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| Author(s) | Kagiya, Shinnosuke; Utsumi, Shunsuke |
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| 1 | Spatial heterogeneity in genetic diversity and composition of bacterial |
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| 4 | Shinnosuke Kagiya ^{1*} , Shunsuke Utsumi ² |
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| 6 | ¹ Graduate School of Environmental Science, Hokkaido University, North 10, West 5, |
| 7 | Sapporo, Hokkaido 060-0810, Japan (e-mail: slotapir-1035@fsc.hokudai.ac.jp) |
| 8 | ² Field Science Center of Northern Biosphere, Hokkaido University, North 9, West 9, |
| 9 | Sapporo, Hokkaido 060-0809, Japan (e-mail: utsumi@fsc.hokudai.ac.jp) |
| 10 | |
| 11 | |
| 12 | *Corresponding author |
| 13 | Shinnosuke Kagiya |
| 14 | Graduate School of Environmental Science, Hokkaido University, North 10, West 5, |
| 15 | Sapporo, Hokkaido 060-0810, Japan |
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17 Abstract

18 Aims

19 Revealing genetic diversity in a root nodulation symbiosis under field conditions is 20 critical to understand the formation of ecological communities of organisms associated 21 with hosts and the nitrogen cycle in natural ecosystems. However, our knowledge of 22 genetic diversity of bacterial mutualists on a local scale is still poor because of the 23 assumption that the genetic diversity of mutualistic bacteria is constrained by their hosts.

24

25 Methods

We thoroughly investigated genetic diversity of *Frankia* in a local forest stand. We collected root nodules from 213 *Alnus hirsuta* seedlings covering the spatial range of the continuous population, which means that *Alnus* individuals occurred in a relatively homogeneous distribution in a continuous forest. Then, a phylogenetic analysis was performed for the *nif*D-K IGS region, including global *Frankia* sequences from *Alnus* hosts.

32

33 Results

The genetic diversity of *Frankia* detected even on a local scale measured as high as that shown by previous studies conducted on a regional scale. Moreover, a genetic structure analysis revealed a spatially mosaic-like distribution of genetic variation in *Frankia* despite the small spatial scale.

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39 Conclusions

The genetic diversity and composition of bacterial mutualists are heterogeneous on a local scale. Our findings demonstrate that genetically different bacterial symbionts simultaneously interact with a single host population and interaction partnerships spatially vary. The standing variation could produce dynamic ecological and evolutionary outcomes in a heterogeneous forest ecosystem.

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46 Key words

47 Alnus hirsuta, Frankia, genetic diversity, local scale, nifD-K IGS region, nitrogen-fixing

48 bacteria, root nodule symbiosis

50 Introduction

Most terrestrial plants interact with microsymbionts in the rhizosphere. Root nodule 51 52 symbiosis between plants and nitrogen-fixing bacteria has a significant impact on the nitrogen cycle in terrestrial ecosystems. In both evolutionary and applied biology, genetic 53 54 variation in rhizobial mutualism has attracted considerable attention (Barrett et al. 2012; Miller and Sirois 1982; Robinson et al. 2000). Typically, experimental inoculation studies 55 using different strains of bacterial symbionts have reported different effects of the 56 57 mutualistic interactions, such as growth, nitrogen contents, and leaf size of the host plants, as well as nitrogen-fixation activity of the bacteria (Dillon and Baker 1982; Hooker and 58 Wheeler 1987; Prat 1989; Sellstedt et al. 1986). In other words, it has been widely 59 60 acknowledged that intraspecific variation in mutualistic nitrogen-fixing bacteria greatly affects host plant performance in terms of growth, survival, reproduction, and defense 61 62 (Ballhorn et al. 2017; Barrett et al. 2012; Dean et al. 2014; Miller and Sirois 1982; Pahua et al. 2018; Prat 1989; Robinson et al. 2000), which in turn influences nitrogen-cycling 63 processes. Thus, the knowledge of genetic variation in rhizobial mutualism is essential to 64 65 understand not only the creation and maintenance of a symbiosis but also wider ecosystem processes. 66

Nitrogen-fixing *Frankia* bacteria form nodules on the roots of actinorhizal plants. Many studies have focused on legume–rhizobia symbioses due to the agricultural importance, while interactions between actinorhizal plants and *Frankia* have been poorly examined. Whereas, many legume plants are herbaceous, most actinorhizal plants are woody (Wheeler *et al.*, 2008), and actinorhizal symbiosis is a major contributor to the global nitrogen budget in forest ecosystems, playing a dominant role in forest succession, especially in temperate and polar ecosystems (Kucho et al. 2010; Lawrence et al. 1967). 74 Therefore, the genetics of actinorhizal plants-Frankia bacteria mutualism may be more 75 important than legume-rhizobia interactions on ecosystem processes in non-agricultural 76 fields. Unraveling spatial structure of genetic diversity of mutualistic bacteria in nonagricultural field is likely to be important toward an understanding of nitrogen cycling, 77 78 associated community dynamics, and coevolutionary dynamics in nodulation symbiosis. Nevertheless, there is only a small body of literature on the genetic diversity of Frankia 79 in natural ecosystems (Anderson et al. 2009; Ben Tekaya et al. 2018; Benson and Hanna 80 1983; Clawson et al. 1998; Clawson et al. 1999; Huguet et al. 2001; Kennedy et al. 2010; 81 Mishra et al. 2015; Pozzi et al. 2018a; Pozzi et al. 2015; Pozzi et al. 2018b; Ridgway et 82 83 al. 2004; Roy et al. 2017; Simonet et al. 1994; Simonet et al. 1989; Vanden Heuvel et al. 84 2004; Wilcox and Cowan 2016).

Researchers recently have begun to reveal Frankia genetic diversity in wide 85 geographic ranges. For example, Nouioui et al. (2014) investigated the genetic structure 86 87 of Frankia on a global scale. The maximum distance of their study sites was approximately 19,000 km. Kennedy et al. (2010) and Wilcox and Cowan (2016) surveyed 88 in regions where the maximum distances were 336.8 km and 165.0 km respectively. Some 89 previous studies have investigated genetic diversity of Frankia on small spatial scales 90 91 and/or from a single host species (Benson and Hanna, 1983; Clawson et al. 1999; Khan et al. 2007; Mishra et al. 2015; Pokharel et al. 2011; Pozzi et al., 2015; Simonet et al. 92 1994; Simonet et al. 1989). However, the above studies assessed genetic diversity of 93 Frankia with small sample size per study sites. Therefore, distribution of Frankia genetic 94 95 diversity within a small spatial scale has been overlooked.

The most important reason why the knowledge of genetic diversity of *Frankia* is still limited on a small spatial scale may be the assumption that the genetic diversity of 98 mutualistic partners is low on a local scale and in a single host species. The traditional 99 mutualistic theory has suggested that the genetic diversity of mutualistic partners is 100 constrained by hosts and could be decreased by the hosts' stabilizing mechanisms, such 101 as partner choice and sanction (Archetti et al. 2011; Heath and Stinchcombe 2014).

102 It should also be noted that most previous studies compared genetic variation in Frankia among host plant species. This is because the focus has mainly been on the 103 symbiotic host specificity in this mutualism (i.e., differences in the infectivity of rhizobial 104 symbionts among host plant species; Baker 1987; Jiabin et al. 1985; Mirza et al. 2009). 105 106 In natural ecosystems, different host species commonly associate with phylogenetically 107 different Frankia strains (Du and Baker 1992; Normand et al. 1996). For this reason, most 108 previous studies have compared genetic variation of Frankia among multiple host species to an understanding of coevolutionary history and effects of actinorhizal mutualism. 109

However, the knowledge of genetic diversity of Frankia on a small spatial scale (e.g., 110 111 seed dispersal range: many seeds of Alnus individuals dispersed within c. 140 m along a river (Cunnings et al. 2016)) should be required to understand outcomes of considerable 112 variation in current actinorhizal ecological interactions, such as the effectiveness of the 113 114 bacteria in growth and survival of the host plants. This is because effects of rhizobial mutualism often depend not only on genetic variation of mutualistic bacteria but also on 115 intraspecific variation of host species (Caldwell 1966; Hayashi et al. 2012; Heath and 116 117 Tiffin 2007; Yamakawa et al. 2003). For example, nodulation rates of Frankia strains could also differ among intraspecific host individuals (Hahn et al. 1988). In fact, large 118 genetic variation in a host plant population, including actinorhizal and legume species, is 119 also ubiquitous in a natural forest stand (Ager et al. 1993; Kagiya et al. 2018; King and 120 Ferris 1998; Wickneswari and Norwati 1993) Our previous study revealed large genetic 121

variation in a single *Alnus* species in a continuous natural forest ($20 \text{ km} \times 70 \text{ km}$; Kagiya et al., 2018). Leaf traits, such as C:N ratio, leaf mass per area (LMA), and herbivory rate, varied with the genetic variation and localities with the forest. Therefore, we should pay attention to genetic diversity of rhizobial bacteria and its spatial heterogeneity on a small spatial scale, which may be crucial to determining outcomes of ecological interaction between rhizobial bacteria and actinorhizal host plants under natural ecosystem conditions.

In this study, our goal is to elucidate spatial structure in genetic diversity and 129 composition of mutualistic bacteria even in a local population of single host species (a 130 131 single-host–population scale). Specifically, we sought to answer the following questions: 132 (1) how diverse genetically is Frankia bacteria within and across local sites, (2) do the genetic compositions of Frankia bacteria differ among local sites, and (3) what is the 133 spatial genetic structure of *Frankia* in a natural forest. For the purposes of the study, we 134 135 focused on the A. hirsuta-Frankia symbiosis in a natural forest in northern Hokkaido, Japan. Actinorhizal populations in this forest region are dominated by a single Alnus 136 species, A. hirsuta. The genetic variation of A. hirsuta within the forest has been 137 138 determined by a genome-wide analysis (Kagiya et al. 2018). We continuously investigated the genetic diversity of *Frankia* bacteria in the *Alnus* populations at intervals 139 140 of c. 100 m (the maximum distance between host populations is 43.476 km; 213 seedlings in total). 141

142

143 Materials and Methods

144 Host species and nitrogen-fixing bacteria

145 Alnus hirsuta (Betulaceae; Alnus incana ssp. hirsuta Spach; Chen and Li 2004; Ren et al. 2010) is a deciduous broadleaf tree and an early successional species. It is widely 146 147 distributed in temperate riparian forests of Japan, northeastern China, Korea, and Russia. Alnus trees have the following characteristics as foundation species in a riparian forest 148 149 ecosystem (Ellison et al. 2005; 2010): (1) they are a dominant species in early succession forests, (2) they support diverse arthropod species (Kagiya et al. 2018; Nyeko et al. 2002), 150 151 and (3) they are actinorhizal species able to form partnerships with nitrogen-fixing 152 actinobacteria, Frankia sp. (Frankiaceae) forming nodules in their roots, which seem to 153 greatly affect ecosystem processes such as nutrient cycling. Frankia bacteria have the 154 ability to convert atmospheric nitrogen into ammonia, and are free-living soil microbes 155 but some are obligate symbionts (Benson and Dawson 2007).

156

157 Root nodule sampling

Our study sites are located in and around the Uryu Experimental Forest (44° 030–290N, 158 159 142° 010–200E) of Hokkaido University in northern Hokkaido, Japan. This experimental 160 forest is a continuously mixed conifer-broadleaf forest of c. 25000 ha. One nodule was collected from each of the roots of 213 A. hirsuta seedlings (DBH: < 2 cm) from five 161 riparian areas of the forest (BT, DRE, DRW, SE, and UT; the maximum distance between 162 163 our areas was 43.5 km; Fig 1) because the host trees are mainly found along rivers, and streams are considered one of the primary dispersal pathways of Frankia bacteria (Arveby 164 and Huss-Danell 1988; Huss-Danell et al. 1997). Seedlings from which we collected root 165 166 nodules were selected from 17 sites from the riparian areas. Sampling root nodules from A. hirsuta seedlings would allow us to collect samples continuously from a whole forest 167 and to estimate genetic diversity and composition of *Frankia* which interact with a single 168

169 host plant population at present. Sampling was conducted from June to September 2016.

170 The distance between sampling points in each site was more than 100 m. *Alnus hirsuta* is

171 the only actinorhizal species in this forest.

172

173 Molecular Analyses

174 The collected nodules were surface-sterilized using 10% (v/v) Clorox bleach. DNA was 175 extracted from root nodules using DNeasy Blood & Tissue Kit (Qiagen). The nodules were sliced using sterilized razor blades and crushed using sterilized homogenization 176 sticks. The crushed lobes were heated to 37 °C for 30 min. with a 25 µl Proteinase K. 177 178 Polymerase chain reaction (PCR) was performed to amplify a 496-bp fragment of a *nif*D-179 K intergenic spacer region with the *Frankia*-specific primer pair, *nif*D1310frGC (5'-CGC CAG ATG CAC TCC TGG GAC TAC T-3'), and nifKR331frGC (5'-CGG GCG AAG 180 TGG CTG CGG AA-3'). We focused on intrageneric variation of Frankia based on the 181 nifD-K IGS region. The genetic marker is considered to be one of the useful genetic 182 183 markers for resolution at the species level of Frankia because the genetic region includes 184 higher variable than ribosomal RNA (Anderson et al. 2009; Mishra et al. 2015). PCR amplification was performed as follows: 1 cycle at 95 °C for 2 min, followed by 35 cycles 185 186 of 95 °C for 1 min and 64 °C for 5min, and a final step of 1 cycle at 72 °C for 5 min. All successful PCR products were cleaned using an ExoSAP master mix containing 0.5 µl 187 188 Exonuclease I (TaKaRa), 0.5 µl Shrimp Alkaline Phosphatase (SAP; New England BioLabs), and 2.0 µl sterile deionized water. Incubation using a thermal cycler was 189 conducted with ExoSAP at 37 °C for 20 min and at 80 °C for 15 min. These products 190 were sequenced with an automated sequencer (3730xl DNA Analyzer, Applied 191 Biosystems). DNA sequence chromatograms were manually checked using FinchTV 192

193 1.4.0 (Geospiza, Inc.; Seattle, WA, USA; http://www.geospiza.com), and sequences were
aligned using MEGA 7.0.21 (Kumar et al. 2016). Finally, nucleotides in obtained
sequences were checked to remove sequences of low reliability. In total, 201 sequences
were used for subsequent analyses.

197 Operational taxonomic unit (OTU) separation was performed to classify the 201 sequences at a 97.0%-threshold, using the CD-Hit program (Li and Godzik 2006). This 198 199 threshold was decided based on the statistical method detailed in Põlme et al. (2014). These sequence data were deposited in the DNA Data Bank of Japan (DDBJ) with 200 201 accession no. LC482655-LC482672. Phylogenetic trees were constructed using 202 Maximum Likelihood (ML; bootstrap analyses with 1000 replications; Fig 2) and Neighbor-Joining (NJ; bootstrap analyses with 10000 replications; Fig S1) based on the 203 Kimura 2-parameter evolutionary model (Kimura, 1980) with a discrete gamma 204 distribution selected by evolutionary model selection procedure in MEGA 7.0.21 (Kumar 205 206 et al. 2016). The phylogeny included the nifD-K locus of uncultured Frankia bacteria obtained from two Alnus species, A. incana (ssp. tenuifolia (Anderson et al. 2009) and 207 ssp. rubra) and A. viridis (Anderson et al. 2009), and nine varieties of Myrica rubra (var. 208 209 biji, var. baimei, var. dongkui, var. muye, var. shuimei, var. wandao, var. wumei, var. yangliu, var. zaoda; He et al. 2004), as well as a cultured ACN14a strain, whose host is 210 211 A. viridis (Normand et al. 2007). Other Frankia sequences from different actinorhizal 212 species, Hippöphae salicifolia (Mishra et al. 2015), three Coriaria species (C. myrtifolia, C. japonica, C. arborea; Nouioui et al. 2014), Elaeagnus angustifolia, Datisca glomerate, 213 and Casuarina equisetifolia (Normand et al. 2007), were also included as outgroups. 214 215 These sequences covered Frankia nifD-K sequences of almost actinorhizal hosts in GenBank. These sequences were obtained from GenBank. All positions with less than 216

95% site coverage were eliminated. Both ML and NJ phylogenetic trees were generated
with MEGA 7.0.21 (Kumar et al. 2016). To describe relationships between *Frankia* OTUs
and areas, we also generated the ML phylogeny with 1000 bootstraps, excluding the
sequences obtained from the database (Fig 4).

To analyze the spatial genetic structure of *Frankia* in a natural habitat, analysis of molecular variance (AMOVA) was performed with 9999 permutations, using GenAlex 6.5 (Peakall and Smouse 2012). The five riparian areas were divided into 2–5 sites each to analyze the genetic structure within the areas.

225

226 Analysis of *Frankia* diversity

To estimate spatial heterogeneity of the *Frankia* composition on a single-host-population 227 scale, Bray-Curtis dissimilarity index was calculated. The total number of each Frankia 228 OTU was used for the data set at the site- and area-level. The data set was standardized 229 to unified scale [0; 1], dividing by total number of each site/area, because sample sizes 230 231 were different among sites/areas. The significance of Frankia composition dissimilarity 232 among areas was analyzed using permutation MANOVA (PERMANOVA) with 9,999 permutations. To visually summarize the dissimilarity among sites, non-metric 233 234 multidimensional scaling (NMDS) in a two-dimensional space was performed. All calculations in above were performed using the R package vegan 2.5-6 (Oksanen et al. 235 236 2019) with the R 3.6.1 software.

To consider spatial autocorrelation in the *Frankia* OTU compositions, we calculated spatial distance among sites and areas. Spatial distance was calculated based on location of each site/area with Great Circle distance method, using the R package sp 1.3-1(Pebesma et al. 2018). The location was calculated centroid location as the averaged latitude and longitude of each *A. hirsuta* seedling. Spatial autocorrelation of *Frankia*compositions was analyzed by Mantel tests with 9,999 permutations. These tests were
performed using the vegan package.

244

245 **Results**

246 Genetic diversity of *Frankia* on a single-host–population scale

To classify Frankia in the study forest, we used OTU methods based on the genetic 247 similarity of the *nif*D-K loci. A total of 18 OTUs were obtained at a 97.0% similarity 248 based on the nifD-K loci of Frankia in the forest (Table S1, Fig 2). This 97.0% threshold 249 for OTUs is likely to be relevant to represent the phylogenetic relationship of Frankia 250 strains based on the *nif*D-K ITS region because sequences of all samples were clearly 251 252 clustered to each clade of single OTU (Fig S2). The phylogenetic trees indicated that 253 obtained OTUs widely spread in the almost range of Alnus-infective clade (Fig 2, S1). Each of the three most common OTUs, OTU01, OTU02, and OTU03, was placed into 254 phylogenetically different clades (Fig 2, S1). Thus, our result demonstrated that 255 genetically diversified strains co-occurred in the forest. In addition, both OTU06 and 256 OTU07 were genetically close to OTU02, and both OTU04 and OTU05 were genetically 257 close to OTU03. Some Frankia sequences in this study were phylogenetically close to 258 259 bacteria from A. incana ssp. tenuifolia or A. viridis, which were belonging to different clades between the host species (Anderson et al. 2009; 2013). Additionally, noted that 260 seven of these OTUs (OTU01-OTU07) were obtained from multiple samples, while the 261 rest (OTU08-OTU18) was rare singleton (Fig 4, S2). 262

To illustrate differences in *Frankia* OTU diversity (i.e., the total number of OTUs in each site) among the sites with the standardization of sample size, we generated a rarefaction curve for each site (Fig 3). OTU diversity was greater in BT, DRE, and DRW areas than in SE and UT areas. While the rarefaction curves of SE and UT areas show the saturation of the total OTU number, those of BT, DRE and DRW areas indicated there was likely to be still undetected OTUs in each area.

269

270 Differences in *Frankia* composition

The three most abundant OTUs (i.e., OTU01, OTU02, and OTU03) were commonly found throughout the entire sampling areas (Fig 4), indicating a sympatric coexistence of these haplotypes. However, other OTUs were localized to parts of different sampling areas. OTU04, OTU05, OTU06, and OTU07 were detected in seedlings from three riparian areas (BT, DRW, and DRE). These results indicated the spatial heterogeneity in *Frankia* compositions within a single-host–population scale.

This finding was supported by NMDS community ordination, in which *Frankia* compositions were diversified among areas within a single-host–population scale (Fig 5). The stress of the NMDS was 0.070, indicating a good representation of the data in twodimensional ordination plot. The significant differences in *Frankia* OTU compositions were detected (PERMANOVA; P < 0.05).

282

283 Spatial genetic structure of *Frankia*

The AMOVA indicated a significant genetic differentiation in *Frankia* genetic communities among riparian areas and sites (Table 1). In addition, no significant correlations were detected between the dissimilarity of *Frankia* compositions and spatial distance (at site level: r = -0.1037, P = 0.6967; at area level: r = 0.4897, P = 0.1833). This suggested that spatial structure of *Frankia* compositions was unlikely to be resulted from
spatial autocorrelation.

Overall, the spatial heterogeneity in *Frankia* genetic variation was due to the differences in both OTU diversity and composition of *Frankia* strains.

292

293 **Discussion**

294 This study clearly demonstrated that multiple Frankia genotypes coexist even in an area 295 within a single-host-population scale. Frankia genetic diversity in a single-hostpopulation scale is comparable with previous studies that analyzed the *nif*D-K genetic 296 region of Alnus-infection Frankia from multiple hosts and/or in regional scales. 297 Rarefaction analysis also showed that the existence of undetected OTUs is also expected 298 in some sites. Furthermore, differences in Frankia genetic diversity and composition were 299 detected even within the small spatial scale (Fig 4, 5). These results suggest that 300 actinorhizal host individuals within the population can interact with different Frankia 301 302 genotypes.

303

304 *Frankia* diversity and composition on small spatial scales

The maintenance mechanisms of sympatric coexistence of phylogenetically distant *Frankia* strains can contribute to understand the spatial heterogeneity in *Frankia* diversity and composition on local scales. Three factors can explain why various genotypes of mutualistic partners coexist in the same habitat (Heath and Stinchcombe 2014): (1) a different selection on each genotype of the partners by genetic variation in hosts (i.e., G 310 × G: genotype–by–genotype interactions), (2) genetic trade-offs between bacterial strains,
311 and (3) different functions of multiple *Frankia* genotypes.

312 First, in natural ecosystems, it is likely that the genetic structure of *Frankia* is greatly restricted by the hosts' phylogenetics (Anderson et al. 2009; Põlme et al. 2014; Pozzi et 313 314 al. 2018). Mutualistic benefits for host plants from root-nodulating symbionts also differ among different genotypes within a host species, as well as exhibiting interspecific 315 variation (Caldwell 1966; Hayashi et al. 2012; Heath and Tiffin 2007; Yamakawa et al. 316 2003). Therefore, the sympatric coexistence of phylogenetically distant *Frankia* strains 317 on a single-host-population scale may be at least partially explained by intraspecific 318 319 variation of the host plant A. hirsuta. Determining the effects of intraspecific variation in 320 a host species with $G \times G$ interactions in mutualism may be important to understanding how a stable coexistence of genetically diverse mutualistic partners is sustained on a small 321 spatial scale. 322

323 Second, genetic trade-offs between different mutualism-related traits, if existent, can contribute to the maintenance of a stable coexistence of different Frankia strains. For 324 example, if mutualistic efficiency is driven by a trade-off with the ability to compete, 325 326 mutualistically efficient Frankia strains can sympatrically coexist with inefficient Frankia strains that have an advantage in intrageneric competition (Ferriere et al. 2002; 327 Hoeksema and Kummel 2003). In actinorhizal symbiosis, Alnus trees often interact with 328 329 different phenotypes of Frankia bacteria, including spore-positive strains hosting 330 abundant sporangia inside plant cells, and sporangia-free, spore-negative strains (Pozzi et al., 2015; Torrey, 1987). Infectivity and nitrogen-fixing activity might be negatively 331 associated between the spore-positive/negative strains (Markham, 2008; Pozzi et al., 332 2015). 333

Third, different functions of Frankia genotypes may contribute to the maintenance of 334 their genetic variation within the same location. Nitrogen resources from rhizobial 335 336 symbionts increases not only the growth of the host plants, but also their resistance to herbivores (Ballhorn et al. 2017; Dean et al. 2014; Thamer et al. 2011). In addition, 337 338 nitrogen fixed by associated rhizobacteria, including *Frankia*, can be stored in nodules and transported to aerial parts as these specific forms, such as amides and ureides (Berry 339 et al. 2011). If the genetic variation of Frankia strains is responsible not only for the 340 nitrogen supply but also the different forms of nitrogen, multiple functions of genetically 341 diverse *Frankia* may complementally improve the overall host plant fitness in a complex 342 343 ecosystem.

Thus, these three interpretations which are not mutually exclusive could complimentary contribute to explain the mosaic-like, spatial genetic structure of *Frankia*. In future studies, phenotypes and functions of different OTUs detected in this study should be investigated.

348

349 Spatial structure in local genetic communities of *Frankia*

The results also revealed a complex, mosaic-like, genetic structures of Frankia on a 350 351 single-host-population scale (spatial differentiation of *Frankia* OTU components; Fig 3) 352 that did not depend on geographic distance. The explanations mentioned in the above 353 section can also contribute to the understanding of the spatial mosaic-like patterns observed in Frankia genetic communities. The heterogeneous spatial structure of the 354 355 interactions between genetically diverse hosts and rhizo-microorganisms can exert selective pressure resulting in the spatial differentiation of Frankia communities. In fact, 356 we detected a genetic differentiation of the alder host not only among the studied riparian 357

areas but also within each area (Kagiya et al. 2018), as well as the spatial genetic structure 358 of Frankia. However, a part of patterns in genetic structure of Frankia were inconsistent 359 360 with the pattern of host genetic structure. For example, Frankia compositions were different between BT and SE area (Fig. 5), while A. hirsuta populations were closely-361 362 related. BT area is completely covered with natural forest stands but SE area is close to agriculture field and its riverside landscape is artificially modified. The differences in 363 abiotic and biotic environments could affects the selection outcomes (e.g., $G \times G \times E$: 364 genotype-by-genotype-by-environment interactions), which may contribute the 365 observed mosaic-like structures of the Frankia communities. 366

367 Furthermore, the dispersion processes of *Frankia* may play a key role in generating 368 these spatial patterns. The significant genetic differentiation of Frankia among riparian areas (Table 1) may be due to the dispersal of *Frankia* bacteria by the waterways (Arveby 369 and Huss-Danell 1988; Huss-Danell et al. 1997). In addition, massive snow-melt in the 370 371 study site (snow depth: > 200 cm) may also drive soil bacteria dispersion by transporting soil components along the complex river landscape. Previous studies suggested that 372 herbivorous mammals (Chaia et al. 2012), birds (Paschke and Dawson 1993), and 373 374 invertebrates such as earthworms (Reddell and Spain 1991) can also drive the dispersion of Frankia, carrying their propagules. Frankia propagules did not lose their activity to 375 infect their hosts despite going through the digestive tracts of such animals (Burleigh and 376 Dawson 1995; Chaia et al. 2012). Thus, the genetic mosaic-like structure can be, at least 377 partially, the result of both biotic (e.g., deer, birds, and earthworms) and abiotic dispersion 378 379 processes (snowmelt and waterways).

To our knowledge, ours is the first study that demonstrated spatial heterogeneity in genetic diversity and composition of *Frankia* bacteria in a single-host–population scale.

Our findings suggest that actinorhizal host individuals can interact with different Frankia 382 strains within a population. The interactions with different genotypes of mutualistic 383 384 bacteria widely influences phenotypes of host plants (Ballhorn et al. 2017; Barrett et al. 2012; Dean et al. 2014; Miller and Sirois 1982; Pahua et al. 2018; Prat 1989; Robinson 385 et al. 2000). The variation in mutualistic partnerships on small spatial scales could 386 increase the heterogeneity of ecosystem processes and/or associated community 387 dynamics in forest ecosystems. Understanding genetic structure of nitrogen-fixing 388 bacterial symbionts holds the key to elucidating these dynamics in forest ecosystems. 389 Further research is required to shed more light on the mechanisms that create the spatial 390 391 heterogeneity in genetic diversity and composition of actinorhizal symbionts on a local 392 scale.

393

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400

401 Author contribution

402 SK and SU designed and conducted the investigation, performed the molecular403 analysis, analyzed the data, and wrote the manuscript.

404

405 **Data availability statement**

- 406 The sequence data are deposited at DDBJ with accession numbers of LC482655-
- 407 LC482672.

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Tables

Table 1. Explanation of genetic structure by study areas/sites based on AMOVA.

| Source | df | Estimate | Ρ |
|--------------|-----|----------|---------|
| Among areas | 4 | 1.463 | 0.015 |
| Among sites | 12 | 2.521 | 0.012 |
| Within sites | 184 | 47.552 | < 0.001 |

659 Figure legends

Fig 1. Map of the sampling points in the Uryu Experimental Forest of Hokkaido University, northern Hokkaido, Japan, where *A. hirsuta* seedlings root nodules were collected. Different colors indicate different areas: (a) overall map, (b–f) individual areas (b: UT; c: DRW; d: BT; e: SE; f: DRE). Marker shapes in magnified map areas (b-f) indicate sites within each area.

665

Fig 2. Phylogenetic tree based on the nifD-K spacer region in Frankia. A maximum-666 likelihood phylogeny was generated based on 97 sequences of the nifD-K locus, obtained 667 from three subspecies of Alnus incana (ssp. hirsuta: operational taxonomic units 'OTUs' 668 in this study; ssp. rubra: 'AR_'; ssp. tenuifolia: 'AT_'), A. viridis ('AV_'), nine varieties 669 of Myrica rubra (var. biji: 'Mbj_'; var. baimei: 'Mbm_'; var. dongkui: 'Mdk_'; var. muye: 670 'Mmy_'; var. *shuimei*: 'Msm_'; var. *wandao*: 'Mwd_'; var. *wumei*: 'Mwy_'; var. *yangliu*: 671 'Myl '; var. zaoda: 'Mdz '), and the ACN14a strain as an Alnus infection clade. Other 672 673 sequences obtained from Hippöphae salicifolia ('Hsli'), three Coriaria species (C. myrtifolia: 'Cm '; C. japonica: 'Cj '; C. arborea: 'Ca '), Elaeagnus ('EAN1pec'), 674 Datisca glomerate, and Casuarina equisetifolia ('CcI3') were also included in the 675 phylogeny as outgroups. These sequences were obtained from GenBank. The characters 676 677 in parentheses indicate accession numbers on GenBank. Branch labels indicate significant bootstrap values. The tree is drawn to scale, with branch lengths measured according to 678 the number of substitutions per site. 679

680

Fig 3. *Frankia* diversity along sampling size in each site. Different colors indicate different areas

683

Fig 4. Relationships between *Frankia* operational taxonomic units (OTUs) and areas. The phylogeny was generated using the maximum-likelihood method. Branch values indicate significant bootstrap values. Circles indicate the presence of each OTU in each site. Circle sizes represent numbers of OTUs in each site (see also Table 1).

688

Fig 5. Non-metric multidimensional scaling (NMDS) of *Frankia* OTU compositions at
 the site-levels. Colors of points indicate areas. The numbers in points indicate ID of sites.





0.10



Sample Size





ΒT DRE DRW



Article title: High genetic diversity of bacterial symbionts in a single host species population: an alder–Frankia system in the field

Journal name: Plant and Soil

Author names: Shinnosuke Kagiya¹, Shunsuke Utsumi²

Affiliation:

¹Graduate School of Environmental Science, Hokkaido University, North 10, West 5, Sapporo, Hokkaido 060-0810, Japan ²Field Science Center of Northern Biosphere, Hokkaido University, North 9, West 9, Sapporo, Hokkaido 060-0809, Japan

E-mail: Shinnosuke Kagiya (e-mail: slotapir-1035@fsc.hokudai.ac.jp)

| Area | | В | Т | | DR | Ε | | DR | W | | | S | E | | | UT | | |
|----------|----|----|----|----|----|---|----|----|----|----|----|----|---|---|----|----|----|------------------|
| Site No. | 1 | 2 | 3 | 4 | 1 | 2 | 1 | 2 | 3 | 4 | 1 | 2 | 3 | 4 | 1 | 2 | 3 | Accession No. |
| OTU01 | 8 | 3 | 9 | 9 | 6 | 2 | 4 | 11 | 5 | 9 | 11 | 9 | 7 | 6 | 8 | 8 | 9 | Ahi01 (LC482655) |
| OTU02 | 4 | 1 | 0 | 0 | 3 | 2 | 2 | 2 | 2 | 2 | 0 | 0 | 0 | 1 | 4 | 0 | 0 | Ahi02 (LC482656) |
| OTU03 | 0 | 2 | 1 | 1 | 1 | 0 | 2 | 1 | 0 | 2 | 0 | 0 | 1 | 0 | 3 | 2 | 0 | Ahi03 (LC482657) |
| OTU04 | 0 | 1 | 1 | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | Ahi04 (LC482658) |
| OTU05 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 2 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | Ahi05 (LC482659) |
| OTU06 | 0 | 6 | 0 | 0 | 1 | 0 | 1 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | Ahi06 (LC482660) |
| OTU07 | 0 | 0 | 0 | 1 | 2 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | Ahi07 (LC482661) |
| OTU08 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 1 | 2 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | Ahi08 (LC482662) |
| OTU09 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | Ahi09 (LC482663) |
| OTU10 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | Ahi10 (LC482664) |
| OTU11 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | Ahi11 (LC482665) |
| OTU12 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | Ahi12 (LC482666) |
| OTU13 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | Ahi13 (LC482667) |
| OTU14 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | Ahi14 (LC482668) |
| OTU15 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | Ahi15 (LC482669) |
| OTU16 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | Ahi16 (LC482670) |
| OTU17 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | Ahi17 (LC482671) |
| OTU18 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | Ahi18 (LC482672) |
| Total | 13 | 14 | 11 | 11 | 16 | 4 | 10 | 17 | 15 | 15 | 12 | 10 | 9 | 9 | 15 | 10 | 10 | |

Table S1. Numbers of OTUs obtained from each site.

| | | I | | | |
|--------------|-----------|---------|----------|----------|-------------|
| | Region | Shannon | Evenness | Richness | Sample size |
| | BT1 | 0.8587 | 0.7817 | 3 | 13 |
| | BT2 | 1.5367 | 0.8577 | 6 | 14 |
| | BT3 | 0.6002 | 0.5463 | 3 | 11 |
| | BT4 | 0.6002 | 0.5463 | 3 | 11 |
| | DRE1 | 1.8080 | 0.8695 | 8 | 16 |
| | DRE2 | 0.6931 | 1.0000 | 2 | 4 |
| | DRW1 | 1.4708 | 0.9139 | 5 | 10 |
| vel | DRW2 | 1.3157 | 0.7343 | 6 | 17 |
| e-le | DRW3 | 1.6792 | 0.8629 | 7 | 15 |
| Site | DRW4 | 1.0776 | 0.7773 | 4 | 15 |
| | SE1 | 0.2868 | 0.4138 | 2 | 12 |
| | SE2 | 0.3251 | 0.4690 | 2 | 10 |
| | SE3 | 0.6837 | 0.6224 | 3 | 9 |
| | SE4 | 0.3488 | 0.5033 | 2 | 9 |
| | UT1 | 1.0096 | 0.9190 | 3 | 15 |
| | UT2 | 0.5004 | 0.7219 | 2 | 10 |
| | UT3 | 0.3251 | 0.4690 | 2 | 10 |
| Area-level | BT | 1.3153 | 0.6759 | 7 | 49 |
| | DRE 1.692 | | 0.8138 | 8 | 20 |
| | DRW | 1.6392 | 0.6836 | 11 | 57 |
| | SE | 0.5779 | 0.3226 | 6 | 40 |
| | UT | 0.8678 | 0.6260 | 4 | 35 |
| Whole forest | | 1.4373 | 0.4973 | 18 | 201 |
| | | | | | |

Table S2. Shannon diversity indices, evenness, richness and sample size of *Frankia* OTU compositions in each site and area and whole of the Uryu Experimental Forest.

Fig S1. Phylogenetic tree based on the *nif*D-K spacer region in *Frankia*. The neighborjoining phylogeny was generated based on 97 sequences of the *nif*D-K locus, obtained from three subspecies of *Alnus incana* (ssp. *hirsuta*: operational taxonomic units 'OTUs_' in this study; ssp. *rubra*: 'AR_'; ssp. *tenuifolia*: 'AT_'), *A. viridis* ('AV_'), nine varieties of *Myrica rubra* (var. *biji*: 'Mbj_'; var. *baimei*: 'Mbm_'; var. *dongkui*: 'Mdk_'; var. *muye*: 'Mmy_'; var. *shuimei*: 'Msm_'; var. *wandao*: 'Mwd_'; var. *wumei*: 'Mwy_'; var. *yangliu*: 'Myl_'; var. *zaoda*: 'Mdz_'), and the ACN14a strain as an *Alnus* infection clade. Other sequences obtained from *Hippöphae salicifolia* ('Hsli_'), three *Coriaria* species (*C. myrtifolia*: 'Cm_'; *C. japonica*: 'Cj_'; *C. arborea*: 'Ca_'), *Elaeagnus angustifolia* (EAN1pec), *Datisca glomerate* and *Casuarina equisetifolia* ('CcI3') were also analyzed in the phylogeny as outgroups. These sequences were obtained from GenBank. The characters in parentheses indicated accession numbers on GenBank. Branch labels indicate significant bootstrap values. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site.



Fig S2. Phylogenetic tree based on all the *nif*D-K sequences of *Frankia*. A maximumlikelihood phylogeny was generated based on 280 sequences of the *nif*D-K locus, obtained from three subspecies of *Alnus incana* (ssp. *hirsuta*: operational taxonomic units 'KS_' in this study; ssp. *rubra*: 'AR_'; ssp. *tenuifolia*: 'AT_'), *A. viridis* ('AV_'), nine varieties of *Myrica rubra* (var. *biji*: 'Mbj_'; var. *baimei*: 'Mbm_'; var. *dongkui*: 'Mdk_'; var. *muye*: 'Mmy_'; var. *shuimei*: 'Msm_'; var. *wandao*: 'Mwd_'; var. *wumei*: 'Mwy_'; var. *yangliu*: 'Myl_'; var. *zaoda*: 'Mdz_'), and the ACN14a strain as an *Alnus* infection clade. Other sequences obtained from *Hippöphae salicifolia* ('Hsli_'), three *Coriaria* species (*C. myrtifolia*: 'Cm_'; *C. japonica*: 'Cj_'; *C. arborea*: 'Ca_'), *Elaeagnus* ('EAN1pec'), *Datisca glomerate*, and *Casuarina equisetifolia* ('CcI3') were also included in the phylogeny as outgroups. These sequences were obtained from GenBank. The characters in parentheses indicate accession numbers on GenBank. Branch labels indicate significant bootstrap values. The tree is drawn to scale, with branch lengths measured according to the number of substitutions per site. We described OTU groups of each sequence.

