) Stoichiometry and nutrition of plant growth in natural communities. Annual 755 756 review of ecology, evolution, and systematics: 153-170. Alphei J, Bonkowski M, Scheu S (1996) Protozoa, Nematoda and Lumbricidae in the 757 rhizosphere of Hordelymus europeaus (Poaceae): faunal interactions, response of 758 microorganisms and effects on plant growth. Oecologia 106: 111-126. 759 Anderson R, Gould W, Woods L, Cambardella C, Ingham R, Coleman D (1983) Organic and 760 761 inorganic nitrogenous losses by microbivorous nematodes in soil. Oikos: 75-80. Anderson RV, Coleman DC (1981) Population development and interactions between 2 762 species of bacteriophagic nematodes. Nematologica 27: 6-19. 763 Anderson RV, Elliott ET, McClellan JF, Coleman DC, Cole CV, Hunt HW (1978) Trophic 764 interactions in soils as they affect energy and nutrient dynamics. 3. Biotic interactions 765 766 of bacteria, amebas, and nematodes. Microb Ecol 4: 361-371.

767	Baath E, Lohm U, Lundgren B, Rosswall T, Soderstrom B, Sohlenius B (1981) Impact of
768	microbial-feeding animals on total soil activity and nitrogen dynamics - A soil
769	microcosm experiment. Oikos 37: 257-264.
770	Bardgett RD, Cook R, Yeates GW, Denton CS (1999) The influence of nematodes on below-
771	ground processes in grassland ecosystems. Plant and Soil 212: 23-33.
772	Barrios E (2007) Soil biota, ecosystem services and land productivity. Ecol Econ 64: 269-285.
773	Benizri E, Dedourge O, Dibattista-Leboeuf C, Piutti S, Nguyen C, Guckert A (2002) Effect of
774	maize rhizodeposits on soil microbial community structure. Applied Soil Ecology 21:
775	261-265.
776	Bjornlund L, Liu MQ, Ronn R, Christensen S, Ekelund F (2012) Nematodes and protozoa
777	affect plants differently, depending on soil nutrient status. European Journal of Soil
778	Biology 50: 28-31.
779	Bjornlund L, Ronn R (2008) 'David and Goliath' of the soil food web - Flagellates that kill
780	nematodes. Soil Biology & Biochemistry 40: 2032-2039.
781	Blanc C, Sy M, Djigal D, Brauman A, Normand P, Villenave C (2006) Nutrition on bacteria
782	by bacterial-feeding nematodes and consequences on the structure of soil bacterial
783	community. European Journal of Soil Biology 42: S70-S78.
784	Bonkowski M (2004) Protozoa and plant growth: the microbial loop in soil revisited. New
785	Phytologist 162: 617-631.
786	Bonkowski M, Clarholm M (2012) Stimulation of plant growth through interactions of
787	bacteria and protozoa: Testing the auxiliary microbial loop hypothesis. Acta
788	Protozoologica 51: 237-247.
789	Bonkowski M, Geoghegan IE, Birch ANE, Griffiths BS (2001a) Effects of soil decomposer
790	invertebrates (protozoa and earthworms) on an above-ground phytophagous insect
791	(cereal aphid) mediated through changes in the host plant. Oikos 95: 441-450.

792	Bonkowski M, Griffiths B, Scrimgeour C (2000) Substrate heterogeneity and microfauna in
793	soil organic 'hotspots' as determinants of nitrogen capture and growth of ryegrass.
794	Applied Soil Ecology 14: 37-53.
795	Bonkowski M, Jentschke G, Scheu S (2001b) Contrasting effects of microbial partners in the
796	rhizosphere: interactions between Norway Spruce seedlings (Picea abies Karst.),
797	mycorrhiza (Paxillus involutus (Batsch) Fr.) and naked amoebae (protozoa). Applied
798	Soil Ecology 18: 193-204.
799	Bonkowski M, Koller R, Jousset AJF (2011) How protozoa structure microbial communities
800	in the rhizosphere of plants. Journal Of Phycology 47: S7-S7.
801	Bonkowski M, Schaefer M (1997) Interactions between earthworms and soil protozoa: A
802	trophic component in the soil food web. Soil Biology & Biochemistry 29: 499-502.
803	Bonkowski M, Villenave C, Griffiths B (2009) Rhizosphere fauna: the functional and
804	structural diversity of intimate interactions of soil fauna with plant roots. Plant and
805	Soil 321: 213-233.
806	Borkott H (1989) Elementgehalte (C, N, P, K) wirbelloser Bodentiere. Zeitschrift für
807	Pflanzenernährung und Bodenkunde 152: 77-80. doi: 10.1002/jpln.19891520114.
808	Bossio D, Girvan MS, Verchot L, Bullimore J, Borelli T, Albrecht A, Scow K, Ball AS,
809	Pretty J, Osborn AM (2005) Soil microbial community response to land use change in
810	an agricultural landscape of western Kenya. Microb Ecol 49: 50-62.
811	Brown LK, George TS, Barrett GE, Hubbard SF, White PJ (2013) Interactions between root
812	hair length and arbuscular mycorrhizal colonisation in phosphorus deficient barley
813	(Hordeum vulgare). Plant and Soil 372: 195-205.
814	Brussaard L (1997) Biodiversity and ecosystem functioning in soil. Ambio: 563-570.
815	Brussaard L, De Ruiter PC, Brown GG (2007) Soil biodiversity for agricultural sustainability.
816	Agriculture, ecosystems & environment 121: 233-244.

817	Cappellazzo G, Lanfranco L, Fitz M, Wipf D, Bonfante P (2008) Characterization of an
818	amino acid permease from the endomycorrhizal fungus Glomus mosseae. Plant
819	Physiol 147: 429-437.
820	Chen X, Liu M, Hu F, Mao X, Li H (2007) Contributions of soil micro-fauna (protozoa and
821	nematodes) to rhizosphere ecological functions. Acta Ecologica Sinica 27: 3132-3143.
822	Cheng YH, Jiang Y, Griffiths BS, Li DM, Hu F, Li HX (2011) Stimulatory effects of
823	bacterial-feeding nematodes on plant growth vary with nematode species. Nematology
824	13: 369-372.
825	Clarholm M (1985a) Interactions of bacteria, protozoa and plants leading to mineralization of
826	soil-nitrogen. Soil Biology & Biochemistry 17: 181-187.
827	Clarholm M (1985b) Possible roles for roots, bacteria, protozoa, and fungi in supplying
828	nitrogen to plants. In: AH Fitter (ed) Ecological interactions in soil. Blackwell Scient.
829	Publ.
830	Cleveland CC, Liptzin D (2007) C: N: P stoichiometry in soil: is there a "Redfield ratio" for
831	the microbial biomass? Biogeochemistry 85: 235-252.
832	Cole CV, Elliott ET, Hunt HW, Coleman DC (1978) Trophic interactions in soils as they
833	affect energy and nutrient dynamics. Phosphorus transformations. Microb Ecol 4: 381-
834	387.
835	Coleman DC, Anderson RV, Cole CV, Elliott ET, Woods L, Campion MK (1978) Trophic
836	interactions in soils as they affect energy and nutrient dynamics. Flows of metabolic
837	and biomass carbon. Microb Ecol 4: 373-380.
838	Coleman DC, Cole CV, Anderson RV, Blaha M, Campion MK, M. C, Elliott ET, Hunt HW,
839	Shaefer B, Sinclair J (1977) An analysis of rhizosphere-saprophage interactions in
840	terrestrial ecosystems. Ecological Bulletins 25: 299-309.

- Coleman DC, Wall DH (2015) Soil fauna: occurrence, biodiversity, and roles in ecosystem
 function. In: EA Paul (ed) Soil Microbiology, Ecology and Biochemistry (Fourth
 Edition). Academic Press, Boston.
- Courty PE, Pouysegur R, Buée M, Garbaye J (2006) Laccase and phosphatase activities of the
 dominant ectomycorrhizal types in a lowland oak forest. Soil Biology & Biochemistry
 38: 1219-1222.
- 847 Couteaux MM, Darbyshire JF (1998) Functional diversity amongst soil protozoa. Applied
 848 Soil Ecology 10: 229-237.
- Cowling AJ (1994) Protozoa distribution and adaptation. In: JF Darbyshire (ed) Soil protozoa.
 CAB International, London.
- Crotty FV, Adl SM, Blackshaw RP, Murray PJ (2013) Measuring soil protist respiration and
 ingestion rates using stable isotopes. Soil Biology and Biochemistry 57: 919-921.
- 853 Darbyshire JF (1994) Soil protozoa. Cab International.
- Barbyshire JF, Davidson MS, Chapman SJ, Ritchie S (1994) Excretion of nitrogen and
 phosphorus by the soil Ciliate *Colpoda steinii* when fed the soil bacterium
- 856 *Arthrobacter sp.* Soil Biology & Biochemistry 26: 1193-1199.
- B57 De Ley P (1992) The nematode community of a marginal soil at Camberene, Senegal, with
- special attention to functional morphology and niche partitioning in the family
- 859 Cephalobidae. Mededelingen van de Koninklijke Academie voor Wetenschappen,
- Letteren en Schone Kunsten van België, Klasse der Wetenschappen 53: 109-153.
- B61 De Telegdy-Kovats L (1932) The growth and respiration of bacteria in sand cultures in the
- presence and absence of protozoa. Annals of Applied Biology 19: 65-86.
- de Vries FT, Thebault E, Liiri M, Birkhofer K, Tsiafouli MA, Bjornlund L, Jorgensen HB,
- Brady MV, Christensen S, de Ruiter PC, d'Hertefeldt T, Frouz J, Hedlund K, Hemerik
- L, Hol WHG, Hotes S, Mortimer SR, Setala H, Sgardelis SP, Uteseny K, van der

866	Putten WH, Wolters V, Bardgett RD (2013) Soil food web properties explain
867	ecosystem services across European land use systems. Proceedings Of The National
868	Academy Of Sciences Of The United States Of America 110: 14296-14301.
869	Decaëns T (2010) Macroecological patterns in soil communities. Global Ecology And
870	Biogeography 19: 287-302.
871	Djigal D, Baudoin E, Philippot L, Brauman A, Villenave C (2010) Shifts in size, genetic
872	structure and activity of the soil denitrifier community by nematode grazing. European
873	Journal of Soil Biology 46: 112-118.
874	Djigal D, Brauman A, Diop TA, Chotte JL, Villenave C (2004) Influence of bacterial-feeding
875	nematodes (Cephalobidae) on soil microbial communities during maize growth. Soil
876	Biology & Biochemistry 36: 323-331.
877	Drevon J-J, Hartwig UA (1997) Phosphorus deficiency increases the argon-induced decline of
878	nodule nitrogenase activity in soybean and alfalfa. Planta 201: 463-469.
879	Dupouey J-L, Dambrine E, Laffite J-D, Moares C (2002) Irreversible impact of past land use
880	on forest soils and biodiversity. Ecology 83: 2978-2984.
881	Ekelund F (1996) Growth kinetics of five common heterotrophic soil flagellates. European
882	journal of soil biology 32: 15-24.
883	Ekelund F, Ronn R (1994) Notes on protozoa in agricultural soil with emphasis on
884	heterotrophic flagellates and naked amebas and their ecology. Fems Microbiology
885	Reviews 15: 321-353.
886	Ekelund F, Saj S, Vestergard M, Bertaux J, Mikola J (2009) The "soil microbial loop" is not
887	always needed to explain protozoan stimulation of plants. Soil Biology &
888	Biochemistry 41: 2336-2342.

Comment citer ce document : Trap, J., Bonkowski, M., Plassard, C., Villenave, C., Blanchart, E. (2015). Ecological importance of soil bacterivores for ecosystem functions. Plant and Soil, 1-24. DOI : 10.1007/s11104-015-2671-6

- Elliott E, Cole C, Coleman D, Anderson R, Hunt H, McClellan J (1979) Amoebal growth in
 soil microcosms: a model system of C, N, and P. Trophic dynamics. International
 Journal of Environmental Studies 13: 169-174.
- Elliott ET, Anderson RV, Coleman DC, Cole CV (1980) Habitable pore-space and microbial
 trophic interactions. Oikos 35: 327-335.
- Elser J, Acharya K, Kyle M, Cotner J, Makino W, Markow T, Watts T, Hobbie S, Fagan W,
 Schade J (2003) Growth rate–stoichiometry couplings in diverse biota. Ecol Lett 6:
 936-943.
- Elser J, Sterner R, Gorokhova E, Fagan W, Markow T, Cotner J, Harrison J, Hobbie S, Odell
 G, Weider L (2000) Biological stoichiometry from genes to ecosystems. Ecol Lett 3:
 540-550.
- Elser JJ, Urabe J (1999) The stoichiometry of consumer-driven nutrient recycling: theory,
 observations, and consequences. Ecology 80: 735-751.
- 902 Fanin N, Fromin N, Buatois B, Hattenschwiler S (2013) An experimental test of the
- 903 hypothesis of non-homeostatic consumer stoichiometry in a plant littermicrobe904 system. Ecol Lett 16: 764-772.
- Fenner M, Lee W (1989) Growth of seedlings of pasture grasses and legumes deprived of
 single mineral nutrients. J Appl Ecol: 223-232.
- 907 Ferris H, Venette RC, Lau SS (1997) Population energetics of bacterial-feeding nematodes:
 908 Carbon and nitrogen budgets. Soil Biology & Biochemistry 29: 1183-1194.
- 909 Ferris H, Venette RC, van der Meulen HR, Lau SS (1998) Nitrogen mineralization by
- 910 bacterial-feeding nematodes: verification and measurement. Plant and Soil 203: 159-

911 171.

Finlay RD (2008) Ecological aspects of mycorrhizal symbiosis: with special emphasis on the
 functional diversity of interactions involving the extraradical mycelium. Journal of

914 Experimental Botany.

- Foissner W (1999) Soil protozoa as bioindicators: pros and cons, methods, diversity,
 representative examples. Agric Ecosyst Environ 74: 95-112.
- 917 Foley JA, DeFries R, Asner GP, Barford C, Bonan G, Carpenter SR, Chapin FS, Coe MT,

918 Daily GC, Gibbs HK (2005) Global consequences of land use. science 309: 570-574.

- Fürst von Lieven A (2003) Functional morphology and evolutionary origin of the three-part
 pharynx in nematodes. Zoology 106: 183-201.
- 921 Gallet-Budynek A, Brzostek E, Rodgers VL, Talbot JM, Hyzy S, Finzi AC (2009) Intact

amino acid uptake by northern hardwood and conifer trees. Oecologia 160: 129-138.

- Gardi C, Jeffery S, Saltelli A (2013) An estimate of potential threats levels to soil biodiversity
 in EU. Global Change Biology 19: 1538-1548.
- Geisen S, Fiore-Donno AM, Walochnik J, Bonkowski M (2014) Acanthamoeba everywhere:
 high diversity of Acanthamoeba in soils. Parasitology Research 113: 3151-3158.
- 927 Geisen S, Tveit AT, Clark IM, Richter A, Svenning MM, Bonkowski M, Urich T (2015)

928 Metatranscriptomic census of active protists in soils. The ISME journal.

- Glücksman E, Bell T, Griffiths RI, Bass D (2010) Closely related protist strains have different
 grazing impacts on natural bacterial communities. Environ Microbiol 12: 3105-3113.
- Gould WD, Coleman DC, Rubink AJ (1979) Effect of bacteria and amoebae on rhizosphere
- 932 phosphatase activity. Appl Environ Microbiol 37: 943-946.
- 933 Griffiths B (1994a) Soil nutrient flow. In: JF Darbyshire (ed) Soil protozoa. CAB

934 International, London.

- 935 Griffiths BS (1994b) Soil nutrient flow. In: J Darbyshire (ed) Soil Protozoa. CAB
- 936 International, Wallingford.

- Griffiths BS, Bonkowski M, Dobson G, Caul S (1999) Changes in soil microbial community
 structure in the presence of microbial-feeding nematodes and protozoa. Pedobiologia
 43: 297-304.
- Griffiths BS, Ritz K (1988) A technique to extract, enumerate and measure protozoa from
 mineral soils. Soil Biology & Biochemistry 20: 163-173.
- Griffiths BS, Spilles A, Bonkowski M (2012) C: N: P stoichiometry and nutrient limitation of
 the soil microbial biomass in a grazed grassland site under experimental P limitation
 or excess. Ecological Processes 1: 1-11.
- Gusewell S (2004) N : P ratios in terrestrial plants: variation and functional significance. New
 Phytologist 164: 243-266.
- Hanley M, Fenner M (1997) Seedling growth of four fire-following Mediterranean plant
 species deprived of single mineral nutrients. Funct Ecol 11: 398-405.
- Hedges LV, Gurevitch J, Curtis PS (1999) The meta-analysis of response ratios in
 experimental ecology. Ecology 80: 1150-1156.
- 951 Herdler S, Kreuzer K, Scheu S, Bonkowskia M (2008) Interactions between arbuscular
- 952 mycorrhizal fungi (*Glomus intraradices*, Glomeromycota) and amoebae
- 953 (Acanthamoeba castellanii, Protozoa) in the rhizosphere of rice (Oryza sativa). Soil
- Biology & Biochemistry 40: 660-668.
- Hinsinger P (2001) Bioavailability of soil inorganic P in the rhizosphere as affected by rootinduced chemical changes: a review. Plant And Soil 237: 173-195.
- 957 Hinsinger P, Brauman A, Devau N, Gerard F, Jourdan C, Laclau J-P, Le Cadre E, Jaillard B,
- 958 Plassard C (2011) Acquisition of phosphorus and other poorly mobile nutrients by
- roots. Where do plant nutrition models fail? Plant And Soil 348: 29-61.
- 960 Hodge A, Robinson D, Fitter A (2000) Are microorganisms more effective than plants at
- 961 competing for nitrogen? Trends in Plant Science 5: 304-308.

- Ingham RE, Trofymow J, Ingham ER, Coleman DC (1985) Interactions of bacteria, fungi, and
 their nematode grazers: effects on nutrient cycling and plant growth. Ecol Monogr 55:
 119-140.
- 965 Irshad U, Brauman A, Villenave C, Plassard C (2012) Phosphorus acquisition from phytate
 966 depends on efficient bacterial grazing, irrespective of the mycorrhizal status of Pinus
 967 pinaster. Plant and Soil 358: 148-161.
- 968 Irshad U, Villenave C, Brauman A, Plassard C (2011) Grazing by nematodes on rhizosphere
 969 bacteria enhances nitrate and phosphorus availability to *Pinus pinaster* seedlings. Soil
 970 Biology & Biochemistry 43: 2121-2126.
- Jentschke G, Bonkowski M, Godbold DL, Scheu S (1995) Soil protozoa and forest tree
 growth Non-nutritional effects and interaction with mycorrhizae. Biology and
 Fertility of Soils 20: 263-269.
- Jones DL, Nguyen C, Finlay RD (2009) Carbon flow in the rhizosphere: carbon trading at the
 soil-root interface. Plant And Soil 321: 5-33.
- Jousset A (2011) Ecological and evolutive implications of bacterial defences against
 predators. Environ Microbiol 14: 1830-1843.
- Jousset A, Lara E, Wall LG, Valverde C (2006) Secondary metabolites help biocontrol strain
 Pseudomonas fluorescens CHA0 to escape protozoan grazing. Appl Environ
 Microbiol 72: 7083-7090.
- Jousset A, Rochat L, Péchy-Tarr M, Keel C, Scheu S, Bonkowski M (2009) Predators
- 982 promote defence of rhizosphere bacterial populations by selective feeding on non983 toxic cheaters. The ISME journal 3: 666-674.
- Jousset A, Rochat L, Scheu S, Bonkowski M, Keel C (2010) Predator-prey chemical warfare
- 985 determines the expression of biocontrol genes by rhizosphere-associated *Pseudomonas*
- 986 *fluorescens*. Appl Environ Microbiol 76: 5263-5268.

987	Jousset A, Scheu S, Bonkowski M (2008) Secondary metabolite production facilitates
988	establishment of rhizobacteria by reducing both protozoan predation and the
989	competitive effects of indigenous bacteria. Funct Ecol 22: 714-719.
990	Kibblewhite MG (2012) Definition of priority areas for soil protection at a continental scale.
991	Soil Use And Management 28: 128-133.
992	Kibblewhite MG, Ritz K, Swift MJ (2008) Soil health in agricultural systems. Philosophical
993	Transactions Of The Royal Society B-Biological Sciences 363: 685-701.
994	Koller R, Robin C, Bonkowski M, Ruess L, Scheu S (2013a) Litter quality as driving factor
995	for plant nutrition via grazing of protozoa on soil microorganisms. FEMS
996	microbiology ecology 85: 241-250.
997	Koller R, Rodriguez A, Robin C, Scheu S, Bonkowski M (2013b) Protozoa enhance foraging
998	efficiency of arbuscular mycorrhizal fungi for mineral nitrogen from organic matter in
999	soil to the benefit of host plants. New Phytologist 199: 203-211.
1000	Koller R, Scheu S, Bonkowski M, Robin C (2013c) Protozoa stimulate N uptake and growth
1001	of arbuscular mycorrhizal plants. Soil Biology and Biochemistry 65: 204.
1002	Kreuzer K, Adamczyk J, Iijima M, Wagner M, Scheu S, Bonkowski M (2006) Grazing of a
1003	common species of soil protozoa (Acanthamoeba castellanii) affects rhizosphere
1004	bacterial community composition and root architecture of rice (Oryza sativa L.). Soil
1005	Biology & Biochemistry 38: 1665-1672.
1006	Krome K, Rosenberg K, Bonkowski M, Scheu S (2009a) Grazing of protozoa on rhizosphere
1007	bacteria alters growth and reproduction of Arabidopsis thaliana. Soil Biology &
1008	Biochemistry 41: 1866-1873.
1009	Krome K, Rosenberg K, Dickler C, Kreuzer K, Ludwig-Muller J, Ullrich-Eberius C, Scheu S,
1010	Bonkowski M (2009b) Soil bacteria and protozoa affect root branching via effects on
1011	the auxin and cytokinin balance in plants. Plant and Soil 328: 191-201.

- Kuikman PJ, Jansen AG, Vanveen JA (1991) N-15-nitrogen mineralization from bacteria by
 protozoan grazing at different soil-moisture regimes. Soil Biology & Biochemistry 23:
 1014 193-200.
- Kuikman PJ, Jansen AG, Vanveen JA, Zehnder AJB (1990) Protozoan predation and the
 turnover of soil organic-carbon and nitrogen in the presence of plants. Biology And
 Fertility Of Soils 10: 22-28.
- Kuikman PJ, Vanveen JA (1989) The impact of protozoa on the availability of bacterial
 nitrogen to plants. Biology And Fertility Of Soils 8: 13-18.
- 1020 Kuzyakov Y, Xu X (2013) Competition between roots and microorganisms for nitrogen:

1021 mechanisms and ecological relevance. New Phytologist 198: 656-669.

- Lambshead P (1993) Recent developments in marine benthic biodiversity research. Oceanis
 1023 19: 5-5.
- Lambshead P, Chen Z, Chen W, Chen S, Dickson S (2004) Marine nematode diversity.

1025 Lavelle P, Spain AV (2001) Soil Ecology. Kluwer, Dordrecht.

- Leake JR, Johnson D, Donnelly DP, Muckle GE, Boddy L, Read DJ (2004) Networks of
- 1027 power and influence: the role of mycorrhizal mycelium in controlling plant
- 1028 communities and agroecosystem functioning. Can J Bot-Rev Can Bot 82: 1016-1045.
- 1029 Lennox LB, Alexander M (1981) Fungicide enhancement of nitrogen fixation and
- 1030 colonization of *Phaseolus vulgaris* by *Rhizobium phaseoli*. Appl Environ Microbiol
 1031 41: 404-411.
- 1032 Levrat P, Pussard M, Alabouvette C (1992) Enhanced bacterial metabolism of a Pseudomonas
- 1033 strain in response to the addition of culture filtrate of a bacteriophagous amoeba.
- European journal of protistology 28: 79-84.
- 1035 Louche J, Ali MA, Cloutier-Hurteau B, Sauvage F-X, Quiquampoix H, Plassard C (2010)
- 1036 Efficiency of acid phosphatases secreted from the ectomycorrhizal fungus *Hebeloma*

1037	cylindrosporum to hydrolyse organic phosphorus in podzols. Fems Microbiology
1038	Ecology 73: 323-335.
1039	Manzoni S, Taylor P, Richter A, Porporato A, Ågren GI (2012) Environmental and
1040	stoichiometric controls on microbial carbon-use efficiency in soils. New Phytologist
1041	196: 79-91.
1042	Marmeisse R, Guidot A, Gay G, Lambilliotte R, Sentenac H, Combier JP, Melayah D,
1043	Fraissinet-Tachet L, Debaud JC (2004) Hebeloma cylindrosporum - a model species to
1044	study ectomycorrhizal symbiosis from gene to ecosystem. New Phytologist 163: 481-
1045	498.
1046	Matz C, Kjelleberg S (2005) Off the hook-how bacteria survive protozoan grazing. Trends
1047	Microbiol 13: 302-307.
1048	Mazzola M, de Bruijn I, Cohen MF, Raaijmakers JM (2009) Protozoan-induced regulation of
1049	cyclic lipopeptide biosynthesis is an effective predation defense mechanism for
1050	Pseudomonas fluorescens. Appl Environ Microbiol 75: 6804-6811.
1051	Meier IC, Avis PG, Phillips RP (2013) Fungal communities influence root exudation rates in
1052	pine seedlings. Fems Microbiology Ecology 83: 585-595.
1053	Mooshammer M, Wanek W, Hämmerle I, Fuchslueger L, Hofhansl F, Knoltsch A, Schnecker
1054	J, Takriti M, Watzka M, Wild B (2014a) Adjustment of microbial nitrogen use
1055	efficiency to carbon: nitrogen imbalances regulates soil nitrogen cycling. Nature
1056	communications 5.
1057	Mooshammer M, Wanek W, Zechmeister-Boltenstern S, Richter A (2014b) Stoichiometric
1058	imbalances between terrestrial decomposer communities and their resources:
1059	mechanisms and implications of microbial adaptations to their resources. Frontiers in
1060	microbiology 5.

- Nasholm T, Kielland K, Ganeteg U (2009) Uptake of organic nitrogen by plants. New
 Phytologist 182: 31-48.
- 1063 Neff JC, Chapin III FS, Vitousek PM (2003) Breaks in the cycle: dissolved organic nitrogen
 1064 in terrestrial ecosystems. Front Ecol Environ 1: 205-211.
- Neidig N, Jousset A, Nunes F, Bonkowski M, Paul RJ, Scheu S (2010) Interference between
 bacterial feeding nematodes and amoebae relies on innate and inducible mutual
 toxicity. Funct Ecol 24: 1133-1138.
- Olsson PA, Baath E, Jakobsen I, Söderstrom B (1996) Soil bacteria respond to presence of
 roots but not to mycelium of arbuscular mycorrhizal fungi. Soil Biology &
 Biochemistry 28: 463-470.
- 1071 Parry JD (2004) Protozoan grazing of freshwater biofilms. Advances in applied microbiology
 1072 54: 167-196.
- Peters JL, Sutton AJ, Jones DR, Abrams KR, Rushton L (2006) Comparison of two methods
 to detect publication bias in meta-analysis. Jama-Journal of the American Medical
 Association 295: 676-680.
- 1076 Pimentel D (2006) Soil erosion: a food and environmental threat. Environment, development1077 and sustainability 8: 119-137.
- Plassard C, Bonafos B, Touraine B (2000) Differential effects of mineral and organic N
 sources, and of ectomycorrhizal infection by Hebeloma cylindrosporum, on growth
 and N utilization in Pinus pinaster. Plant Cell Environ 23: 1195-1205.
- Plassard C, Dell B (2010) Phosphorus nutrition of mycorrhizal trees. Tree Physiol 30: 11291139.
- 1083 Plassard C, Louche J, Ali MA, Duchemin M, Legname E, Cloutier-Hurteau B (2011)
- 1084 Diversity in phosphorus mobilisation and uptake in ectomycorrhizal fungi. Ann For1085 Sci 68: 33-43.

1086	Postma-Blaauw M, de Vries FT, De Goede R, Bloem J, Faber J, Brussaard L (2005) Within-
1087	trophic group interactions of bacterivorous nematode species and their effects on the
1088	bacterial community and nitrogen mineralization. Oecologia 142: 428-439.
1089	Puglisi E, Pascazio S, Suciu N, Cattani I, Fait G, Spaccini R, Crecchio C, Piccolo A, Trevisan
1090	M (2013) Rhizosphere microbial diversity as influenced by humic substance
1091	amendments and chemical composition of rhizodeposits. Journal of Geochemical
1092	Exploration 129: 82-94.
1093	Pussard M, Alabouvette C, Levrat P (1994) Protozoan interactions with the soil microflora
1094	and possibilities for biocontrol of plant pathogens.
1095	Ramirez C, Alexander M (1980) Evidence suggesting protozoan predation on Rhizobium
1096	associated with germinating seeds and in the rhizosphere of beans (Phaseolus vulgaris
1097	L.). Appl Environ Microbiol 40: 492-499.
1098	Ronn R, McCaig AE, Griffiths BS, Prosser JI (2002) Impact of protozoan grazing on bacterial
1099	community structure in soil microcosms. Appl Environ Microbiol 68: 6094-6105.
1100	Ronn R, Vestergard M, Ekelund F (2012) Interactions between bacteria, protozoa and
1101	nematodes in soil. Acta Protozoologica 51: 223-235.
1102	Rosenberg K, Bertaux J, Krome K, Hartmann A, Scheu S, Bonkowski M (2009) Soil
1103	amoebae rapidly change bacterial community composition in the rhizosphere of
1104	Arabidopsis thaliana. Isme J 3: 675-684.
1105	Saleem M, Fetzer I, Dormann CF, Harms H, Chatzinotas A (2012) Predator richness increases
1106	the effect of prey diversity on prey yield. Nature communications 3: 1305.
1107	Sardans J, Rivas-Ubach A, Penuelas J (2012) The elemental stoichiometry of aquatic and
1108	terrestrial ecosystems and its relationships with organismic lifestyle and ecosystem
1109	structure and function: a review and perspectives. Biogeochemistry 111: 1-39.

Version postprint

1111	community composition and individual biomass in northern hardwood forests of North
1112	America. Applied Soil Ecology 83: 159-169.
1113	Sinsabaugh RL, Manzoni S, Moorhead DL, Richter A (2013) Carbon use efficiency of
1114	microbial communities: stoichiometry, methodology and modelling. Ecol Lett 16:
1115	930-939.
1116	Smith SE, Read DJ (2008) Mycorrhizal Symbiosis - Third Edition. Academic Press, London.
1117	Somasundaram S, Bonkowski M, Iijima M (2008) Functional role of mucilage-border cells: a
1118	complex facilitating protozoan effects on plant growth. Plant production science 11:
1119	344-351.
1120	Sundin P, Valeur A, Olsson S, Odham G (1990) Interactions between bacteria-feeding
1121	nematodes and bacteria in the rape rhizosphere - Effects on root exudation and
1122	distribution of bacteria. Fems Microbiology Ecology 73: 13-22.
1123	Tao J, Chen X, Liu M, Hu F, Griffiths B, Li H (2009) Earthworms change the abundance and
1124	community structure of nematodes and protozoa in a maize residue amended rice-
1125	wheat rotation agro-ecosystem. Soil Biology & Biochemistry 41: 898-904.
1126	Tibbett M, Sanders FE (2002) Ectomycorrhizal symbiosis can enhance plant nutrition through
1127	improved access to discrete organic nutrient patches of high resource quality. Annals
1128	of Botany 89: 783-789.
1129	Venette RC, Ferris H (1998) Influence of bacterial type and density on population growth of
1130	bacterial-feeding nematodes. Soil Biology & Biochemistry 30: 949-960.
1131	Vitousek PM, Cassman K, Cleveland C, Crews T, Field CB, Grimm NB, Howarth RW,
1132	Marino R, Martinelli L, Rastetter EB, Sprent JI (2002) Towards an ecological
1133	understanding of biological nitrogen fixation. Biogeochemistry 57: 1-45.

Schlaghamersky J, Eisenhauer N, Frelichc LE (2014) Earthworm invasion alters enchytraeid

1110

Version postprint

1134	Wagg C, Bender SF, Widmer F, van der Heijden MGA (2014) Soil biodiversity and soil
1135	community composition determine ecosystem multifunctionality. Proceedings Of The
1136	National Academy Of Sciences Of The United States Of America 111: 5266-5270.
1137	Wall DH, Bardgett RD, Kelly EF (2010) Biodiversity in the dark. Nat Geosci 3: 297-298.
1138	Wallenda T, Read DJ (1999) Kinetics of amino acid uptake by ectomycorrhizal roots. Plant
1139	Cell Environ 22: 179-187.
1140	Weekers PH, Bodelier PL, Wijen JP, Vogels GD (1993) Effects of grazing by the free-living
1141	soil amoebae Acanthamoeba castellanii, Acanthamoeba polyphaga, and Hartmannella
1142	vermiformis on various bacteria. Appl Environ Microbiol 59: 2317-2319.
1143	Weisse T (2002) The significance of inter- and intraspecific variation in bacterivorous and
1144	herbivorous protists. Antonie Van Leeuwenhoek International Journal of General and
1145	Molecular Microbiology 81: 327-341.
1146	Wright DJ (1975) Elimination of nitrogenous compounds by Panagrellus redivivus, Goodey,
1147	1945 (Nematoda: Cephalobidae). Comparative Biochemistry and Physiology Part B:
1148	Comparative Biochemistry 52: 247-253.
1149	Xiao HF, Griffiths B, Chen XY, Liu MQ, Jiao JG, Hu F, Li HX (2010) Influence of bacterial-
1150	feeding nematodes on nitrification and the ammonia-oxidizing bacteria (AOB)
1151	community composition. Applied Soil Ecology 45: 131-137.
1152	Xu X, Thornton PE, Post WM (2013) A global analysis of soil microbial biomass carbon,
1153	nitrogen and phosphorus in terrestrial ecosystems. Global Ecology and Biogeography
1154	22: 737-749.
1155	Yeates G (2007) Abundance, diversity, and resilience of nematode assemblages in forest
1156	soils. Canadian journal of forest research 37: 216-225.
1157	Yeates GW (2003) Nematodes as soil indicators: functional and biodiversity aspects. Biology
1158	and Fertility of Soils 37: 199-210.

1159	Zechmeister-Boltenstern S, Keiblinger KM, Mooshammer M, Peñuelas J, Richter A, Sardans
1160	J, Wanek W (2015) The application of ecological stoichiometry to plant-microbial-
1161	soil organic matter transformations.
1162	Zwart KB, Darbyshire JF (1992) Growth and nitrogenous excretion of a common soil
1163	flagellate Spumella sp a laboratory experiment. Journal Of Soil Science 43: 145-
1164	157.

Table 1. Effects of bacterivores on plant root parameters (% of control).

References	Years	Plant species	Bacterivores	Modalities within studies	Root architecture (min – max)*			
					Length (m)	SRL (m g^{-1})	Area (m ²)	Tips (number)
Jentschke et al.	1995	Picea abies	Protist	- Lactarius rufus	135 - 180	38 - 70		130 - 220
				+ Lactarius rufus	22 - 45	2-36		30 - 45
Bonkowski et al.	2001	Picea abies	Protist	- Paxillus involutus	62	56		50
				+ Paxillus involutus	32	-32		32
Bonkowski & Brandt	2002	Lepidium sativum	Protist	Main roots	15			
				Lateral roots	421			286 ^{\$}
Kreuzer et al.	2006	Oryza sativa	Protist	Total roots	17			-37 ^{\$}
				Root diameter > 4mm	-43			-56 ^{\$}
				Root diameter < 4mm	195			1140 ^{\$}
Herdler et al.	2008	Oryza sativa	Protist	- Glomus intraradices		-27		
				+ Glomus intraradices		-15		
Krome et al.	2009	Lepidum sativum	Protist	Agar medium				100
				Soil medium	28 - 62			88 - 91
Irshad et al.	2011	Pinus pinaster	Nematode	-			-18	3
Cheng et al.	2011	Oryza sativa	Nematode	Day 14	14 – 22			26 - 43
				Day 20	2 – 15			-3 - 12
Irshad et al.	2012	Pinus pinaster	Nematode	- Hebeloma cylindrosporum	37 – 41		20 - 35	31 - 46
				+ Hebeloma cylindrosporum	20 - 32		16 - 20	18 - 36
Koller et al.	2013b	Plantago lanceolata	Protist				31	
Ranoarisoa et al.	2015	Pinus pinaster	Nematode	- Hebeloma cylindrosporum	7 - 32		9 - 22	1 – 15
				+ Hebeloma cylindrosporum	6 - 31		2 - 23	4 - 23
Mean					56	16	16	96

*Specific root length (SRL)

2

^{\$}Number of tips estimated from number of roots

Comment citer ce document : Trap, J., Bonkowski, M., Plassard, C., Villenave, C., Blanchart, E. (2015). Ecological importance of soil bacterivores for ecosystem functions. Plant and Soil, 1-24. DOI : 10.1007/s11104-015-2671-6 Figure 1. Number of studies included in the meta-analysis (total and per criteria).



Comment citer ce document : Trap, J., Bonkowski, M., Plassard, C., Villenave, C., Blanchart, E. (2015). Ecological importance of soil bacterivores for ecosystem functions. Plant and Soil, 1-24. DOI : 10.1007/s11104-015-2671-6 **Figure 2.** Mean effect sizes (ES_m) of bacterivores on ecosystem functions. White and black circles indicate significant (different from zero) and non-significant ES_m , respectively. Horizontal bars correspond to 95% confidence intervals. Bold numbers above circles specify the ES_m expressed as % of control (%CC). Italic numbers between brackets specify the total number of observations for each function. M_{mic} : microbial biomass; B_{num} : bacterial number; Q_{mic} : microbial metabolic quotient; N_{min} : net N mineralization; P_{min} : net P mineralization; S:R : shoot:root ratio.



Version postprint

Figure 3. Changes in bacterivore-induced effect size (mean) on soil net N (A) and P (B) mineralization according to media types used in experimental studies. Letters (a and b) indicate significant differences in ES_m between media types according to a Kruskal-Wallis test at *P*<0.05.



Figure 4. Changes in bacterivore-induced effect size (mean) on shoot N mineralization (A), root biomass (B) and shoot N concentration (C) according to the presence of bacterial-feeding protists (Pr), nematodes (Nem) alone or together (Pr + Nem). Letters (a and b) indicate significant difference in ES_m among media types according to a Kruskal-Wallis test at P < 0.05.



Figure 5. Multiple linear regressions performed on individual effect-size (ES_i). (A) ES_i of total N amount in shoot (response variable) as a function of ES_i of shoot biomass and ES_i of shoot N concentration (explanatory variables). (B) ES_i on total P amount in root (response variable) as a function of ES_i on root biomass and ES_i on root P concentration (explanatory variables). $%_{expl.}$ indicates the percentage of total variance of the response variable explained by each explanatory variable and by their interaction. ***P<0.001 (n= number of

7 observations).



Comment citer ce document : Trap, J., Bonkowski, M., Plassard, C., Villenave, C., Blanchart, E. (2015). Ecological importance of soil bacterivores for ecosystem functions. Plant and Soil, 1-24. DOI : 10.1007/s11104-015-2671-6 1 Figure 6. Changes in bacterivore-induced effect size (mean) on shoot (A) and root (B)

- 2 biomass according to the presence of mycorrhizal fungi (-M : non mycorrhizal plant; +M:
- 3 mycorrhizal plant). Different letters (a and b) indicate significant changes according to a
- 4 Kruskal-Wallis test (*P*<0.05, n=14).
- 5



Comment citer ce document : Trap, J., Bonkowski, M., Plassard, C., Villenave, C., Blanchart, E. (2015). Ecological importance of soil bacterivores for ecosystem functions. Plant and Soil, 1-24. DOI : 10.1007/s11104-015-2671-6

- Figure 7. Theoretical framework describing the relative dominance* of nutritional and non-nutritional mechanisms involved in bacterivoreeffects on plant performance.
- 3 (A) Relationship between plant relative growth rate and plant C:nutrient ratios (modified from Ågren (2008)). The solid line describes the
- 4 C:limiting-nutrient (N or P) ratio while the broken lines indicate three different variations in the C:nutrient ratio of a second element (N or P) as a
- 5 function of relative growth rate. We here defined the *critical* C:limiting-nutrient ratio (R_{cri}) above which the relative growth rate is null.
- 6 (B) Diagram illustrating four different cases of plant behavior in the presence of bacterivores along a soil C:N and C:P gradients. Solid lines
- 7 describe the C:N_{cri} and C:P_{cri} according to soil C:N:P ratio, while the broken line indicates a hypothetical zone where the dominant mechanism
- 8 changes. Grey arrows describe the hypothetical trends in microbial nitrogen- and phosphorus-use efficiency (NUE, PUE) along the soil C:N:P
- 9 gradient. *the term "dominance" describes the relative importance of mechanisms to initiate and drive the effects of bacterivores on plants at a
- 10 given moment in the experiment.



13

Comment citer ce document : Trap, J., Bonkowski, M., Plassard, C., Villenave, C., Blanchart, E. (2015). Ecological importance of soil bacterivores for ecosystem functions. Plant and Soil, 1-24. DOI : 10.1007/s11104-015-2671-6

Comment citer ce document : Trap, J., Bonkowski, M., Plassard, C., Villenave, C., Blanchart, E. (2015). Ecological importance of soil bacterivores for ecosystem functions. Plant and Soil, 1-24. DOI : 10.1007/s11104-015-2671-6 Supplementary Material

1 **Appendix 1.** List of studies (in chronological order) used in the meta-analysis.

References	Year	Country	Medium	Bacterivores		Nutrients		Plant (species)	Mycorrhiza	Journals	
				Protist	Nematode	Nitrogen	Phosphorus				
Coleman et al.	1977	USA	Soil	Yes	Yes	Yes	Yes	No	No	Ecological Bulletins	
Cole et al.	1978	USA	Soil	Yes	Yes	No	Yes	No	No Microbial Ecology		
Elliott et al.	1979	USA	Soil	Yes	No	Yes	Yes	No	No	No Intern. J. Environmental Studies	
Cole et al.	1978	USA	Soil	Yes	Yes	No	Yes	No	No	Microbial Ecology	
Anderson et al.	1981	USA	Soil	No	Yes	Yes	Yes	No	No	Microbial Ecology	
Baath et al.	1981	Sweden	Humus	No	Yes	Yes	No	Pinus sylvestris	No	Oïkos	
Woods et al.	1982	USA	Soil	Yes	Yes	Yes	No	No	No	Soil Biology & Biochemistry	
Trofymow et al.	1983	USA	Soil	No	Yes	Yes	No	No	No	Oecologia	
Anderson et al.	1983	USA	Soil	No	Yes	Yes	No	No	No	Oïkos	
Clarholm	1985	Sweden	Soil	Yes	No	Yes	No	Triticum sp.	No	Soil Biology & Biochemistry	
Inghamet al.	1985	USA	Soil	No	Yes	Yes	Yes	Bouteloua gracilis	No	Ecological Monographs	
Griffiths	1986	Scotland	Soil	Yes	Yes	Yes	Yes	No	No	Soil Biology & Biochemistry	
Kuikman & Van Veen	1989	Netherlands	Soil	Yes	No	Yes	No	Triticum aestivum	No	Biology and Fertility of Soils	
Kuikman et al.	1990	Netherlands	Soil	Yes	No	Yes	No	Triticum aestivum	No	Biology and Fertility of Soils	
Setälä et al.	1991	Finland	Humus	No	Yes	Yes	Yes	No	No	Pedobiologia	
Kuikman et al.	1991	Netherlands	Soil	Yes	No	Yes	No	Triticum aestivum	No	Soil Biology & Biochemistry	
Rutherford et al.	1992	Canada	Soil	Yes	No	Yes	No	No	No	Canadian Journal of Soil Science	
Bouwman et al.	1994	Netherlands	Soil	No	Yes	Yes	No	No	No	Biology and Fertility of Soils	
Jentschke et al.	1995	Germany	Sand + nutrient solution	Yes	No	Yes	Yes	Picea abies	Yes	Biology and Fertility of Soils	
Alphei et al.	1996	Germany	Soil	Yes	Yes	Yes	Yes	Hordelymus europaeus	No	Oecologia	
Ferris et al.	1998	USA	Sand-OM substrate	No	Yes	Yes	No	No	No	Plant and Soil	
Setälä et al.	1999	Finland	Sand-OM substrate	No	Yes	Yes	No	Pinus sylvestris	Yes	Oikos	
Bardgett & Chan	1999	UK	Soil	No	Yes	Yes	Yes	Nardus stricta	No	Soil Biology & Biochemistry	
Bonkowski et al.	2000	UK	Soil-sand-OM substrate	Yes	Yes	Yes	No	Lolium perenne	No	Applied Soil Ecology	
Bonkowski et al.	2001a	UK	Soil-sand-OM substrate	Yes	No	Yes	Yes	Triticum aestivum	No	Oïkos	
Bonkowski et al.	2001b	Germany	Humus	Yes	No	Yes	Yes	Picea abies	Yes	Applied Soil Ecology	
Djigal et al.	2004a	Senegal	Soil	No	Yes	Yes	Yes	Zea mays	No	Plant & Soil	
Djigal et al.	2004b	Senegal	Soil	No	Yes	Yes	Yes	Zea mays	No	Soil Biology & Biochemistry	
Postma-Blaauw et al.	2005	Netherlands	Soil-OM mixture	No	Yes	Yes	No	No	No	Oecologia	
Herdler et al.	2008	Germany	Soil	Yes	No	Yes	Yes	Oryza sativa	Yes	Soil Biology & Biochemistry	
Somasundaram et al.	2008	Japan	Soil	Yes	No	No	No	Oryza sativa	No	Plant Production Science	
Ekelund et al.	2009	Netherlands	Soil-OM mixture	Yes	No	Yes	No	Holcus lanatus	No	Soil Biology & Biochemistry	
Krome et al.	2009a	Germany	Sand-OM mixture	Yes	No	Yes	No	Arabidopsis thaliana	No	No Soil Biology & Biochemistry	
Krome et al.	2009b	Germany	Agar/Sand	Yes	No	Yes	No	Lepidium sativum & Arabidopsis thaliana	No	Plant & Soil	
Xiao et al.	2010	China	Soil-OM mixture	No	Yes	Yes	No	No	No	Applied Soil Ecology	
Cheng et al.	2011	China	Soil	No	Yes	Yes	No	Oryza sativa	No	Nematology	
Irshad et al.	2011	France	Agar	No	Yes	Yes	Yes	Pinus pinaster	No	Soil Biology & Biochemistry	
Bjornlund et al.	2012	Denmark	Soil-OM mixture	No	Yes	Yes	No	Hordeum vulgare	No	European Journal of Soil Biology	
Irshad et al.	2012	France	Agar	No	Yes	No	Yes	Pinus pinaster	Yes	Plant and Soil	
Koller et al.	2013a	Germany	Soil-sand mixture	Yes	No	Yes	No	Plantago lanceolata	Yes	Soil Biology & Biochemistry	
Koller et al.	2013b	Germany	Soil-sand mixture	Yes	No	Yes	No	Plantago lanceolata	No	FEMS Microbiology Ecology	
Ranoarisoa et al.	2015	France	Soil-sand mixture	No	Yes	Yes	Yes	Pinus pinaster	Yes	Personnal communication	