

Asymmetric outcome of community coalescence of seed and soil microbiota during early seedling growth

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1 Asymmetric outcome of community coalescence of seed and soil microbiota during early

2 seedling growth

- 3 Running title: Asymmetric coalescence of seed and soil microbiota
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10 Abstract

11 Seed microbial community constitutes a primary inoculum for plant microbiota assembly. 12 Still, the persistence of seed microbiota when seeds encounter soil during plant emergence 13 and early growth is barely documented. Here, we characterized the interchange event or ¹⁴coalescence of seed and soil microbiota and how it structured seedling bacterial and fungal 15 communities. We performed eight contrasted coalescence events to identify drivers 16 influencing seedling microbiota assembly: four seed lots of two Brassica napus genotypes 17 were sown in two soils of contrasted diversity. We found that seedling root and stem ¹⁸microbiota were influenced by soil diversity but not by initial seed microbiota composition. A 19 strong selection on the two-source communities occurred during microbiota assembly, with 20 only 8-32% of soil taxa and 0.8-1.4% of seed-borne taxa colonizing seedlings. The 21 recruitment of seedling microbiota came mainly from soil (35-72% of diversity) and not from 22 seeds (0.3-15%). The outcome of seed and soil microbiota coalescence is therefore strongly 23 asymmetrical with a dominance of soil taxa. Interestingly, seedling microbiota was primarily

- 24 composed of initially rare taxa (from seed, soil or unknown origin) and sub-dominant soil
- 25 taxa. Our results suggest that plant microbiome engineering success based on native seed or
- 26 soil microbiota will rely on rare and sub-dominant taxa in source communities.
- 27 Keywords: microbiota, coalescence, seed, root, stem, soil, Brassica napus

28 Introduction

29 Plants live in complex associations with a wide variety of microorganisms that can modulate 30 their fitness $(1-3)$. The plant microbiota is mainly acquired horizontally through different 31 environmental sources including soil (4), air (5), rainfall (6) and insects (7). However, some ³²members of the plant microbiota are also acquired vertically through vegetative propagation 33 (8) or sexual reproduction via seeds (9).

³⁴In spermatophyte, seed-associated microbial community constitutes the primary ³⁵inoculum for the next plant generation (10). The seed microbiota can have a crucial role for 36 crop installation by modulating dormancy (11,12), germination (13,14), seedling 37 development (15–17) and recruitment of plant symbionts (18). Seed transmission of some ³⁸specific plant-beneficial or phytopathogenic microbial strains is well documented (19–23). ³⁹However, little knowledge is available on the fraction of the plant microbiota that is acquired 40 through seeds. Pioneer studies indicate that a fraction of seed microbiota persists during 41 germination and emergence and actively colonizes seedlings (24–26). Still, most of these 42 studies have been carried out in gnotobiotic conditions in absence of other environmental 43 sources, like soil.

44 The encounter between two microbial communities and how they then evolve 45 together has been called community coalescence (27-29). The outcomes of community ⁴⁶coalescence can vary in a continuum between two categories from symmetric to asymmetric 47 outcomes (30). In symmetrical outcomes, the initial communities contribute quite equally to 48 the resultant coalesced community, whereas in asymmetrical outcomes one initial 49 community becomes dominant if considering the prevalence of the members. In the latter 50 case, the resident community is generally favored at the expense of the invasive community.

51 This resident advantage can be explained by (i) a higher population size of the resident 52 community (i.e. mass effects) (31), (ii) prior use of resource and space (i.e. priority effects) 53 (32) or (iii) local adaptation of the resident community to the reception niche (i.e. 54 community monopolization effect) (33).

55 To date, seed or soil microbiota have been extensively studied in isolation but their 56 coalescence, the assembly mechanisms at play remain broadly unexplored and their 57 coalescence type remains to be characterized. Seed and soil coalescence present a singular 58 situation if the seedling community is considered as the outcome community. The seed 59 community is extremely reduced in terms of abundance compared to soil community, but its ⁶⁰members have potentially a 'home advantage' as they have been selected by the plant. We 61 therefore hypothesized that despite its lower microbial richness and diversity, seed ⁶²microbiota being already present and more adapted to the plant environment, would 63 benefit from priority effects over soil microbiota for the colonization of plant compartments 64 (18). Nevertheless, adaptation to seed does probably not recover all features for adaptation 65 to a growing seedling. The initial soil microbial diversity level is probably another key driver 66 of seedling microbiota and persistence of seed-borne microbial taxa. Indeed, we propose 67 that a soil of high microbial diversity would induce a weaker colonization of the seed ⁶⁸microbiota on seedlings due to stronger competition and higher functional redundancy in 69 this soil (34).

70 10 In the present work, we monitored the outcome of seed and soil communities' 71 coalescence during early seedling growth. More specifically, we investigated (i) what is the 72 main source of microbial transmission to seedlings; (ii) does seed microbiota composition 73 and soil diversity impact the assembly of seedling microbiota; and (iii) does the initial taxon

74 abundance in the source influence its transmission success? To answer these questions, we 75 selected seed lots of different genotypes of the winter oilseed rape Brassica napus, which 76 harbored contrasting seed microbiota (35). We simulated a total of eight different 77 coalescence events by sowing the four seed lots in two soils of different microbial diversities. 78 Bacterial and fungal community structure was characterized by amplicon sequencing of the 79 source seed and soil communities and of the roots and stems sampled at different seedling 80 growing stages. This study gained new insights into the soil and plant drivers that influence 81 the coalescence of soil and seed microbiota during the first steps of seedling microbiota 82 assembly.

84 Materials and Methods

85 Soil preparation

86 In 2014, soil was sampled at a depth of 10-30 cm in a 20-year wheat experimental plot (La 87 Gruche, Pacé, France, 48°08'24.5"N 1°48'01.0"W). Soil was sterilized and recolonized with 88 two levels of soil microbial diversity according to the experimental procedure described in 89 Lachaise et al. (36). In short, a fraction of the soil sample was ground and sieved at 2 mm. ⁹⁰Three sub-samples (900 g each) were soaked in 8L of sterile water. The resulting soil 91 suspensions were 1:10 serially diluted in sterile water up to the 10⁶. Another fraction (~80 92 kg) of the soil sample was ground, sieved at 4 mm and mixed with 1/3 washed sand. This soil 93 was dispatched in 2.5 kg bags and gamma-irradiated (35 kGy, Ionisos, France). Each bag was 94 inoculated with 320 mL of undiluted or $10⁶$ or soil suspensions, therefore resulting in soil 95 with high and low microbial diversity, respectively. The soil recolonization was repeated 3 96 times (A, B, C). The bags were mixed and aerated in sterile conditions twice a week during 97 the 39 incubation days at 20° C, to homogenize gaseous exchanges and recolonization. 98 Microbial recolonization dynamics of soil was monitored by sampling 30g of soil at several 99 times. For bacteria, each sample was serial diluted and plated on 1/10 strength Tryptic Soy 100 Agar (17 g.L⁻¹ tryptone, 3 g.L⁻¹ soybean peptone, 2.5 g.L⁻¹ glucose, 5 g.L⁻¹ NaCl, 5 g.L⁻¹ K₂HPO₄ 101 and 15 g.L¹ agar) supplemented with Nystatin (0.025 g.L¹), and the number of colony-102 forming units was measured after 3 days of incubation at 27°C. For fungi, following the pour-103 blate method, serial diluted samples were mixed with molten Acid Malt Agar (10 g.L⁻¹ malt 104 extract, 15 g.L⁻¹ agar and 0.25 g.L⁻¹ citric acid) supplemented with streptomycin (0.15 g.L⁻¹) 105 and penicillin (0.075 g.L¹) and the number of colony-forming units was measured after 7 106 days of incubation at 20°C. After 39 days of incubation, a portion of each soil (high and low

111 Sowing and sampling

112 After 39 days of soil incubation, individual pots were filled with a 5 mm layer of sterile 113 vermiculite and 80g of soil of high or low microbial diversity. Soils were saturated with tap 114 water by sub-imbibition one day before sowing, in order to reach approximately 80% of 115 humidity (retention capacity) on the day of sowing. During the experiment, plants were 116 watered twice (at 5 and 12 days) with tap water.

117 Seed samples from two genotypes of Brassica napus (Boston and Major) collected 118 during two consecutive years (Y1 and Y2) were selected for this work, resulting in four 119 contrasted initial seed microbiota used for the coalescence experiment. At d0, seeds of each 120 genotype and year were individually sown at a depth of 5mm and grown under controlled 121 conditions (14h day/ 10h night period, 20°C). Seven days (d07) and fourteen days (d14) after 122 sowing, 10 and 20 plants were sampled per modality (2 genotypes x 2 harvesting years x 2 123 soils), respectively. Roots were cut from stems and gently soaked 10 seconds in sterile water 124 to remove residual soil. Otherwise, stems were size-equalized on 2cm from the root basis. 125 Therefore, the resulting sampled root habitat was composed of the inner root and 126 rhizoplane and the stem habitat was composed of inner stem and stem surface. The 10 (d07) 127 or 20 (d14) roots and stems were pooled separately and stored as root and stem 128 compartments at -80°C until DNA extraction.

129 Microbial DNA sample preparation

130 Seed samples were prepared for extraction as previously detailed in Rochefort et al., (2019). 131 Briefly, seeds were soaked in phosphate-buffered saline (PBS, Sigma-Aldrich) supplemented 132 with Tween[®] 20 (0.05 % v/v, Sigma-Aldrich) during 2h30 at 4°C. DNA extraction was 133 performed with the DNeasy PowerSoil HTP 96 Kit (Qiagen) following manufacturers 134 procedure. Soil, root and stem samples were lyophilized before DNA extraction. Samples 135 were mixed with 1mm and 3mm beads, and crushed 2 x 30s at 5m/s with FastPrep (MP 136 Biomedicals). DNA extraction was performed with the DNeasy PowerSoil HTP 96 Kit.

137 Library construction and sequencing

138 Amplicon libraries were constructed with the primer sets gyrB aF64/gyrB aR553 (24) and 139 ITS1F/ITS2 (37), following the experimental procedure previously described in Rochefort et 140 al. (35). A blank extraction kit control and a PCR-negative control were included in each PCR 141 plate. Amplicon libraries were mixed with 10% PhiX and sequenced with two MiSeq reagent 142 kit v3 600 cycles.

143 Sequence processing

144 Primer sequences were removed with cutadapt version 1.8 (38). Fastq files were processed 145 with DADA2 version 1.6.0 (39), using the following parameters: truncLen=c(190, 200), 146 maxN=0, maxEE=c(1,1), truncQ=5 for gyrB reads. ITS1 reads were processed with the same 147 parameters, except that reads were not trimmed to a fixed length and that trunQ was set to ¹⁴⁸2. Chimeric sequences were identified and removed with the removeBimeraDenovo function 149 of DADA2. Taxonomic affiliations of amplicon sequence variants (ASV) were performed with 150 a naive Bayesian classifier (40) implemented in DADA2. ASVs derived from gyrB reads were 151 classified with an in-house gyrB database (train set gyrB v4.fa.gz) available upon request. 152 ASVs derived from ITS1 reads were classified with the UNITE v7.1 fungal database (41).

153 Microbial community analyses

¹⁵⁴Analyses of diversity were conducted with the R package Phyloseq version 1.26.0 (42). Since 155 the primer set gyrB aF64/gyrB aR553 can sometimes co-amplify parE, a paralog of gyrB, the 156 gyrB taxonomic training data also contained parE sequences. Hence, ASV affiliated to parE or 157 unclassified at the phylum-level were removed. Sequences were aligned with DECIPHER ¹⁵⁸2.14.0 (43) and neighbor-joining phylogenetic trees were constructed with Phangorn 2.5.5 ¹⁵⁹(44). Identification of sequence contaminant was assessed with decontam version 1.4 (45) 160 using the "prevalence" method at a threshold of 0.1. Fungal ASVs unassigned at the phylum 161 level were removed. For the ITS1 dataset, we created a hybrid-gene phylogenetic tree (i.e. 162 ghost tree) (46) based the 18S rRNA gene sequence database (Silva 132) and ITS database 163 (UNITE v8) in order to perform phylogenetic analyses.

164 Data were normalized based on sequencing depth. For soil analyses, data were 165 rarefied at 30 000 and 25 000 reads per sample for gyrB and ITS1, respectively. For plant 166 (root and stem) analyses, data were rarefied at 1 000 reads per sample for gyrB and ITS1 ¹⁶⁷(Fig. S1), a validated choice to keep an optimal number of reliable samples. Faith's 168 phylogenetic diversity was calculated on gyrB and ITS1 datasets with the R package picante 169 version 1.8 (47). Differences in alpha-diversity estimators were assessed with Analysis of 170 Variance. Differences were considered as significant at a p-value < 0.05.

171 Changes in microbial community composition were assessed on $log_{10}+1$ transformed 172 values with Bray-Curtis (BC) index. Principal coordinate analysis (PCoA) was used for 173 ordination of BC index. Homogeneity of group dispersions was assessed with betadisper 174 function of vegan 2.5.3 (48). Difference in dispersion between groups was assessed with a 175 Wilcoxon non-parametric test. To quantify the relative contribution of seed microbiota, soil

176 diversity, development stage and habitat in microbial community profiles, permutational 177 multivariate analysis of variance (PERMANOVA) (49) was performed with the function 178 adonis2 of the R package vegan 2.5.3 (48). The relative abundance of ASVs belonging to 179 ammonia-oxidizing bacteria (AOB) and nitric-oxidizing bacteria (NOB) according to their 180 taxonomic classification were investigated in soils of high and low microbial diversities. 181 These include Nitrosospira sp. and Nitrosovibrio sp. for AOB and Nitrobacter sp. and 182 Nitrospira sp. for NOB (50,51). Differences in relative abundance of AOB and NOB were 183 considered as significant with T-test at a p-value < 0.1 .

184 Transmission analyses

185 A prevalence matrix of common ASVs between seeds, soils, roots and stems was constructed 186 on non-rarefied data to have the most exhaustive presence-absence analysis of all taxa. 187 Owing to the microbial variability between individual seeds, an ASV was recorded as present 188 if detected in at least one of the 3 seed sample repetitions. Visualization of bacterial and 189 fungal ASVs shared between biological habitats or specific to one was assessed with the R 190 package UpSetR 1.4.0 (52).

191 The datasets supporting the conclusions of this article are available in the European 192 Nucleotide Archive under the accession number PRJEB41004.

194 Results

195 Selected seed lots have distinct microbial communities

196 Harvesting year and plant genotypes are significant drivers of the diversity structure of the B . 197 napus seed microbiota (35). Two B. napus genotypes (Boston and Major) collected during 198 two harvest years (Y1 and Y2) were selected for this work. The most prevalent seed-borne 199 bacterial ASVs were affiliated to Sphingomonas and Frigoribacterium genera (Fig. S2A). 200 Fungal communities were dominated by ASVs affiliated to *Cladosporium* and Alternaria 201 genera (Fig. S2B).

²⁰²Estimated bacterial richness (Chao1) was on average at least two times higher in Y2 ²⁰³(250 ASVs) than in Y1 (100 ASVs) for both genotypes, and phylogenetic diversity (Faith's PD) 204 was also significantly ($p < 0.05$) higher in Y2 (Fig. 1A).

²⁰⁵While estimated fungal richness was also higher in Y2 (~50 ASVs) than in Y1 (~35 206 ASVs), phylogenetic diversity was higher in Y1 (Fig. 1B). According to distance from centroid, 207 variation in bacterial and fungal community composition (BC index) was significantly lower (p 208 \lt 0.05) in Y2 in comparison to Y1 (Fig. 1C). Structure of microbial communities was 209 significantly (p <.001) impacted by harvesting year with 24% and 62% of variance explained 210 by this factor for bacteria and fungi, respectively (Fig. 1D). Moreover, seed genotype was 211 also significantly ($p < 0.001$, 13% of variance) impacting bacterial community composition 212 but not fungal community composition (Fig. 1D). In brief, the structure of microbial 213 communities was different between the seed lots selected for this study and harvest year 214 was the most important driver of these changes.

216 Production of soils with contrasting levels of diversity

217 To obtain two soils with contrasted levels of microbial diversity, gamma-irradiated soil was 218 inoculated with undiluted and diluted soil suspensions (Materials & Methods). After 39 days 219 $\,$ of incubation, a plateau of 10 9 bacterial CFU and 10 5 fungal CFU per gram of soil was reached 220 for each soil (Fig. S3). The acidity level of low-diversity soils significantly ($p = 0.003$) increased 221 by 0.65 pH unit compared to high-diversity soil. In addition the amount of nitric nitrogen was 222 twice as large in high-diversity soils ($p < 0.001$), whereas ammoniacal nitrogen significantly (p 223 \leq 0.001) decreased (Table 1). Relative abundance of AOB was significantly (p < 0.1) higher in 224 the soil of high diversity in comparison with the soil of low diversity. Moreover, no NOB was 225 detected in the soil of low diversity while the soil of high diversity was composed on average 226 of 0.25% NOB.

227 The most prevalent soil-borne bacterial ASVs were affiliated to Massilia, Nitrosospira 228 and Sphingomonas (Fig. S2A). The most prevalent soil-borne fungal ASVs were affiliated to 229 Mortierella, Trichoderma and Exophiala (Fig. S2B).

230 According to alpha-diversity indexes (Chao1 and Faith's index), high-diversity soils 231 presented a higher (p <0.05) bacterial richness (~800 ASVs) and phylogenetic diversity 232 compared to low-diversity soils (\sim 500 ASVs) (Fig. 1A). With regard to fungal communities, 233 only phylogenetic diversity was greater in high-diversity soils (Fig. 1B). Variation in microbial 234 community composition was more important for low-diversity soils compared to high-235 diversity soils (Fig. 1C). The initial level of dilution employed explained 40% and 39% of the 236 variance observed in bacterial and fungal communities, respectively (Fig. 1D). Hence, the 237 dilution approach employed in this work resulted in two soils with contrasted levels of 238 phylogenetic diversity and distinct community structure.

239 Initial level of soil diversity impacted the diversity and structure of the plant microbiota

240 The outcome of community coalescence between seed and soil microbiota was investigated 241 on two plant compartments (stem and root) collected at two distinct plant developmental 242 stages (d07 and d14). On the seedling roots and stems, the most prevalent bacterial and 243 fungal ASVs presented abundance patterns clearly influenced by initial soil diversity (e.g. 244 Devosia or Fusicolla aquaeductuum, Fig. S2). Diverse bacterial (e.g. Afipia, Ensifer, 245 Fictibacillus, Nocardiodes) and fungal genera (e.g. Peziza, Fusicolla, Metarhizium) were 246 dominant on seedling roots. The stems were dominated by bacterial ASVs affiliated to 247 Bacillus, Nocardioides, Devosia and Fictibacillus, while the most prevalent fungal genera 248 were Fusarium, Fusicolla and Acremonium (Fig. S2).

249 Seed genotypes and harvest years did neither impact estimated richness nor 250 phylogenetic diversity in stem and root (Fig. 2).

251 In contrast, the initial level of soil diversity significantly ($p < 0.05$) influenced microbial 252 richness (Fig. 2A and 2C) and phylogenetic diversity (Fig. 2B and 2D) in stems at d07 and d14, 253 with a higher richness and diversity in stems collected from high-diversity soils. No 254 significant change in bacterial richness and phylogenetic diversity was detected in roots 255 between the two soils (Fig. 2A and 2B) while fungal richness and phylogenetic diversity were 256 almost two times higher in roots from soil of high diversity at each development stage (Fig. 257 2C and 2D).

258 Moreover, there was a little but significant increase in bacterial richness and 259 phylogenetic diversity in stems at stage d14 compared to stage d07. No differences were 260 observed on fungal diversity across stages (Fig. 2).

261 Composition of the plant microbiota was not influenced by the seed genotype and 262 the harvesting year (Fig. 2E-H and Table 2). In contrast, the initial soil diversity significantly 263 ($p < 0.001$) contributed to the variance observed in stem (14.6%) and root (36.8%) bacterial 264 communities (Fig. 2I-J) as well as the variance in stem $(24.1%)$ and root $(22.1%)$ fungal 265 communities (Fig. 2K-L and Table 2). Stem- and root-associated microbial community 266 compositions were also significantly impacted by the plant developmental stage (Table 2).

267 Relative contribution of seed-borne and soil taxa to seedling microbiota

268 To describe the outcome of community coalescence between seed and soil microbiota 269 during seedling growth, we characterized the proportion of bacterial and fungal ASVs from 270 seed, soil or unknown origin that composed the stem and root seedling microbiota. A total 271 of 39 bacterial ASVs and 16 fungal ASVs were shared between seeds, roots and stems (Fig. 272 3). Among these ASVs, 16 bacterial and 4 fungal ASVs were also found in soil samples.

²⁷³Between 35 and 72% of seedling-associated bacterial ASVs were detected in the soil, 274 while 20 to 45% of seedling fungal ASVs came from the soil (Fig. 3). These soil-derived ASVs 275 represented between 40 and 98% of the total seedling microbiota. On the contrary, few 276 seedling-associated ASVs were detected in the seed (0.3-15%) representing less than 1% of 277 microbial relative abundance in roots and stems (Fig. 3). Lastly, a fraction of the seedling 278 microbiota was of undetermined origin as undetected in the initial seed and soil pools. ASVs 279 of undetermined origin either may correspond to unsampled taxa in soil or seed 280 compartments (i.e. rare taxa) or may derive from other environmental sources. The 281 proportion of bacterial ASVs of undetermined origin was higher for plants grown in the high-282 diversity soil (approximately 45-65% of community membership and composition) than in 283 the low-diversity soil (20-30%) (Fig. 3 and Fig. $S4A$). This result suggests that seedlings are

284 primarily colonized by rare soil taxa that were initially undetected. In contrast, fungal ASVs 285 and detected in soil and seeds represented approximately 40-60% of community 286 membership in plants but their impacts on community $(i.e.$ their relative abundance) was 287 quite low with less than 20% regardless of the soil diversity (Fig. 3 and Fig. S4B).

288 Contrasting microbial taxonomic composition was observed between the initial soil 289 and seed pools. Concerning bacteria, Enterobacterales and Sphingomonadales were more 290 abundant in seeds, while Burkholderiales were more abundant in soils (Fig. 4A). Changes in 291 bacterial order relative abundance were also detected between stem and root, especially at 292 d07 where Bacillales dominated the bacterial fraction of the stem microbiota (Fig. 4A). The 293 taxonomic composition of the fungal fraction of the seed and soil microbiota was highly 294 contrasted at the order level. Seeds were mainly inhabited by Pleosporales, Capnodiales and 295 Tremellales, when soils were principally constituted of Mortierellales and Hypocreales (Fig. 296 4B). This latter order was however more prevalent in low-diversity soils in comparison to 297 high-diversity soils (Fig. 4B). As in soil, Burkholderiales remained the dominant order in roots 298 but were in part outcompeted by an increasing proportion of Rhizobiales. Xanthomonadales 299 were reduced in roots especially in low-diversity soil and more drastically after 14 days. 300 Bacillales were the transitory most dominant order at 7 days in stems before to regress at 14 301 days. On the opposite Pseudomonadales and Rhizobiales were transitory limited in stems 302 especially in low-diversity soil before to increase at 14 days (Fig. 4A). For fungi, Hypocreales 303 were highly prevalent in seedlings when Mortierellales were more or less prevalent at 7 days 304 in roots and stems, then disappeared in roots and were not represented in stems at 14 days 305 (Fig. 4B). These pools represented only a small proportion of transmitted ASVs from seed to 306 seedling compared to the pools originating from soil (Fig. 3). Mainly bacterial ASVs belonging

307 to Bacillales, Burkholderiales and Pseudomonadales and many fungal ASVs belonging to ³⁰⁸Hypocreales were transmitted from soil to seedlings.

309 Seedling transmission success of seed-borne and soil microbiota

310 The contrasted seed lots and soils employed in the experimental design enabled the 311 investigation in the influence of the initial diversity of the colonizing pools on the 312 transmission success to the seedling (Fig. 5).

313 For bacteria and fungi, we found no clear relationships between the level of diversity 314 of the initial pool (seed or soil) and the percentage of ASVs transmission in seedlings (Fig. 5). 315 Soil microbiota always presented the highest rates of ASVs transmission to seedlings with 10 316 to 30% of the initial ASV pool. Overall, the transmission of soil-borne microbiota to seedlings 317 was higher for the bacterial community (113-180 ASVs) than for the fungal community (3-18 318 ASVs, Fig. 5A, 5B). For bacteria, transmission to seedlings was higher in low-diversity soil 319 $(32\% + 0.84)$ compared to high-diversity soil (15.6% + 1.49, Fig. 5A). For fungi, the opposite 320 pattern was observed with a higher percentage of ASVs transmitted to seedlings from high-321 diversity soils (19.8% + 3.24) compared to low-diversity soil (8.5% + 1.41, Fig. 5B). A low 322 percentage of ASVs transmission from seeds to seedlings was observed for bacteria (0.8% + 323 0.93) and fungi (1.45% + 1.44), with no clear influence of the genotype and harvest year (Fig. 324 5). Finally, overall ASV transmission to seedlings was similar on roots and stems, except for 325 the fungal pool of the high soil diversity where transmission rate in roots was higher (22.1%) 326 \pm 3.18) than transmission rate in stems (17.7% \pm 1.2, Fig. 5).

327 Emergence of rare and sub-dominant taxa in seedlings

³²⁸We next examined the influence of the initial ASV relative abundance on its seedling 329 transmission success and resulting abundance in roots and stems at d14. ASVs were grouped

330 in three arbitrary abundance classes: rare (average relative abundance <0.01%), sub-331 dominant (between 0.01 and 1%) and dominant (>1%). An interesting finding was that 332 dominant ASVs in the source pool (seed or soil) almost never became dominant in roots and 333 stems (below 1:1 line) or were not even transmitted to seedlings (Fig. 6). The majority of 334 transmitted seed-borne bacterial ASVs were initially rare taxa, with nine out 11 ASVs in roots 335 and 11 out of 16 ASVs in stems (Fig. 6A, 6C). Only one dominant seed ASV (out of 65 in the 336 initial pool) was found to be transmitted to roots and stems (Afipia). One sub-dominant ASV 337 was successfully transmitted to both roots and stems (Blastomonas) and three sub-dominant 338 ASVs to stems solely (Pseudomonas lurida, Cutibacterium acnes, Hyphomicrobium) (Table 339 S1). It should be noted that none of these dominant and sub-dominant transmitted seed-340 borne bacterial ASVs became dominant in seedlings (relative abundance <0.3%).

341 The most abundant root bacterial ASVs (>1%) originated from soil-borne ASVs 342 classified as sub-dominant ASVs (n=26 ASVs) and dominant ASVs (5 ASVs) (Fig. 6B, 6D). For 343 the stems, most of the abundant ASVs originated from sub-dominant soil ASVs (n=24), a few 344 dominant ASVs (n=6) and one rare ASV. When considering the entire seedling bacterial 345 microbiota and not only the most abundant taxa, we found that 95% of the dominant soil 346 ASVs, 40% of the sub-dominant ASVs and 28% of the rare ASVs were transmitted to roots. 347 For the stems, 86% of dominant ASVs, 40% of sub-dominant ASVs and 29% of rare ASVs 348 were transmitted.

349 For the fungal community, four seed-borne ASVs were transmitted to both roots and 350 stems, composed of two dominant ASVs (Alternaria infectoria, Cladosporium delicatulum), 351 one sub-dominant (Alternaria sp.) and one rare (Gibberella avenacea). In addition to these 352 four ASVs, three additional dominant ASVs, all affiliated to A. *infectoria*, were transmitted to

361 Altogether, these results indicate that seed-borne taxa transmitted to seedlings are 362 predominantly rare taxa, while transmitted soil-borne taxa are primarily sub-dominant and 363 dominant taxa. Moreover, this analysis shows that the transmission rates of fungi is two to 364 four times lower than bacteria for all ASV abundance classes (dominant: 41-53% vs 86-95%; ³⁶⁵sub-dominant: 10-12% vs 40%; rare: 11-12% vs 28-29%).

367 Discussion

368 Asymmetric outcome of seed and soil community coalescence in favor of soil microbiota

369 Soil microbiota had a great influence on the plant microbiota composition since 70% and 370 30% of soil-borne bacterial and fungal taxa were detected in seedlings, respectively. In 371 contrast, transmission of seed-borne microorganisms to roots and stems of B. napus was 372 lower than soil with on average 2% of bacterial ASVs and 12% of fungal ASVs detected in 373 seedlings. The remaining seedling-associated ASVs were not identified in both seed and soil 374 habitats. It is highly plausible that most of these taxa were soil-borne and were initially not 375 sampled due to the high heterogeneity of soil spatial structure (28). Indeed, there were on 376 average many more bacterial ASVs of undetermined origin in the seedlings growing in the 377 soil of high diversity (> 50%) than in those from a soil of low diversity (< 30%). These data 378 suggest that a large part of these ASVs come from the soil but that they were not sampled in 379 high-diversity soils. Overall, these results highlighted an asymmetric outcome of the 380 coalescence between the soil and the seed microbiota (30), where the soil microbiota take 381 precedence over the seed microbiota.

382 This asymmetric coalescence could be due the high population size of soil-borne taxa 383 in comparison to seed-borne microbial population, a process known as mass-effect (31). ³⁸⁴Although seed microbial taxa can be considered as plant residents that are adapted to 385 available niches (53), the successful invasion of root and stem by soil-borne taxa is 386 undoubtedly facilitated by their population sizes. Alternatively, the dominance of soil 387 associated microbial communities in seedlings could be explained by the higher amount of 388 phylogenetic diversity in soil in comparison to seeds. Higher phylogenetic diversity could 389 indeed increase the functional capabilities of soil associated microbial communities and then

390 limit niche overlap between microbial entities (54). The level of diversity of the initial seed 391 and soil pools were however not directly correlated to the number of seedling transmitted 392 ASVs (Fig. 5). In other words, the diversity of the regional species pool (i.e. soils and seeds) 393 were not a good predictor of the diversity of local communities (i.e. seedlings).

³⁹⁴Weak transmission of seed-borne microorganisms to the rhizosphere has been 395 previously reported in Asteraceae (55), Cucurbitaceae (56,57) and Poaceae (25,58). Survival 396 on barley roots of seed-borne bacteria occurred in the non-growing part of the root system 397 but not in emerging roots that are colonized principally by soil-borne bacteria (58). While we 398 did not investigate the fate of seed-borne microorganisms in different parts of the root 399 system of B. napus, the same proportion of seed-borne taxa were detected within stems or 400 roots. Hence, the weak transmission of seed-borne taxa to seedlings was not a consequence 401 of microhabitat heterogeneity.

402 Seedling microbiota assembly is driven by the level of soil diversity but not the structure of 403 the seed microbiota

⁴⁰⁴The initial level of soil diversity was the key driver in the assembly of the stem and root 405 microbiota. Indeed, stem microbial diversity and root fungal diversity were lower in the low-406 diversity soils compared to high-diversity soils (Fig. 2). Moreover, soil diversity explained 407 between 15% to 37% of variance in microbial community composition in stem and root (Fig. 408 2). The impact of soil diversity on plant microbiota composition increased over time in roots 409 and stems therefore confirming soil resident time and plant age in root microbiota dynamics ⁴¹⁰(59,60).

411 The soil recolonization approach employed in this work did result in two soils with 412 contrasted levels of bacterial and fungal diversity. This modification of soil microbial diversity

413 also impacts the functional diversity and ultimately the soil physicochemical parameters. 414 Changes in NH₄⁺ and NO₃ concentrations and pH between low and high-diversity soils were 415 indeed monitored. A similar drastic switch in the net balance between mineral nitrogen 416 forms, from dominant nitrate forms in the high diversity soil to an altered N-cycle with 417 dominant ammonium forms in the low diversity soil is reported in another soil dilution 418 \quad experiment (61). A lower NO₃ amount is consistent with a reduced abundance of bacterial 419 – and archaeal ammonia oxidizers. A higher NO_3 amount is linked with a decrease in the 420 abundance of denitrifiers that can reduce $NO₃$ into N₂. Nitrification process is associated 421 $\;\;\;\;$ with pH reduction prior to NO₂ accumulation period, with a potential decrease of 0.75 pH 422 unit (62). Hence, the impact of soil diversity on seedling microbiota structure could not be 423 solely due to the level of microbial diversity but to local changes of physicochemical 424 parameters.

425 Contrary to what has been observed in the soil, we did not detect any significant 426 impact of the initial seed microbiota composition on the overall structure of the stem and 427 root microbiota. This observation reflected not only a low transmission of seed-borne taxa to 428 roots and stems of B. napus but also a weak, if not an absence, of historical contingency. 429 Historical contingency is mediated by priority effects that correspond to the impact of 430 species on one another depending on its order of arrival within the local community (32). 431 Importance of historical contingency in assembly of the plant microbiota was recently 432 highlighted in wheat, where the identity of seed-transmitted fungal taxa can modify 433 colonization of roots by dark septate endophytes, three weeks following germination (18). 434 While we cannot conclude there is an absence of priority effects between microbial species 435 within the root and stem microbial communities of B. napus, historical contingency is not 436 promoted during seedling community assembly.

437 Seedling microbiota is primarily composed of initially rare and sub-dominant taxa in source

439 The initial abundance of microbial taxa in the regional species pool was not positively 440 correlated to their seedling transmission. This was particularly marked for bacterial ASV 441 where rare seed-borne taxa and sub-dominant soil-borne taxa were mostly transmitted to 442 the seedling (Fig. 6, Fig. S5). An increase in the abundance of rare taxa in seedlings could be 443 explained by several attributes including more favorable environmental conditions for their 444 growth and awakening from dormancy (63). Many seed-borne bacteria enter a viable but 445 non-culturable (VBNC) state (64–66). Nutrient-rich environment, such as the spermosphere 446 (67), can elicit the resuscitation of VBNC cells (68). In any case, the rare taxa that are 447 selected on seedlings must carry specific genetic determinants that are responsible for their 448 higher fitness that deserve to be investigated further. The emergence of rare taxa seems to 449 be a shared coalescence outcome between different ecosystems, *i.e.* for the mixing of 450 freshwater and marine microbiome (29), the unequal mixing of two soils with different 451 physicochemical and microbial compositions (69) or the mixing of soils for the outcome of 452 rhizobial communities on rooibos nodules (70). Rare taxa becoming dominant can provide 453 essential or new functions in nutrient cycling or plant growth, which replace or compensate 454 for the function deficiency of abundant species (71).

455 Interestingly, dominant taxa in the source pool (seed or soil) almost never became 456 dominant in roots and stems. None of the dominant seed taxa (e.g. P. agglomerans) is 457 transmitted maybe because these taxa are well adapted to ripe seed and low osmotic seed 458 environment and not to growing seedling in competition with soil microbiota. Conversely, 459 the better transmission of dominant and sub-dominant soil-borne taxa likely combines soil

⁴⁶⁰microbiota mass effect and a plant selective process among soil bacteria that are more 461 adapted to root and stem habitats.

463 Conclusion

⁴⁶⁴In conclusion, we highlighted that the outcome of seed and soil microbiota 465 coalescence was strongly asymmetrical with a clear dominance of soil microbiota. The level 466 of soil diversity was an important driver of the structure of the seedling microbiota. Our 467 approach enabled us to quantify the relative contribution of seed-borne and soil-borne taxa 468 to seedling microbiota assembly and provided an estimation of transmission rates of each 469 source microbiota. Out of all soil taxa detected, only 8 to 32% were able to colonize 470 seedlings, while the proportion was of 0.8 to 1.4% for seed-borne taxa, which indicates a 471 strong selection during seedling microbiota assembly. We demonstrate a high transmission 472 of rare seed-borne and sub-dominant soil-borne taxa to seedlings, which is also a key feature 473 of this coalescence (Fig. SS). These results provide an important foundation for the 474 development of plant microbiome engineering through the modification of native seed or 475 soil microbiota. Particularly, we encourage future work using natural or synthetic 476 communities to integrate rare microbial taxa, as we showed that focusing only on dominant 477 taxa in the source-communities is not informative to understand future seedling microbiota 478 composition.

479 The community coalescence framework just started to emerge in the field of 480 microbial ecology and we show here that this approach accelerates discoveries related to 481 plant microbiota assembly and microbiome engineering. In this dynamic, further studies are 482 needed to understand the ecological processes involved in this coalescence between seed

483 and soil and how direct inoculation of microorganisms or modification of environmental 484 conditions might alter its outcome.

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694 Figure legend

695 Figure 1 Diversity of initial seed and soil microbial community pools before coalescence. (A) 696 Bacterial alpha-diversity. (B) Fungal alpha-diversity. (C) Microbial dispersion to centroid. (D) 697 Microbial beta-diversity. Bacterial and fungal richness were estimated with Chao1. Bacterial 698 and fungal phylogenetic diversities were calculated with Faith's phylogenetic diversity (PD). 699 PCoA ordination of Bray-Curtis index was calculated for bacteria and fungi. Seed microbiota 700 (blue) was described in two B. napus genotypes (Boston, Major) harvested during two 701 consecutive years (Y1, Y2). Soil microbiota (green) was described in two soils originating 702 from dilution-extinction serials (High-, Low-diversity). 703 Figure 2 Structure of seedling microbiota after coalescence with contrasted seed and soil

⁷⁰⁴microbiota. (A, B) Bacterial alpha-diversity. (C, D) Fungal alpha diversity. (E, F, I, J) Bacterial 705 beta-diversity in stems and roots. (G, H, K, L) Fungal beta-diversity in stems and roots. Open 706 and closed dots are B. napus genotypes. Dots are colored either by harvest year (A-H) or by 707 soil diversity (I-L). Bacterial and fungal richness were estimated with Chao1. Bacterial and 708 fungal phylogenetic diversities were calculated with Faith's phylogenetic diversity. PCoA 709 ordination of Bray-Curtis index was calculated for bacteria and fungi.

710 Figure 3 Origin of microbial taxa within stem and root assemblages. Bacterial (A) and fungal ⁷¹¹(B) ASVs shared between seeds, soil, roots and stems at each harvesting stage (d07 or d14) 712 and soil diversity (High or Low). For each plot, upper-right bar charts summarize the 713 proportion and relative abundance of plant-associated ASVs that were detected in soil 714 and/or seeds or not detected (unknown) within these habitats.

727

Seed microbiota

Soil microbiota

E - Stem bacterial composition

I - Stem bacterial composition

B - Bacterial phylogenetic diversity

- **High** Low **High** Low
- F Root bacterial composition

J - Root bacterial composition

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 $30[°]$

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 $10₁$

 Ω

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 $30[°]$

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 $10 -$

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G - Stem fungal composition

K - Stem fungal composition

- D Fungal phylogenetic diversity $d07$ $d14$ $5 \mathbf b$ \overline{a} $\mathbf b$ \overline{a} $\overline{4}$ $3 -$ Stem Genotype $2¹$ ◯ Boston Θ ● Major Ω $5 \mathsf b$ b a a Year \bullet Y1 3 \bullet Y2 Root $\overline{2}$ Ω High Low High Low H - Root fungal composition Genotype $d07$ $d14$ ◯ Boston $[146%]$ ● Major Year Axis 2 \bullet Y1 \bullet Y2 Axis 1 [34.1%] L - Root fungal composition $d07$ $d14$ $[146%]$
	- Soil diversity \bullet High **O** Low

Axis 2

B

A.

B.

Mean % of ASV transmitted from the initial pool

transmitted

of ASV

Mean %

from

the initial pool

Mean number of ASV in initial pool

Mean number of ASV in initial pool

Table 1: Soil physicochemical analyses and relative abundance of bacterial nitrifiers. Values for high and low soil microbial diversities are the average of the 3 replicates ± standard errors. T-test was performed on data, and significant p-values are indicated with an asterisk. AOB = Ammonia-oxidizing

bacteria. NOB = Nitrite-oxidizing bacteria.

Table 2: Permutational multivariate analysis of variance on seedling roots and stems microbial betadiversity. Linear model was built with the adonis2 function on data separated by root or stem compartment, integrating soil diversity (high and low), sampling stages (d07 and d14), GxY (interaction between plant genotype and year), and interaction between soil and GxY Significant values and their associated percentage of variance

 $(R²)$ are in bold and followed by an asterisk (p <0.05).