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Soil biota in boreal urban greenspace: responses

2to plant type and age

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29Key words: Microbial biomass (PLFA), Urban greenspace, Nematodes,

30Earthworms

32Abstract

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34Plant functional type influences the abundance and distribution of soil biota. 35With time, as root systems develop, such effects become more apparent. The 36 relationship of plant type and time with the structure and abundance of soil 37microbial and invertebrate communities has been widely investigated in a 38variety of systems. However, much less is known about long-term soil 39community dynamics within the context of urban environments. In this study, 40we investigated how soil microbes, nematodes and earthworms respond to 41 different plant functional types (lawns only and lawns with deciduous or 42evergreen trees) and park age in 41 urban parks in southern Finland. As non-43urban controls we included deciduous and evergreen trees in 5 forest sites. We 44 expected that microbial biomass and the relative abundance of fungi over 45bacteria would increase with time. We also expected major differences in soil 46microbial and nematode communities depending on vegetation: we 47hypothesized that i) the presence of trees, and evergreens in particular, would 48support a greater abundance of fungi and fungal-feeding nematodes over 49bacteria and bacterial-feeding nematodes and ii) the fungi to bacteria ratio 50would be lowest in lawns, with deciduous trees showing intermediate values. 51In contrast to our predictions, we showed that old deciduous trees, rather than 52evergreens, supported the highest fungal abundances and fungal-feeding 53nematodes in the soil. Consistent with our predictions, microbial biomass in 54urban park soils tended to increase with time, whereas – in contrast to our 55hypotheses – fungal-feeding nematode abundance declined. Even in the oldest 56parks included in the current study, microbial biomass estimates never 57approximated those in the minimally managed natural forests, where biomass 58estimates were three times higher. Anecic earthworm abundance also 59 increased with time in urban parks, whereas abundances of fungal-feeding, 60plant-feeding and omnivorous nematodes, as well as those of epigeic and 61endogeic earthworms remained constant with time and without any distinct 62differences between urban parks and the control forests. Our findings highlight 63that although urban park soils harbor diverse soil communities and

64considerable microbial biomass, they are distinct from adjacent natural sites in 65community composition and biomass.

67Highlights

- 69• In cities, soil communities under different plant types diverge
- 70 increasingly with time
- 71• Microbial biomass associated with deciduous trees increases with park
- 72 age
- 73• Earthworm biomass increases with park age
- 74• Adjacent non-urban sites maintain a greater microbial biomass than
- 75 urban parks

761. Introduction

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78Urbanized areas are rapidly expanding at the expense of natural habitats. 79Urban green spaces, including public parks, assume a pivotal role as surrogates 80for these lost natural environments. These green spaces provide a vast array of 81ecosystem services (Costanza et al., 1997; Bolund and Hunhammar, 1999) 82including carbon and nitrogen sequestration (Raciti et al., 2011; Setälä et al., 832016), storm water interception and purification (Valtanen et al., 2015), 84biodiversity and climate regulation (Bolund and Hunhammar, 1999). These 85ecosystem services depend strictly on the soil, and in turn on the soil biota 86hosted therein.

Soil microbes and invertebrates are directly linked to biogeochemical 88processes that take place in the soil and thus promote a variety of soil-derived 89ecosystem services (Lavelle et al. 2006; Balser and Firestone, 2005; Blouin et 90al., 2013 Ledin, 2000; Haritash and Kaushik, 2009). Yet, the abundance and 91distribution of soil biota are often directly linked with the distribution of plant 92species, and indirectly to soil properties that are modified by the plants 93(Wardle et al., 2004) over time (Bardgett et al., 2005). Although the 94successional trajectories of soil communities in relation to plant type and soil 95characteristics over time have been widely investigated in, e.g., primary 96(Ohtonen et al., 1999; Doblas-Miranda et al., 2008; Brown and Jumpponen 972014) and secondary succession (Pižl, 1992; Maharning et al., 2009), much less 98is known about these dynamics in urban park soils.

99 The linkage between plant functional types and soil microbial and 100invertebrate communities has received substantial interest in contemporary 101soil- and ecosystem ecology (Orwin et al., 2010; Thomson et al., 2010). There is 102also a growing body of literature describing soil biota in urban parks 103(microbes: Baxter et al., 1999; Xu et al., 2014; Ramirez et al., 2014; Hui et al. 1042017a, b; nematodes: Pavao-Zuckerman and Coleman 2007; Amossé et al., 1052016; earthworms: Steinberg et al., 1997; Smetak et al., 2007; Amossé et al., 1062016). In a nutshell, these studies collectively provide evidence that 107urbanization can substantially change in soil microbial and faunal 108communities. Yet, there is a paucity of studies that simultaneously account for

109different trophic groups in large and well-replicated experimental designs 110within the context of urban ecosystems.

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111 Plant species identity is linked to the quantity and quality of inputs 112provided to the soil, either via litter deposition or through root exudates, which 113in turn are largely responsible for the composition of soil microbial (Grayston 114et al., 1998; Bardgett and McAlister, 1999; Marschner et al., 2004), nematode 115(Ilieva-Makulec et al., 2006) and earthworm communities (Curry, 2004). Whilst 116 plants that produce labile, nitrogen-rich litter, such as grasses, herbs and 117 deciduous trees often support bacterial-dominated soil microflora, plants 118producing more recalcitrant litter, such as evergreens, more commonly 119support fungal-based soil food webs (Wardle et al. 2004). Evergreen trees are 120adapted to low nutrient availability and thus have leaves with low nutrient 121 contents (Kattge et al. 2011). In contrast, deciduous broadleaf trees have 122higher foliar nutrient content (Kattge et al. 2011) with *Tilia cordata* and *Acer* 123platanoides particularly in particular producing high quality litter (Aerts and 124Chapin 2000, Hobbie et al. 2014). Moreover, soil acidification promoted by 125evergreen trees (Setälä et al. 2016) is also associated with the prevalence of 126fungi over bacteria (Bååth and Anderson 2003). Finally, the quantity of root 127exudates, in boreal systems, tends to be higher in evergreen trees than 128deciduous trees (Gower et.al 2001), suggesting that evergreen trees may 129allocate a higher percentage of net primary productivity to, i.e. ectomycorrhizal 130fungi than do deciduous trees. Such plant-associated distinctions in soil 131microbial communities are reflected in the relative proportions of bacterial-132and fungal-feeding fauna (Trofymow and Coleman, 1982; de Vries et al., 2013). 133This is particularly interesting as fungal-based food webs are characterized by 134slow nutrient cycling and a high capacity to retain nutrients, whereas bacterial 135dominated food webs are characterized by high nutrient turnover and nutrient 136leaching (de Vries et al., 2006; de Vries et al., 2012). Consequently, the ratio 137between fungi and bacteria in the soil is crucial when considering fundamental 138ecological processes such as the rate of organic matter decomposition 139(Coleman et al., 1983; Wardle, 2002; Moore et al., 2005; Paterson et al., 2008) 140and thus carbon and nutrient sequestration.

141The type and availability of plant-derived resources can also change142during plant community succession (Berendse, 1990; Knops and Tilman,1432000). As soil organic matter (OM), C and N accumulate over time, so too do the144biomasses of soil microbes (Zak et al., 1990; Ohtonen et al., 1999), nematodes145(Háněl, 2010) and earthworms (Pižl, 1992). Changes in microbial abundances146are not only quantitative, but also qualitative. For instance, the relative147abundance of fungi over bacteria tends to increase with time (Ohtonen et al.,1481999; Zeller et al., 2001; Bardgett and Walker, 2004). Furthermore, temporal149changes in the fungal to bacterial ratio during succession are often also150mirrored by changes in the ratio of fungal feeding to bacterial feeding151nematodes (Brzeski, 1995; Ferris and Matute, 2003; Háněl, 2010). Only a few152studies have investigated such successional trajectories of soil biota in urban153parks (Smetak et al., 2007; Amossé et al., 2016; Hui et al., 2017a, b).

This study is part of a larger project that aims to shed light on the 155 influence of divergent plant types on the physico-chemical and biological soil 156 characteristics in urban parks of diverging ages. Our previous work has shown 157 that plant type and park age are strong determinants of soil characteristics 158 (Setälä et al., 2016; Setälä et al. 2017) and soil microbial community 159 composition (Hui et al., 2017a, b). However, the response of microbial biomass, 160 a measure that strongly relates to the functional activity and capacity of soils, 161 to plant type and park age remains unresolved. Moreover, in this study, we 162 incorporate the microbial consumer responses – key actors in providing soil-163 based ecosystem services and contributing to nutrient turnover (Wardle et al., 1642004). We aim to investigate how park age and plant functional type affect the 165 biomass of soil microbes and two important functional groups of soil fauna: 166 nematodes and earthworms.

Given the clear effects that plant functional type and park age have on Given the clear effects that plant functional type and park age have on Setälä et al. 2016; Setälä et al. 2017) we tested the Setälä et al. 2017) we tested the Helpfollowing hypotheses: plant functional type affects the soil food web so that i) Proevergreen trees producing recalcitrant litter promote an increase of fungal Proventies over bacterial biomass and ii) deciduous trees, the lawn in particular, Producing more labile litter, promote the establishment of bacterial biomass. Proventies in the soil microbial community also cascade up to

174higher trophic levels. We expect that higher densities of fungal feeding 175nematodes (compared to bacterial feeders) associate with evergreen trees and 176higher densities of earthworms associate with lawns and deciduous trees. Also, 177we test iv) whether time since park construction promotes changes in soil 178microbial and invertebrate community structure and abundance. We 179hypothesize that soil biota in old parks resemble natural communities more 180than in young parks. This is because the capacity of plants (especially trees) to 181modify soils is park-age dependent (Setälä et al. 2016, 2017) with young parks 182not having had the time to develop plant-soil interactions that are typical of 183natural forests.

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1862. Materials and Methods

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1882.1. Study area

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190The study was conducted in two municipalities in the boreal forest zone in 191southern Finland; Helsinki metropolitan area (60° 10′15′′N 24° 56′ 15′′E, with 192a population of ca. 1.4 million people) and Lahti (60° 58' 57 N 25° 39' 41 E, 193population ca. 110 000). Winters are cold and wet, while rainfall is moderate 194all year round. The annual mean temperature is 5.3°C in Helsinki and 4.5°C in 195Lahti; the annual mean precipitation is 628 mm in Helsinki and 636 mm in 196Lahti. Summer lasts for approximately 110-120 days, winter 135-145 days and 197the temperature can span from -35 to +35°C (Finnish Meteorological Institute). 198The Helsinki and Lahti regions are classified by NRCS (National Resource 199Conservation Service) as having soils of primarily the Spodosol suborder. 200However, all urban parks in the two cities are constructed and none showed 201detectable signs of podzolisation at the time of sampling.

We selected 41 parks in the two cities and five additional control forests 203(see Setälä et al., 2016 and Hui et al., 2017a for details). Parks of three ages 204were selected: young parks (between 7 and 15 years old), intermediate parks 205(ca. 50 years old) and old parks (> 100 years). Control forests, situated in the 206outskirts of the city of Lahti, are typified as unmanaged, conifer and linden

207dominated forests (> 80 years of age). Park size varied from one to several 208hectares. The selected parks were subjected to routine maintenance, including 209mowing (mowing residues not removed) and raking of tree leaves in the fall. 210However, the parks were not irrigated or commonly fertilized. Until the early 2111990s, some of the older parks in the city of Lahti were occasionally fertilized -212commonly with saltpeter (N, P, K, S), while some of the park lawns in Helsinki 213have received and still receive light refurbishment fertilization.

214 In each park we selected, where possible, three vegetation-types: 215deciduous (represented by Tilia x vulgaris 93% and Acer platanoides 7%) and 216evergreen trees (spruce, Picea sp. 43.3%, Abies sp. 20%, Pseudostuga menziesii 21713.3%, Pinus sylvestris 13.3%, Larix sp. 10%) and a non-treed lawn with grass 218(including herbs such as Trifolium pratense, Plantago major). Lawn cover 219 extended also under tree canopies. In some cases we selected parks including 220only deciduous trees and lawn or only evergreen trees and lawn. The control 221 sites never had lawns, but deciduous (forest linden, Tilia cordata) and 222evergreen trees (Norway spruce, Picea abies) were always present at each site. 223In the parks, distance between the two tree types was always greater than the 224height of the nearest tree. Plant age was considered as coinciding with park 225age, except for young parks, where ca. 10 year old saplings were planted at the 226time of park construction. In order to have at least 10 replicates per park age 227and plant functional type, we selected 41 urban parks, resulting in 91 sampling 228locations as described by Setälä et al. (2016) plus five control forests with 10 229additional sampling locations, totaling 101 sampling locations. 230

2312.2. Soil sampling and measurements for edaphic responses 232

233In urban parks, soils were sampled at the edge of the canopy projection of each 234tree and in the middle of the lawn area (when present) in October 2014. In the 235control forests, soils were sampled under the canopy projection of each tree in 236May 2015. Samples were obtained using a metal corer that was sterilized in 237ethanol between samples. At each sampling point, 3 subsample soil cores were 238collected and then pooled. The soil from each sample was homogenized and 239larger stones, roots and fresh or recognizable plant material removed. Samples

240for microbial analysis were stored in resalable plastic bags on ice in the field 241and frozen at -20 °C in the laboratory. Soil pH was measured in a 1/5 (vol/vol) 242soil/distilled water suspension. Soil was weighed and analyzed for percent 243moisture by drying for 48 hr at 105 °C. Total carbon (hereafter C) and nitrogen 244(N) were obtained by dry combustion at 1350 °C using a LECO CNS2000 245Elemental Analyzer (0.07% C and 0.09% N detection limits) and reported as 246percentage dry mass.

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2482.3. Soil microbial analysis, PLFA

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250Before analysis, the frozen samples were thawed at room temperature and 251 sieved through a 2 mm mesh to remove leaf litter, rocks, large particles and 252roots. Then, part of the soil was freeze-dried and an aliquot of 4 g for urban 253park soil and 2 g for forest soil was taken for phospholipid fatty acids (PLFA) 254analysis. Fatty acids (FA) were extracted following a modified procedure as 255described in Macnaughton et al. (1997), Bligh and Dyer (1959), Frostegård et 256al. (1991) and White et al. (1979). The extraction was performed using a 257Dionex ASE 350 machine (Thermo Scientific). Briefly, the samples were mixed 258 with diatomaceous earth and placed in the extraction vessels between two 259cellulose filters prewashed in chloroform. FA were extracted with Bligh and 260Dyer (methanol/chloroform/citrate buffer, 2/1/0.8 vol/vol/vol) at 80 °C and 261the samples underwent two static cycles of 15 min each. Once FA acids were 262extracted, chloroform and citrate buffer were added, for a final volume of 263chloroform/methanol/citrate buffer of 1/1/0.9, vol/vol/vol. The extracts were 264left overnight to separate the aqueous phase from the lipidic phase. The 265supernatant was discharged and the lower phase collected, which was then 266evaporated under a flux of N₂ at 40 °C. The resulting pellet was suspended in 267chloroform and applied to silica columns (Bond Elut LRC, Agilent). Chloroform, 268acetone and methanol were applied in sequence to the column to separate 269neutral lipids, glycolipids and phospholipids. The extracts were then dried 270under N₂. The phospholipid fraction so obtained was methylated and the fatty 271acid methyl esters extracted with hexane/chloroform (4/1, vol/vol), dried 272under N₂ and redissolved in 1 ml hexane. We used 19:0 (methyl

273nonadecanoate) as an internal standard. All the solvents used were of GC grade 274and glassware was baked in an oven at 450 °C for 4 h prior to use.

275 The samples were analyzed in a Shimadzu GCMSQP2010 Ultra with an 276Agilent J&WDB23 column. The column oven temperature was set to 60 °C and 277the injection temperature was 250 °C in splitless mode. MS ion source 278temperature was 200 °C and the interface temperature was 250 °C. As 279reference library we used the bacterial acid methyl ester (BAME) mix 280(SigmaAldrich), while the Supelco 37 Component FAME mix (SigmaAldrich) 281was used for calibration for PLFA biomass calculation. Peaks were cross-282checked against the NIST spectral library. A total of 29 PLFAs were identified. 2830f the identified peaks, the following were considered to be of bacterial origin: 284i15:0, 15:0, 17:0, 17:0cy (eubacteria); a15:0, i16:0, i17:0 (gram-positive 285bacteria); 16:1 ω 9, 18:1 ω 7 (gram-negative bacteria). The PLFA 18:2 ω 6 was 286considered to be mainly of fungal origin (Frostegård and Bååth, 1996). PLFAs 287were expressed as $\mu g g^{-1}$ soil dry weight. The totality of identified peaks was 288used to investigate coarse microbial community changes, as explained in 289section 2.5 below. The microbial to fungal (F/B) ratio was calculated as the 290ratio of PLFA 18:206 to bacterial PLFAs.

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2922.4. Soil Fauna

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2942.4.1. Nematodes

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296Nematodes were collected in early June 2016 in Lahti, and in July-August 2016 297in Helsinki. Similarly to the soil sampling, nematodes were collected at the edge 298of the canopy projection of each tree and in the middle of the lawn (when 299present) area. In the control forests, soils were sampled under the canopy 300projection of each tree type. Each sample consisted of six pooled 5 cm diameter 301soil cores of the uppermost 10 cm. The samples were pooled to make one 302composite sample and immediately placed in plastic bags and stored in a 303cooler. In the laboratory, the soil was gently mixed and 10 g (fresh weight) of 304soil was used for nematode extraction. Nematodes were extracted for 48 h 305without lights/heating following the wet funnel method described in Sohlenius

306(1979). Nematode feeding guilds were assigned following Yeates et al. (1993). 307We calculated the nematode channel ratio (NCR) as described in Yeates (2003) 308and the maturity index (MI) as in Bongers (1990). NCR is calculated as the ratio 309of bacterial feeding (BF) to bacterial-feeding plus fungal-feeding (FF) 310nematodes: BF/(BF+FF), and is constrained to have values between 1 (totally 311bacterial-mediated) and 0 (totally fungal-mediated) (Yeates, 2003). MI is the 312sum of the relative abundance of each taxon multiplied by their colonizer-313persister (c.p.) value. The c.p. value of each family was assigned following 314Bongers (1990, 1999). MI provides information on the degree of 315environmental disturbance, with higher values indicating a less disturbed 316environment.

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3182.4.2. Earthworms

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320Earthworms were collected in August 2016 close to where soil microbes and 321nematodes were sampled. We followed a modified version of the hot mustard 322liquid method described by Gunn (1992). Each sample consisted of two (in 323public parks) or four (in control sites) 25 × 25 cm frames irrigated with a hot 324mustard liquid. The hot mustard slurry was prepared by mixing 15 g of hot 325mustard powder (Colman's powder) in 100 ml of water, and then allowing it to 326sit for at least 4 h. Immediately prior to sampling, the mustard slurry was 327added to approximately 3.5 L of tap water in a watering can. This 3.5 L of 328mustard water was applied to each frame over a span of 15–20 min. The 329collected worms were immediately placed in tap water for some minutes, then 330placed into plastic bags containing moist tissue paper, and stored in a cooler. In 331the laboratory the earthworms were placed in a refrigerator and stored for 48 332h to empty their guts. The worms were then identified, weighed and stored in 3334% formaldehyde solution.

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3352.5. Data analysis

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337Generalized linear mixed models (GLMM) were used to investigate the 338relationship between our focal response variables (see below) and park age (a

339factor with three levels; young, intermediate, old), plant functional type (a 340factor with three levels; evergreen, deciduous, lawn), soil pH, % soil C, % soil N, 341C/N ratio, soil OM, and % soil moisture. Our response variables were: total 342microbial biomass (PLFA), PLFA fungal/bacterial ratio, PLFA fungal marker $343(18:2\omega 6)$, PLFA gram-/gram+ ratio; nematode NCR ratio, BF nematode, FF 344nematode, PF (plant-feeding) nematode, PO (predators and omnivores) 345nematode, anecic earthworm Lumbricus terrestris abundance and endogeic and 346epigeic earthworms (Lumbricus rubellus, epigeic; Allolopophora caliginosa, 347endogeic; Octolasion cyaneum, endogeic - collected in one sample only; 348*Allolophora chlorotica*, endogeic - collected in one sample only; *Dendrobaena* 349octaedra, epigeic). These GLMMs did not include the control forest samples in 350Lahti. Response variables were log or square-root transformed, when 351necessary, to satisfy assumptions of normality. In the GLMM analysis, city was 352considered a random effect, with park identity nested within city. We 353performed a stepwise model selection procedure by removing insignificant 354predictor variables, one at a time, if their p-values were greater than 0.05 and if 355the AIC subsequently decreased. However, park age and vegetation functional 356type were always retained in the final model, irrespective of significance. Then, 357we compared the control forest with old parks in Lahti, using ANOVA, with 358park age and vegetation functional type (and their interaction) as factors. 359Lawns were excluded from this analysis. Finally we evaluated whether FF 360nematodes, BF nematodes and NCR were correlated, respectively, with PLFA 36118:2ω6, bacterial PLFA and the PLFA fungal/bacterial ratio, using Pearson 362correlation.

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Microbial (based on the relative abundance of PLFA biomasses) and Microbial (based on the relative abundance of PLFA biomasses) and Microbial community structures were assessed using nonmetric MMDS). The earthworm community was Seedominated by one species, resulting in low species richness and deemed Grunsuitable for NMDS. For the NMDS analysis we used Euclidean distances for SeRLFAs and Bray-Curtis distances for nematodes. The "envfit" function was used Seyto assess the significance of the relationship between these communities and Tothe environment (factors: city, park age, plant functional type; abiotic 371parameters: pH, % soil C, % soil N, C/N ratio, soil OM, % soil moisture). To test

372our specific hypotheses, NMDS analyses were performed on three different
373datasets; i) we included all park samples to evaluate the effects of plant
374functional type and park age on microbial and nematode communities; ii) we
375included only trees belonging to intermediate and old parks to focus on
376differences mediated by tree type; iii) we included only old deciduous and
377evergreen trees in Lahti parks and control forests in order to specifically
378explore differences between urban parks and natural forest. R scripts and data
379are provided as supplementary material (Supplementary material, S1-S25).
All statistical analyses were performed in R version 3.2.1. (R Core Team,

3812015), using the packages vegan (Oksanen et al. 2007), lme4 (Bates et al. 3822014), nmle (Pinheiro et al. 2007) and car (Fox et al. 2011).

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3853. Results

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3873.1. Microbial biomass and community composition

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389The GLMM models provided little evidence for systematic responses in PLFA-390based total microbial, bacterial and fungal biomasses (18:2ω6) to plant 391functional type or park age (Table 1, Fig. 1). Rather, these effects highlighted 392interactions between park age and plant type: PLFA biomasses (total, bacterial, 393fungal) were lowest under deciduous trees in young parks, yet highest under 394deciduous trees in old parks (Fig. 1 a, b). Total microbial and bacterial 395biomasses were mainly negatively correlated with the C/N-ratio (Table 1, Fig. 3962a) and positively correlated with % soil C (Fig. 2b). The fungal marker 397responded positively to % soil N (Table 1, Fig. 2c).

In old Lahti parks, total microbial and bacterial biomass 399(Supplementary material, Table S26) of evergreen and deciduous tree soils was 400about a third lower than in control forests (total microbial PLFA, age: $F_{1,16}$ = 40115.18, p < 0.01; bacterial PLFA, age: $F_{1,16}$ = 76,74, p < 0.01). Within forests and 402old parks, total microbial PLFA biomass did not differ under evergreen and 403deciduous trees, but bacterial biomass was higher under deciduous ($F_{1,16}$ = 5.13, 404p = 0.03) than evergreen trees. Fungal biomass (Supplementary material, Table 405S26) showed an interaction between land use type (control forest vs. urban 406park) and tree type ($F_{1,16}$ = 5.13, p = 0.03): fungal biomass was similar among 407deciduous trees in forest and parks, whereas under evergreens it was 408substantially higher in forests than in parks.

The PLFA-based fungi/bacteria (F/B) ratios suggest that, regardless of 410park age and plant type, all soils were bacteria dominated (Fig. 1c). However, 411the relative proportion of fungi over bacteria was higher under deciduous trees 412than evergreen trees and lawns, regardless of park age (Table 1, Fig. 1c). None 413of the measured soil parameters (Supplementary material, Table S26) 414correlated with the F/B-ratio, and the F/B ratio among old parks and forests 415did not differ (Table 1).

The ratio between gram-/gram+ bacteria did not respond to park age or 417to vegetation type, and did not differ between control forests and old Lahti 418parks. However, the gram-/gram+ ratio correlated positively with soil pH 419(Table 1, Fig. 2d).

420 Relative abundances of PLFAs were used in NMDS analyses to 421 investigate changes in the soil microbial communities in parks and control 422 forest. Although microbial biomasses responded, we observed distinct changes 423 in community composition only when the following datasets were analyzed; i) 424 all plant types and park ages and ii) control forest and old deciduous and 425 evergreen vegetation types in Lahti. When all plant types and park ages were 426 included in the NMDSs, plant functional type had no effect on the soil microbial 427 community composition in urban parks (Fig. 3, panel 1). However, park age 428 influenced the composition of these communities ($R^2 = 0.06$, p = 0.02) (Fig. 3, 429 panel 2). The soil microbial communities also differed ($R^2 = 0.11$, p = 0.01) 430 between the two cities. Environmental variables – primarily % soil C ($R^2 = 0.14$, 431 p < 0.01), % soil N ($R^2 = 0.12$, p < 0.01), the C/N-ratio ($R^2 = 0.08$, p = 0.02) and 432 soil OM ($R^2 = 0.07$, p = 0.02) – correlated with the soil microbial community 433 composition in the parks.

434 Comparing natural forest to deciduous and evergreen trees in old parks 435in Lahti showed that plant functional type clearly influenced ($R^2 = 0.14$, p = 4360.04) the soil microbial communities (Fig. 3, panel 5). However, the soil

437microbial community did not differ among old parks and control forests (Fig. 3, 438panel 6).

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4403.2. Nematodes

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442Bacterial and fungal feeding nematodes were more abundant under evergreen 443trees than under lawns (Table 1). However, while the abundance of bacterial 444feeders declined with park age, fungal feeders did not differ across park age 445classes. The abundance of bacterial feeders declined with park age and 446positively correlated with soil % soil C (Table 1). Fungal feeders negatively 447correlated with soil 0M (Table 1). Plant feeders and omnivorous nematodes, 448although not responding to plant type or park age, positively correlated with 449soil 0M and the C/N ratio (Table 1). When forests were compared with old 450parks in Lahti, bacterial feeding nematodes were more abundant ($F_{1,16}$ = 7.06, p 451= 0.01) under evergreen trees than under deciduous trees in old parks and 452control forests, while the other groups did not differ (see supplementary 453material, Table S26).

Plant type affected the nematode channel ratio (NCR) (F = 4.89, p = 4550.01). However, lawn and evergreen or deciduous and evergreen did not differ 456(Table 1, Fig. 4a), but rather, the differences can be attributed to those between 457deciduous trees and lawn. NCR negatively correlated with soil OM in the soil 458(Table 1). The NCR did not differ between old parks and control forest in Lahti. 459The maturity index (MI) showed a significant interaction between plant type 460and park age: when considering only deciduous trees, MI had the lowest values 461in young and old parks and highest values in intermediate parks (Table 1, Fig. 4624b). MI also positively correlated with the soil C/N-ratio (Table 1). MI was 463higher ($F_{1,16}$ = 7.92 p = 0.01) in old Lahti parks than forest controls (see 464supplementary material, Table S26).

We observed no relationships (Pearson correlations) between (i) the
466fungal marker and fungal feeding nematodes, (ii) bacterial biomass and
467bacterial feeding nematodes, and (iii) the PLFA fungal/bacterial-ratio and NCR.
The NMDSs revealed that nematode communities depended neither on
469plant functional type nor park age (Fig. 3, panels 7, 8). Park age did, however,

470have an effect when deciduous and evergreen trees in old and intermediate 471 parks were compared (Fig. 3, panel 10) ($R^2 = 0.63$, p < 0.01). Forests had 472distinct ($R^2 = 0.48$, p = 0.01) nematode communities compared to old Lahti 473parks (Fig. 3, panel 12). When all park samples were included in the analysis, % 474soil C was an important determinant of nematode community structure (R^2 = 4750.07, p = 0.04). However, when only every every and deciduous trees in 476intermediate and old urban parks were investigated, only soil moisture 477 correlated with nematode community structure ($R^2 = 0.15$, p = 0.04). Soil 478characteristics seemed to strongly control soil nematode communities (Fig. 1 479panels 11, 12) when forests were compared to old parks, with only deciduous 480and evergreen trees included in the analysis (moisture: $R^2 = 0.61$, p < 0.01; pH: $481R^2 = 0.56$, p < 0.01; OM: $R^2 = 0.39$, p = 0.01; % soil C: $R^2 = 0.39$, p = 0.01; % soil N: 482R² = 0.33, p = 0.02; C/N-ratio: R² = 0.57, p < 0.01). Nematode community 483 compositions differed among the two cities when the comparison was made 484between all parks ($R^2 = 0.13$, p < 0.01) and when deciduous and evergreen 485trees in old and intermediate parks in Helsinki and Lahti were included in the 486analysis ($R^2 = 0.19$, p = 0.04).

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4883.3. Earthworms

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490GLMM results of the anecic earthworm *Lumbricus terrestris* showed that its 491biomass positively correlated with park age (Table 1), with the highest 492biomasses in intermediate and old parks (Supplementary material, Table S2). 493Plant type was not associated with *L. terrestris* biomass. *L. terrestris* correlated 494negatively with soil OM content (Table 1). The *L. terrestris* biomass did not 495differ between old parks and control forests in Lahti.

The epigeic and endogeic earthworm biomasses responded to plant 497type (Table 1), with highest abundances in lawns (Supplementary material, 498Table S2). Here, soil pH negatively correlated with the biomass of earthworms 499(Table 1). The biomass of these earthworm taxa did not differ between old 500parks and control forests, nor between evergreen and deciduous trees in the 501two habitat types.

503

5044. Discussion

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5064.1. Effects of plant functional type on the soil biota

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508We hypothesized that different plant functional types – due to their 509documented divergent effects on soil characteristics and plant derived 510resources – will lead to changes in soil communities and the abundances of 511species and functional groups. Although overall communities were 512compositionally largely invariable, the biomasses and relative abundances of 513different microbial and nematode functional groups did respond to plant 514functional type.

515 Importantly, the PLFA-based F/B ratio was lowest in lawns, suggesting 516that bacteria dominated in lawn soils. Yet, to our surprise, soils under 517deciduous trees, and not evergreen trees, had the highest F/B and lowest NCR. 518NCR represents the relative abundance of bacterial feeders over fungal feeders, 519and thus mirrors the relative amount of bacteria over fungi in the soil (Yeates 5202003). In general, the relative amount of fungi over bacteria tends to be higher 521under deciduous and especially under evergreen trees than lawn. Thus, our 522data contrast those reported (e.g. Wardle et al., 2004). The reasons for this 523 remain unclear, and given our data, we can only speculate. For instance, a 524 higher amount of fine roots (Giardina et al. 2005) and decaying roots support a 525higher saprophytic fungal biomass (Hobbie 2006) in soil associated with 526deciduous trees than evergreen, which could explain why soils associated with 527old lindens showed such a high F/B ratio. Another important factor potentially 528explaining the similar F/B ratio under lawns and evergreen trees may be the 529 continuous disturbance present in urban parks; in particular, soil compaction 530can lower the F/B ratio (Hedlund et al., 2003).

531 Our third hypothesis focused on plant type mediated changes in higher 532trophic groups, such as nematodes and earthworm. GLMM analyses focusing on 533microbial and nematode feeding guilds revealed clear responses to plant type, 534although in some cases the response was seemingly mediated largely through 535age and soil properties of the parks. Total microbial biomass, as well as

536bacterial and fungal biomasses separately were highest under deciduous trees 537in old parks, whereas no such effect was detectable in younger parks. Bacterial 538feeding nematodes and fungal feeding nematodes were more abundant under 539evergreen and deciduous trees than under lawn. Nematode communities often 540vary according to vegetation, as a direct result of plant-provided resources and 541 indirectly through changes in the quality and quantity of the microbial fauna 542controlled by plant litter and exudate production. The lower abundance of 543 nematodes in lawns may be due to diminished predation/consumption by 544earthworms (Dash 1980), which were less abundant under trees than lawns. 545Moreover, Yates (1981) reported complementary dynamics between 546earthworms and bacterial feeding nematodes, suggesting that nematodes and 547earthworms can compete for resources, with both using microbial biomass as 548food. In line with other studies, total microbial and bacterial biomasses in this 549study were also positively correlated with % soil C. Soil % C was also 550correlated with nematode abundance (both FF and BF) and community 551 structure. Taken together, this is a clear indication that, just as in non-urban 552 soils, the availability of carbon is a key determinant of microbial and secondary 553 consumer biomass in urban soils, but that the effects appear only over time.

554 Contrary to previous findings (Lauber et al. 2009; Hui et al. 2017a), the 555soil microbial (PLFA) community did not correlate with pH. In contrast, the 556gram-/gram+ ratio did. Even though these two groups are not strict functional 557groups, they responded differently to soil pH; and in agreement with previous 558studies, gram- bacteria were relatively more abundant at high pH (Wang et al. 5592016). Further, our results also corroborate Hui et al. (2017a), who showed 560that microbial communities (characterized with high throughput sequencing 561methods) in these urban parks were distinguished by plant functional types 562and correlated with % soil N and pH.

563

5644.2. Effects of park age on the soil biota

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566Our fourth hypothesis stated that time since the establishment of a park (park 567age as well as tree age) will drive changes in the soil microbial and invertebrate 568communities. Our previous study, conducted in the same parks, but using high

569throughput sequencing, showed that microbial communities differed 570 compositionally between young parks and old / intermediate parks (Hui et al., 5712017a). Consistent with those results, our NMDS analyses of microbial 572 communities separated old and intermediate parks from young parks. This 573suggests that the soil microbial community reaches a stable structure between 57410 and 50 years after park construction. Soil microbial community composition 575in old parks overlapped with that in control forests. However, microbial 576biomass was substantially higher in forests than in old parks. This is likely due 577to frequent disturbances in the latter system (Pickett and Cadenasso 2009). 578Malmivaara-Lämsä and Fritze (2003) reported that human soil trampling in an 579urban boreal forest did not affect microbial community composition. 580Accordingly, we also observed that microbial community composition of old 581 parks approximates the non-disturbed forest sites, albeit microbial abundance 582is diminished in urban parks compared to forest sites. However, for logistical 583 reasons, we sampled the urban parks and control sites (forest) at different 584times. Microbial biomass can change across seasons, with peaks in the spring 585and autumn (Diaz-Ravina et al. 1995). Nevertheless, the control sites also had 586higher soil OM and %C (Setälä et al. 2016) suggesting that the observed 587microbial biomass differences are unlikely mere artifacts attributable to 588different sampling times. In addition to biomass, microbial community 589composition can be seasonally dynamic (Moore-Kucera and Dick 2008), but 590despite such potential seasonal dynamics, the communities in control and 591urban park sites overlapped.".

592 Contrary to our expectations, the relative amount of fungi over bacteria 593did not increase as a function of park age. Similarly, NCR, describing the 594relative abundance of bacterial-feeding over fungal-feeding nematodes, 595remained mostly invariant across all park age classes. In successional systems, 596such as agricultural systems under restoration, the F/B ratio increases (Zeller 5972001, Bailey 2002) as a result of changes in soil parameters (Zeller et al. 2001) 598and reduced disturbance (Hedlund et al. 2003). However, in urban parks the 599normal successional course is arrested and disturbance is continuous, possibly 600preventing increases in the relative abundance of fungi. Disturbance may also 601be the reason behind the invariant MI (Nematode Maturity Index) among the 602parks of different ages. Interestingly, MI, which is regarded as a good indicator 603of ecosystem maturity, was higher in old parks compared with forest. In a 604study on the response of a riparian nematode community along a rural-urban 605transect, Pavao-Zuckerman et al. (2007) reported no differences in MI among 606sites, suggesting that the level of pollution was not high enough to influence MI. 607Also in our case, the levels of metal pollution were below ecologically relevant 608values (Setälä et al. 2017), but it is still unclear why MI values in forests are 609lower than in parks.

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We were also unable to detect any shifts in the abundances of different 611nematode functional groups. Similarly to our speculations above, here too 612factors such as disturbance and litter removal may greatly contribute in 613altering resource availability to the soil biota, thus homogenizing these taxa 614across park ages. This could also explain the lack of response to plant traits and 615park age in plant-feeding nematodes, although plant root volumes likely 616increased substantially with time since park construction and establishment.

Nematode communities did respond to park age, but only in the 618rhizospheres of the two tree types; yet no changes in abundance with park age 619were observed. Nematodes can respond to changes in biotic and abiotic soil 620conditions (De Goede and Bongers 1994, Hanel 1995). Therefore, it is not 621surprising that we observed an age related shift in the nematode community 622only in association with trees, i.e. plant types which with time resulted in the 623clearest soil property changes in our study parks (see Setälä et al., 2016), and 624not in lawns.

The nematode community was also extremely responsive when we 626compared old parks and control forest, responding in parallel with the 627microbial communities as reported in Hui et al. (2017a). The nematode 628community structure was also correlated with changes in the soil 629characteristics, which in turn were modified by the different plant functional 630type and time. Community composition correlated with changes in abiotic soil 631parameters, such as pH, C, N and soil moisture. Again, this shows that 632nematodes are sensitive indicators (Neher 2001) of temporally mediated 633changes in soil characteristics. Contrary to previous findings, nematode 634densities were similar in rural forests and in urban parks. Pouyat et al. (1994)

635suggested that a decline in nematode populations in urban soils might be 636linked to higher heavy metal concentrations in urban soils. On the other hand, 637Ohtonen (1992) found no link between metal pollution and nematode density. 638It is possible that pollution, and in particular metal loads, in urban soils in 639Helsinki and Lahti were not extreme enough to affect the soil biota.

640 Abundance of the anecic earthworm L. terrestris was lowest in young 641 parks, increased with park age and reached a plateau in intermediate parks. 642The low abundance of earthworms in young parks can be attributed to two 643 factors: i) lack of time for earthworms to colonize newly constructed habitats, 644and/or ii) the inhospitality of the newly built park soil as a habitat. As 645 resources such as C and N increase in our park soils (see Setälä et al., 2016), so 646too does earthworm biomass. Similar trends in urban parks were observed by 647Smetak et al. (2007). This can have important implications since earthworms 648can ameliorate negative soil characteristics, compensating for and even 649reducing soil compaction with time. It is interesting to note that epigeic and 650endogeic earthworms did respond to pH variation, being more abundant at low 651pH than at high pH. Earthworms are quite sensitive to pH (Edwards and 652Bohlen, 1996). However, variation in pH among parks and plant types was 653rather narrow (Setälä et al., 2016) and among the earthworm species that we 654 recorded, Lumbricus rubellus and Aporrectodea caliginosa were more 655abundant. Considering that pH is highly correlated with park age (Setälä et al., 6562016), the significant relationship between earthworm abundance and pH 657 could also be interpreted as the result of a relationship between park age and 658earthworm abundance.

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661 Conclusions

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663Our data show that microbial biomass in natural forests is much higher than in 664urban parks, irrespective of their age. Thus, the urban greenspaces 665investigated unlikely approximate ecosystem properties or functions of the 666surrounding non-urban areas. Yet, our study provides strong evidence that 667deciduous trees support a greater microbial biomass with time since urban

668green space establishment. Deciduous trees also had the highest relative 669amount of fungi over bacteria in urban parks, that evergreen trees would 670promote fungal rich soil microbial communities. Not only did microbial 671biomass increase with age in urban parks, but so too did the abundance of 672secondary consumers, such as earthworms. These data suggest that although 673urban parks do not approximate natural forests, likely as a consequence of the 674maintenance/disturbance regime they experience, the soil microbial and 675invertebrate communities respond to vegetation and edaphic shifts over time. 676Nevertheless, further studies are needed to assess the correlation and type of 677ecosystem services provided by these trophic groups.

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680Acknowledgments

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1056Figure and Table Captions

Fig. 1 The effects of plant functional type and park age on a) total microbial 1059PLFAs, b) fungal PLFA (18:2 ω 6) and c) the PLFA fungal to bacterial ratio. Mean 1060values ± SE are presented.

Fig. 2 Relation between a) the C/N ratio and predicted values of total microbial 1063PLFAs, b) % soil C and predicted values of total microbial PLFAs, c) % soil N 1064and predicted values of fungal PLFA (18:2 ω 6) and d) pH and predicted values 1065of gram-/gram+ bacteria.

Fig. 3 NMDS plots for bacterial (1-6, on the left) and nematode (7-12, on the 1068right) communities. Microbial communities were grouped by plant functional 1069type (1, 3, 5) and age (2, 4, 6), and nematode communities were grouped by 1070plant functional type (7, 9, 11) and age (8, 10, 12) under lawn, deciduous and 1071evergreen trees in young, intermediate and old parks. Significant effects (p < 10720.05) are indicated with a check mark in the upper panels and significant 1073vectors are shown in the NMDS plots.

1075Fig. 4 The effects of plant functional type and park age on a) the nematode1076channel ratio (NCR), and b) the nematode maturity index (MI). Mean values ±1077SE are presented.

Table 1. GLMM results for PLFAs, nematodes and earthworms. For PLFAs the 1080response variables included total microbial PLFA, bacterial PLFA, fungal PLFAs 1081(18:2 ω 6, saprophytic fungi), fungal to bacterial ratio (fungi/bacteria) and gram + to 1082gram – ratio (gram+/gram-). For nematodes the response variables included 1083bacterial feeders, fungal feeders, plant feeders, predators and omnivorous, the 1084maturity index (MI) and the nematode channel ratio (NCR). For earthworms the 1085response variables were: *Lumbricus terrestris* and other earthworms. For each 1086variable we reported the coefficient, standard error and p-value. Significant effects 1087(p < 0.05) are highlighted in bold and significant interactions are indicated with an 1088asterisk. Young evergreen trees are in the intercept.

Supplementary material, Table S2. Means (± 1 SD) of the following PLFAs: 1091total microbial, bacterial, fungal PLFAs (18:2ω6, saprophytic fungi), 1092fungal/bacterial ratio, gram/gram+ ratio, nematodes: bacterial feeders, fungal 1093feeders, plant feeders, predators and omnivores, maturity index (MI), 1094nematode channel ratio (NCR), earthworms: *Lumbricus terrestris* and other 1095earthworms, for the three different park ages (young, intermediate and old) 1096and plant functional type (evergreen, deciduous, lawn). Values for Lahti old 1097parks and control forest (evergreen and deciduous trees) are also presented.

1122Table 1. GLMM results for PLFAs, nematodes and earthworms. For PLFAs the 1123response variables included total microbial PLFA, bacterial PLFA, fungal PLFAs 1124(8:2 ω 6, saprophytic fungi), fungal to bacterial ratio (fungi/bacteria) and gram + to 1125gram – ratio (gram+/gram-). For nematodes the response variables included 1126bacterial feeders, fungal feeders, plant feeders, predators and omnivorous, the 1127maturity index (MI) and the nematode channel ratio (NCR). For earthworms the 1128response variables were: *Lumbricus terrestris* and other earthworms. For each 1129variable we reported the coefficient, standard error and p-value. Significant effects 1130(p < 0.05) are highlighted in bold and significant interactions are indicated with an 1131asterisk. Young evergreen trees are in the intercept.

	variable	intercept	lawn	deciduous	intermediate	old	age x functional group	рН	% soil C
	Total microbial PLFAs	6.821	0.041	0.040	0.081	0.068			0.064
		0.220	0.092	0.092	0.103	0.100	*		0.023
		<0.001	0.447	0.437	0.791	0.675			<0.00
		5.951	0.091	0.007	0.060	0.046			0.070
	Bacterial PLFAs	0.228	0.094	0.094	0.105	0.102	*		0.023
		<0.001	0.336	0.942	0.565	0.655			<0.00
	Fungal PLFAs	3.414	0.129	-0.156	0.248	-0.320			-0.162
PLFAs		0.290	0.266	0.261	0.272	0.274	*		0.085
		<0.001	0.627	0.549	0.362	0.243			0.056
	Fungi/bacteria	17.296	-4.994	8.044	-2.107	1.353			
		3.141	3.444	3.417	3.417	3.417			
		<0.001	0.147	0.019	0.537	0.692			
	Gram-/gram+	0.327	0.007	0.037	-0.064	-0.133		0.105	
		0.320	0.050	0.052	0.056	0.053		0.051	
		0.308	0.897	0.474	0.260	0.018		0.040	
	Bacterial feeding	8.676	-2.038	-1.782	-2.121	-1.668			0.475
Nematodes		2.279	0.847	0.832	0.860	0.852			0.218
		<0.001	0.016	0.321	0.013	0.050			0.029
	Fungal feeding	5.58	-1.322	-0.245	-0.35	-0.761			
		0.963	0.492	0.485	0.55	0.548			
		<0.001	0.01	0.612	0.524	0.165			
	Plant feeders	4.303	-1.030	-1.386	0.176	0.895			
		1.335	0.918	0.886	0.929	0.924			
		<0.001	0.262	0.118	0.849	0.333			
	Predators and omnivorous	-0.082	0.161	0.770	0.770	-0.109			
		1.049	0.438	0.438	0.462	0.463			
		0.937	0.712	0.104	0.095	0.813			
	MI	0.550	0.209	0.090	0.033	-0.074			
		0.119	0.059	0.060	0.064	0.064	*		
		<0.001	<0.001	0.134	0.600	0.246			

		0.007	< 0.001	0.027	0.480	0.090	0.015
Earthworms –	Other earthworms	1.355	0.190	0.196	0.226	0.217	0.200
	Lumbricus terrestris	3.657	0.794	0.433	0.160	0.368	-0.486
		0.008	0.611	0.448	<0.001	0.007	0.032
		0.725	0.320	0.319	0.406	0.404	0.123
		1.920	-0.163	-0.242	1.452	1.097	-0.265
		<0.001	0.121	0.084	0.212	0.848	
	NCR	0.094	0.055	0.055	0.066	0.065	

Table 2 Means (± 1 SD) of the following PLFAs: total microbial, bacterial, fungal PLFAs (18:2ω6, saprophytic fungi), fungal/bacterial ratio, gram/gram+ ratio, nematodes: bacterial feeders, fungal feeders, plant feeders, predators and omnivores, maturity index (MI), nematode channel ratio (NCR), earthworms: *Lumbricus terrestris* and other earthworms, for the three different park ages (young, intermediate and old) and plant functional type (evergreen, deciduous, lawn). Values for Lahti old parks and control forest (evergreen and deciduous trees) are also presented.

	Variable		Young	SD	Intermediate	SD	Old	SD	Old Lahti	SD	Control	SD
PLFA, µg g ⁴ soil dw	Total microbial PLFAs	Evergreen	380.09	132.83	526.96	208.32	417.16	93.44	318.82	64.09	842.19	319.05
		Deciduous	271.04	93.78	530.15	146.41	616.97	347.73	669.85	439.87	957.27	281.94
		Lawn	460.25	165.97	447.87	128.15	473.87	185.37				
		Evergreen	172.85	66.11	223.54	88.39	181.19	50.83	160.42	46.90	799.71	306.51
	Bacterial PLFAs	Deciduous	116.54	43.01	224.04	59.13	241.89	112.35	270.50	105.90	982.63	252.72
		Lawn	211.31	90.41	203.88	60.37	218.14	82.75				
	Fungal PLFAs	Evergreen	30.98	13.60	45.55	25.43	26.59	15.87	21.74	14.16	127.29	99.25
		Deciduous	24.65	9.00	48.12	28.67	111.03	125.14	141.05	156.35	127.97	100.76
		Lawn	34.26	12.77	23.88	11.22	26.16	16.65				
		Evergreen	20.33	12.39	21.17	11.73	15.96	11.25	14.05	8.43	18.39	15.22
	Fungi/bacteria	Deciduous	22.65	9.65	21.89	10.83	37.53	26.93	43.87	32.64	12.91	8.83
		Lawn	18.76	8.98	11.88	5.29	11.83	4.55				
		Evergreen	2.98	0.45	2.77	0.31	2.49	1.02	3.07	1.02	3.14	0.83
	Gram+/gram-	Deciduous	3.35	0.79	2.70	0.44	2.68	0.53	2.78	0.57	3.53	0.62
		Lawn	2.79	0.40	2.66	0.26	2.73	0.35				
	Bacterial feeders	Evergreen	139	113	100	85	93	47	64	33	74	31
		Deciduous	106	84	51	60	85	73	34	30	38	6
		Lawn	79	83	78	63	49	51				
	Fungal feeders	Evergreen	25	23	18	14	12	12	11	10	40	53
		Deciduous	24	25	20	30	13	9	12	7	13	8
		Lawn	11	11	10	14	7	6				
		Evergreen	63	63	54	55	86	67	68	36	45	27
	Plant feeders	Deciduous	37	32	43	53	48	33	47	36	16	24
Nematodes 10 g		Lawn	129	283	69	55	136	285				
¹ soil	Predators and omnivorous	Evergreen	9	8	12	9	7	11	4	4	19	15
		Deciduous	15	15	14	9	10	9	12	12	18	18
		Lawn	12	14	8	5	9	11				
	м	Evergreen	2.17	0.11	2.30	0	2.35	0.23	2.42	0.03	2.15	0.20
		Deciduous	2.24	0.43	2.84	0.38	2.23	0.44	2.56	0.18	2.34	0.27
		Lawn	2.37	0.44	2.43	0.64	2.54	0.46				
	NCR	Evergreen	0.82	0.13	0.71	0.25	0.79	0.17	0.86	0.08	0.71	0.30
		Deciduous	0.77	0.19	0.85	0.12	0.88	0.09	0.70	0.17	0.76	0.12
		Lawn	0.88	0.12	0.88	0.10	0.87	0.10				
Earthwoms	Lumbricus terrestris	Evergreen	34	43	88	60	62	65	99	65	115	133
		Deciduous	15	32	63	49	66	67	102	77	97	100
		Lawn	27	30	54	67	43	37				
g m ⁻²		Evergreen	8	10	11	12	9	12	12	15	14	11
	Other earthworms	Deciduous	9	13	19	32	19	13	25	13	50	48
		Lawn	16	16	36	44	35	40				

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1157Fig. 3

