The Composition and Phosphorus Cycling Potential of Bacterial Communities Associated With Hyphae of *Penicillium* in Soil Are Strongly Affected by Soil Origin

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Intimate fungal-bacterial interactions are widespread in nature. However the main drivers for the selection of hyphae-associated bacterial communities and their functional traits in soil systems remain elusive. In the present study, baiting microcosms were used to recover hyphae-associated bacteria from two *Penicillium* species with different phosphorus-solubilizing capacities in five types of soils. Based on amplicon sequencing of 16S rRNA genes, the composition of bacterial communities associated with *Penicillium* hyphae differed significantly from the soil communities, showing a lower diversity and less variation in taxonomic structure. Furthermore, soil origin had a significant effect on hyphae-associated community composition, whereas the two fungal species used in this study had no significant overall impact on bacterial community structure, despite their different capacities to solubilize phosphorus. However, discriminative taxa and specific OTUs were enriched in hyphae-associated communities of individual *Penicillium* species indicating that each hyphosphere represented a unique niche for bacterial colonization. Additionally, an increased potential of phosphorus cycling was found in hyphae-associated communities, especially for the gene *phnK* involved in phosphonate degradation. Altogether, it was established that the two *Penicillium* hyphae represent unique niches in which microbiome assemblage and phosphorus cycling potential are mainly driven by soil origin, with less impact made by fungal identity with a divergent capacity to utilize phosphorus.

Keywords: *Penicillium*, soil microcosm, hyphae-associated bacterial community, soil origin, phosphorus cycling gene
INTRODUCTION

In the dynamic soil environment, highly diverse bacterial and fungal communities co-exist (Nazir et al., 2013). Although there is competition between fungi and bacteria for soil resources, the presence of fungal hyphae provides colonization sites for bacteria, and creates new microhabitats with abundant and available carbon substrates (de Boer et al., 2005; Halsey et al., 2016). Approaches to identify bacteria enriched by the presence of fungal hyphae include studies of soil affected by hyphae compared to hyphae-free soils (Nazir et al., 2010, 2013; Nuccio et al., 2013; Wang et al., 2016), as well as baiting approaches for the identification of bacteria colonizing fungal hyphae in laboratory (Scheublin et al., 2010) or close-to-natural soil conditions (Ghodsalavi et al., 2017). According to these studies, environmental differentiation between the hyphosphere and the bulk soil leads to different compositions of hyphae-associated and bulk soil bacterial communities, with genera from β-Proteobacteria, γ-Proteobacteria, and Firmicutes commonly enriched in the hyphosphere or soil influenced by fungal hyphae.

Despite increasing knowledge of the composition of hyphae-associated microbiomes, the factors affecting assembly of these hyphae-associated bacterial communities remain elusive. Ballhausen et al. (2015) found differences between bacteria adhering to hyphae of Trichoderma, Mucor, and Rhizoctonia exposed to bacteria extracted from rhizosphere soil, while weaker effects of the identity of four arbuscular mycorrhizal fungi (AMF) on hyphae-associated bacterial communities have been observed using a related approach (Scheublin et al., 2010). In more realistic soil systems, the identity of AMF and ectomycorrhizal fungi has been shown to influence the composition of bacterial communities associated with hyphal bundles or ectomycorrhizal roots (Boersma et al., 2009; Marupakula et al., 2016). Abiotic factors of the soil environment could also be important for shaping fungal-associated bacterial communities (Filonev and Arora, 1987; Rousk et al., 2009), as shown for ectomycorrhizal roots where soil pH, soil carbon (C), nitrogen (N) and phosphorus (P) content as well as C:N ratios have a strong effect on bacterial communities (Simon et al., 2016; Marupakula et al., 2017).

Fungal hyphae are not only the perfect sites for bacterial colonization and dispersal, but also serve as hotspots for fungal-bacterial interactions and nutrient cycling (Furuno et al., 2012). In the hyphosphere, carbon flow from fungal hyphae to hyphae-associated bacteria could occur through the consumption of hyphal exudates, grazing on living hyphae or degradation of senescent hyphae (Drigo et al., 2010; Jansa et al., 2013). From the perspective of mutualistic interactions, bacteria attracted by hyphae could supply fungi with nutrients such as N and P. Previous studies have shown that fungi select bacterial communities with a higher efficacy for utilizing fungal-derived carbon compounds and also with a greater potential for mineral weathering (Uroz et al., 2007, 2011). Another type of evidence of hyphae serving as hotspots is the frequent appearance of N-fixing bacteria or P-solubilizing bacteria (PSB) enhancing the capacities of nutrient cycling in the hyphosphere (Zhang et al., 2014; Zagryadskaya, 2017). In relation to P turnover, Zhang et al. (2014) demonstrated that the high C:P ratio in the hyphosphere may stimulate the recruitment of bacteria with an enhanced ability to mobilize soil (in)organic P, while other studies suggest competition for P between mycorrhizal fungi and PSB in the hyphosphere soil (Brooks et al., 2011; Wang et al., 2016).

Penicillium fungi are ubiquitous in soil (Egbuta et al., 2016) and are considered a key group in P cycling (Harvey et al., 2009; Richardson et al., 2011). In part this is due to their ability to solubilize inorganic P through the release of organic anions (Richardson et al., 2011), although with considerable differences between species (Wakelin et al., 2004). Elite strains have been developed into commercial biofertilizer inoculants for the improvement of plant P nutrition (Richardson et al., 2011). However, for Penicillium, the effects of soil conditions and fungal identity on bacterial community composition and their potential role in P cycling are largely unknown.

The objectives of the current study were therefore to investigate the impacts of soil origin on (1) the community structure and (2) the P-cycling potential of hyphae-associated bacteria, assembled on two Penicillium isolates with different P-cycling potential. Baiting microcosms with P-solubilizing P. canescens or non-solubilizing P. janthinellum were established to recover hyphae-associated bacteria under close-to-natural conditions. Soils differing in physicochemical properties, including P availability, were used to constitute a diverse base for hyphal selection of bacterial communities.

MATERIALS AND METHODS

Soil Microcosm Setup and Sample Collection

Penicillium canescens ATCC 10419 (Pc) and Penicillium janthinellum ATCC 10455 (Pj) were originally isolated from soil in the United Kingdom and Nicaragua respectively. The strains differ in their P-solubilizing capacities. Pj produced 1,343 ± 40 mg/L PO$_4^{3-}$ in liquid culture with 5.4 g/L hydroxyapatite, while almost no solubilized phosphorus was determined for Pc (0 ± 21 mg/L PO$_4^{3-}$). Strains were grown for 14 days at room temperature with shaking at 300 rpm, in a medium containing: Sucrose 10.0 g/L, NH$_4$Cl 0.4 g/L, KNO$_3$ 0.78 g/L, NaCl 0.1 g/L, MgSO$_4$·7H$_2$O 1.0 g/L, CaCl$_2$·2H$_2$O 0.1 g/L, Hydroxyapatite 5.41 g/L. At this time, soluble PO$_4^{3-}$ was measured by the BioVision Phosphate Colorimetric Assay (BioVision, Mountain View, CA, United States) (data kindly provided by Drs. Dave Greenshields and Michael Furlan, United States).

To ensure distinct bacterial communities for hyphal recruitment, five soil samples from different geographical locations, which had received either inorganic fertilizer or organic manure amendments, and differing in P availability, pH and texture, were used in this study. S$_1$ was sampled from the LTNDT (Long-Term Nutrient Depletion Trial) N$_1$P$_1$K$_1$ field at University of Copenhagen, Denmark (Van der Bom et al., 2017). S$_2$ and S$_3$ were sampled from the topsoil (5–15 cm) and subsoil (30–45 cm), respectively, of fields amended with animal manure at Sjællands Odde in northern Zealand, Denmark (Paulin et al.,...
µtransferred to sterile polyamide mesh bags (50 removed and the glass slips covered with fungal hyphae were plates. After 3 days’ incubation at 26°C, the fungal plugs were placed on sterilized glass slips (25 mm, VWR International, Bridgeport, NJ, United States) with the fungal hyphae facing down. The slips were placed on water agar plates. After 3 days’ incubation at 26°C, the fungal plugs were removed and the glass slips covered with fungal hyphae were transferred to sterile polyamide mesh bags (50 µm, Sintab Produkt AB, Sweden). Mesh bags with hyphae-covered glass slips were buried in the middle of a Petri dish filled with 90–to 100 g of soil, and incubated at 26°C. Glass slips were recovered from soil microcosms and washed twice gently using 500 µl MilliQ water to remove the free-living bacteria. For DNA extraction, Penicillium hyphae and associated bacteria on glass slips were subsequently harvested by scraping off hyphae with a scalpel while washing with sterilized MilliQ water. Biological triplicates were collected for DNA extraction, where the suspensions from 10 glass slips were pooled independently, and at least two glass slips were processed per fungal species. Stained hyphae, spores and bacteria in the suspensions from 10 glass slips were processed as described above and used for soil DNA extraction. Sterilized glass slips without fungal inoculum were used as negative controls.

**Hyphal Viability and Bacterial Colonization**

Glass slips covered with growing hyphae were collected from soil microcosms after 0, 4 and 8 days’ incubation, and used to check hyphal structure and viability. Calcofluor® White M2R (CFW) and the FUN® 1 from LIVE/DEAD® Yeast Viability Kit (Molecular Probes, Eugene, OR, United States) were used separately since the interference of CFW and FUN 1 staining was observed for Penicillium hyphae. To stain fungal cell wall chitin, 30 µl of 25 µM CFW was added on top of the slips to flood fungal hyphae, and incubated in the dark at room temperature for 15 min. Hyphae flooded with 25 µM FUN 1 were incubated at 30°C in the dark for 1 h before checking the metabolic state of the hyphae. To detect bacterial attachment at day 8, 20 µl of SYBR Green solution (100 times dilution of the stock, Invitrogen Life Technologies, Carlsbad, CA, United States) was added and incubated at room temperature for 5 min. Each staining process was carried out independently, and at least two glass slips were processed for each fungal species. Stained hyphae, spores and bacteria were observed using an epifluorescence microscope (Axioskope 2, Zeiss, Oberkochen, Germany) equipped with a FITC filter (Exciter BP 450–490 nm, emission long pass LP 520 nm) and a DAPI filter (Exciter filter G 365 nm, emission long pass LP 420 nm) for FUN 1, SYBR Green, and CFW staining respectively. All images were observed under a 63 × oil objective lens (Carl Zeiss), captured with a Zeiss AxioCam digital camera, and processed with AxioVision Software and the ImageJ program.

**DNA Extraction, Sequencing, and Data Analysis**

Soil DNA was extracted from 0.25 g microcosm bulk soil using the PowerSoil® DNA Isolation Kit (MoBio Laboratories, Inc., Carlsbad, CA, United States). DNA from the suspensions of Penicillium hyphae and associated bacteria were extracted using the DNeasy Blood & Tissue kit (Qiagen, Santa Clarita, CA, United States) following the protocol for Gram-positive bacteria. The 16S rRNA gene V3–V4 region was amplified using primers Bakt_341F (CCTACGGGNGGCWGCAG) and Bakt_805R (GACTACHVGGGTATCTAATCC), purified and sequenced by the Illumina MiSeq platform as described previously (Ghodsalavi et al., 2017). Raw sequence data were submitted to the GenBank Sequence Read Archive with SRA accession: SRP132339.

The paired-end reads (>200 bp) were merged and quality screened using PANDAseq with an assembly quality score of 0.9 (Masella et al., 2012), generating 1,904,155 merged sequences with 462 ± 6 bp. Downstream sequence analysis was processed using the QIME pipeline, including de novo OTU (Operational Taxonomic Unit) clustering by UCLUST (97% similarity threshold), representative sequence picking, and taxonomic assignment by UCLUST against Greengenes 13_8 (Caporaso et al., 2010). The OTU table was filtered to exclude singletons and taxa identified as Chloroplasts and Mitochondria. Phyloseq files were generated using R version 3.3.2 with the phyloseq package (McMurdie and Holmes, 2013), and further used to calculate alpha diversity indexes after rarefaction to an even depth of 8,903 reads. NMDS (non-metric multidimensional scaling ordination) and PERMANOVA (permutational multivariate analysis of variance, 10,000 permutations) were performed based on a weighted UniFrac distance of OTU matrix using vegan package in R. Hierarchical cluster analysis (Ward’s method) was performed based on the Bray–Curtis dissimilarity using the mean relative abundances of bacterial OTUs at phylum and genus level. To identify taxonomic biomarkers with significant differential abundance among interested samples, LEfSe (linear discriminant analysis effect size) was performed using Kruskal-Wallis and pairwise Wilcoxon tests at a significance level of P < 0.05 (Segata et al., 2011). The effect size of differentially abundant taxa was estimated by LDA (linear discriminant analysis), with the logarithmic LDA score above 2 as the threshold for discriminative features. All LEfSe analysis was performed using the Galaxy framework1. Identification and visualization of differentially abundant OTUs enriched in the soil, P. canescens and P. janthinellum hyphae were performed in R using the code published by Żgadzaj et al. (2016), where a negative binomial

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1http://huttenhower.sph.harvard.edu/lefse/
GLM (generalized linear model) was employed with a false discovery rate (FDR) < 0.05, and on the OTUs with relative abundance above 0.5%. Variance partitioning and partial redundancy analysis (pRDA) were carried out using vegan package in R to explain the contribution of soil conditions and fungal identities to the variation in hyphae-associated bacterial communities. Mantel test was used to explore the correlation between the Euclidean distance of edaphic factors and Bray–Curtis dissimilarity of OTU matrix from hyphae-associated bacterial community in R (vegan package). To determine the relationships between relative abundance of bacteria phyla and edaphic factors, redundancy analysis (RDA) was applied using Canoco version 4.5. Student's t-test was used to determine the significant difference in relative abundance of major phyla/genera with relative abundance above 2%, between soil and hyphae-associated samples. P cut-off values were adjusted to account for multiple testing. One-way ANOVA followed by the Tukey's HSD test and Pearson correlation analysis was performed in SPSS Statistics 22 (IBM). Networks of significant positive correlations (P < 0.05) were visualized using Cytoscape 3.5.1 (Shannon et al., 2003). All graphs were annotated using Adobe Illustrator (Adobe Systems, United States).

Quantification of Phosphorus Cycling Genes
16S rRNA genes and genes involved in phosphorus cycling (bpp, phoX, pHoD, phnK, ppX, and ppqC) were quantified by qPCR using the Mx3000P qPCR system (Agilent Technologies, Santa Clara, CA, United States). Twenty-µl reaction mixtures were prepared with 10 µl Brilliant III Ultra-Fast SYBR® Green Low ROX qPCR Master Mix (Agilent Technologies, Santa Clara, CA, United States), 1 mg/ml BSA (bovine serum albumin, Thermo Fisher Scientific, Waltham, MA, United States), 0.4 µM of each primer, and 2 µl of template DNA (1 to 10 ng/µl). Thermal cycling was as follows: an initial cycle of 95°C for 3 min, followed by 40 cycles each of 95°C for 20 s and 58°C for 30 s. A dissociation curve was generated at the end of the qPCR program to verify a specific melting temperature of the amplified product, by including a cycle of 95°C for 1 min, 55°C for 30 s, and finally reaching 95°C by increments of 0.5°C s⁻¹, each followed by a fluorescence acquisition step. Primers used for PCR reactions are listed in Supplementary Table S2. Absolute abundance was calculated based on a standard curve for each target gene. Standard plasmids with the insertion of target genes were constructed using the TOPO® TA Cloning® Kit (Invitrogen, United States). Ten-fold serial dilutions of standard plasmid were included in each qPCR run for each target gene. Detailed descriptions on plasmid construction and calculations can be found in Supplementary Material. The abundance of functional genes was finally normalized to the 16S rRNA gene to calculate the normalized relative abundance (copy numbers per 16S rRNA gene). Student’s t-test was used to analyze significant differences in relative abundance of P-cycling genes between soil and hyphae-associated samples (P < 0.05).

RESULTS
Hyphal Growth, Viability, and Colonization by Bacteria in Soil
The presence of *P. canescens* (*Pc*) and *P. janthinellum* (*Pj*) hyphae on the surface of cover slips was observed during incubation lasting 8 days in S1 microcosms (Figure 1A and Supplementary Figure S1). For *Pc*, germination of spores led to a large amount of thin, metabolically active hyphae forming a densely cross-linked network with old hyphae at day 8 (Figures 1A–b,c). For *Pj*, a mixture of metabolically active and inactive spores was observed, most of which did not germinate even after 8 days (Figures 1A–f). Most old hyphae were found to be inactive/dead (Figures 1A–h), however active hyphae were observed in *Pj* samples by day 8 (Figures 1A–g). Taken together, a mixture of metabolically active and inactive hyphae and spores was observed for both *Penicillium* species, with hyphae and spores of *Pc* appearing more active than those from *Pj*. Despite differences in fungal activity, both *Pc* and *Pj* hyphae were densely colonized by bacteria with various morphologies and growth forms (aggregates, chains, biofilms) after 8 days of incubation in all five soils employed in the study (Figure 1B). Control glass slips without fungal hyphae showed only minor colonization by soil bacteria (data not shown), as also previously observed (Ghodsalav et al., 2017).

Diversity of Soil and Hyphae-Associated Bacterial Communities
Hyphae-associated bacterial communities were compared to soil communities for five different soils (Table 1). Soils included three top soils (S1, S2, and S3) with different texture, pH and soil nutrient contents; further two sub-soils related to the S2 top soil were included, one of which (S3) had received an amendment with inorganic P in the laboratory prior to establishment of soil microcosm (Supplementary Table S1). Bacterial communities associated with hyphae had significantly lower OTU numbers (decreased by 69 ± 19%) and less diversity than soil communities according to both Chao 1 richness and Shannon diversity indices (Table 1). However, neither index varied significantly between *Pc* and *Pj* hyphae-associated samples in any soil. Additionally, P addition to the S3 subsoil did not significantly affect the richness and diversity of the soil (S1 vs. S3, P > 0.05) or hyphae-associated bacterial communities (*Pc/S1* vs. *Pj/S3*, P > 0.05).

The NMDS ordination of OTU-based weighted UniFrac distances revealed that the three top soils S1, S2, and S3 did not cluster, while the S3 and S4 subsoils tended to cluster with the S2 top soil from the same location (Figure 2A). Furthermore, it was revealed that *Penicillium* hyphae recruited bacterial communities that were significantly distinct from soil communities (PERMANOVA, $F = 27.23, \text{P} = 9.99 \times 10^{-5}$) (Figure 2A). Only around 9% of OTUs from bulk soils were present in hyphae-associated bacterial communities (Table 1). PERMANOVA based on a weighted UniFrac distance of the OTU matrix indicated that soil origin exhibited a strong and significant impact on hyphae-associated bacterial communities ($F = 21.57, \text{P} = 9.99 \times 10^{-5}$), while the effect of fungal species was not significant ($\text{F} = 0.89, \text{P} = 0.45$) (Figure 2B). However,
a significant interaction effect was observed between soil origin and fungal species (two-way PERMANOVA, $F = 2.06$, $P = 0.02$), suggesting that the impacts of either soil origin or fungal species on recruitment were interdependent.

**Effect of Edaphic Factors on Assembly of Hyphae-Associated Bacterial Communities**

Variation partitioning, based on pRDA, revealed that soil properties and fungal identities explained 57.7 and 2.7% of the total variation in hyphae-associated bacterial communities respectively. The remaining 39.6% of the variation was unexplained (Supplementary Figure S2). Mantel test further confirmed the correlation between edaphic factors distance and bacterial community dissimilarity ($P = 0.001$). As soil properties accounted for most of the observed variation in hyphae-associated bacterial communities, the effect of specific edaphic factors on relative abundances of phyla associated with hyphae was further investigated by RDA (Figure 2C). The first two axes of the RDA explained 48.5 and 16.4% of the total variation in the data. Soil pH and $\text{Mg}^{2+}$ were positively correlated with the relative abundance of Proteobacteria (Pearson correlation coefficient: $r = 0.74$, $P = 2.97 \times 10^{-6}$ and $r = 0.65$, $P = 9.16 \times 10^{-5}$ for pH and $\text{Mg}^{2+}$ respectively) and Actinobacteria ($r = 0.84$, $P = 5.54 \times 10^{-9}$ and $r = 0.79$, $P = 1.87 \times 10^{-7}$ for
### TABLE 1 | Origin, sequencing information, and alpha diversity of samples used in this study.

| Sample | Origin | Soil microcosms | No. of sequences | No. of all OTUs | Decreased OTUs (%) | No. of shared OTUs with soil (%) | Percentage of shared OTUs with soil (%) | Shannon | Chao1 |
|--------|--------|-----------------|-----------------|----------------|-------------------|---------------------------------|----------------------------------------|---------|-------|
| S1     | Soil 1 | LTNDT N/P/K1    | 34,684 ± 566    | 5,873 ± 88^D  | –                 | –                               | –                                       | 6 ± 0^R | 4,844 ± 57^C |
| S2     | Soil 2 | Sjaelland Odde Org.A | 48,622 ± 1,794 | 10,035 ± 186^A | –                 | –                               | –                                       | 7 ± 0^A | 7,349 ± 106^A |
| S3     | Soil 3 | Sjaelland Odde Org.B | 42,113 ± 1,092 | 8,269 ± 193^B | –                 | –                               | –                                       | 7 ± 0^A | 6,048 ± 220^B |
| S4     | Soil 4 | Sjaelland Odde Org.B + 60 mg/kg P (NaH₂PO₄·H₂O) | 40,130 ± 1,086 | 8,145 ± 149^mIB | –                 | –                               | –                                       | 7 ± 0^A | 6,262 ± 172^B |
| S5     | Soil 5 | Jyndevad A     | 38,111 ± 3,890  | 7,052 ± 701^C | –                 | –                               | –                                       | 7 ± 0^A | 5,577 ± 551^BC |
| P1S1   | P1 Hyphae | P1 + LTNDT N/P/K1 | 61,095 ± 3,704 | 3,003 ± 294^EF | 49                | 411                             | 7                                       | 4 ± 0^F | 2,076 ± 103^DE |
| P1S2   | P1 Hyphae | P1 + Sjaelland Odde Org.A | 51,560 ± 2,302 | 2,858 ± 244^FG  | 51                | 468                             | 8                                       | 4 ± 0^DE | 2,126 ± 223^GE |
| P2S2   | P1 Hyphae | P2 + Sjaelland Odde Org.A | 22,151 ± 3,202 | 1,455 ± 304^HI  | 86                | 425                             | 5                                       | 5 ± 0^CD | 1,247 ± 202^EF |
| P2S3   | P1 Hyphae | P2 + Sjaelland Odde Org.B | 24,937 ± 4,074 | 1,842 ± 264^GH  | 82                | 477                             | 5                                       | 5 ± 0^CD | 1,581 ± 205^EF |
| P3S3   | P2 Hyphae | P3 + Sjaelland Odde Org.B | 19,320 ± 4,167 | 1,255 ± 260^H   | 85                | 354                             | 4                                       | 4 ± 0^E  | 1,110 ± 226^F  |
| P3S4   | P2 Hyphae | P3 + Sjaelland Odde Org.B + 60 mg/kg P (NaH₂PO₄·H₂O) | 20,669 ± 6,349 | 1,296 ± 310^H   | 84                | 338                             | 4                                       | 4 ± 0^E  | 1,169 ± 249^F  |
| P4S4   | P2 Hyphae | P4 + Sjaelland Odde Org.B + 60 mg/kg P (NaH₂PO₄·H₂O) | 21,828 ± 4,759 | 1,106 ± 111^H   | 45                | 1530                            | 22                                      | 4 ± 0^E  | 938 ± 41^F     |

(1) \(P_c\) and \(P_i\) are Penicillium canescens and Penicillium janthinellum respectively. All data are presented as the mean ± standard deviation of three replicates. Different superscript capital letters represent significant difference among samples (one-way ANOVA, \(P < 0.01\)). (2) \(^a\)Compared with OTUs in each corresponding soil sample.
pH and Mg\(^{2+}\) respectively), and negatively correlated with the relative abundance of Bacteroidetes in hyphae-associated bacterial samples \((r = -0.74, P = 2.35 \times 10^{-6}\) and \(r = -0.71, P = 1.11 \times 10^{-5}\) for pH and Mg\(^{2+}\) respectively). Other phyla including Gemmatimonadetes, Verrucomicrobia, Acidobacteria, Planctomycetes and WS3 were associated with Olsen P and TC according to RDA analysis \((\text{Figure 2C})\).

**Composition of Soil and Hyphae-Associated Bacterial Communities**

In soils, Proteobacteria was the most abundant phylum, accounting for 23–39% of all OTUs \((\text{Figure 3A})\). Other dominant phyla, including Gemmatimonadetes, Verrucomicrobia, Bacteroidetes, Actinobacteria, and Acidobacteria, varied significantly with soil origin \((\text{Supplementary Table S3})\), as also reflected in the clustering of the samples \((\text{Figure 3A})\). Hyphae-associated communities were also dominated by Proteobacteria, with other dominant phyla being Actinobacteria, Bacteroidetes, and Firmicutes \((\text{Figure 3A})\). Compared to soil communities, all the hyphae-associated bacterial communities, except for \(P_c/S_1\), presented a significantly higher relative abundance of Proteobacteria \((\text{Supplementary Table S4})\). Bacteroidetes was only found to have significantly higher relative abundances in \(P_j/S_1\) \((P = 1.89 \times 10^{-4}\)), and the relative abundance of Firmicutes was significantly higher in hyphae-associated communities from \(P_c/S_2/S_5\) \((P = 4.85 \times 10^{-4}\) / \(P = 4.85 \times 10^{-4}\)). At the genus level, the relative abundance varied strongly with soil origin \((\text{Figure 3B})\). A significant increase of major genera was observed in the hyphae-associated bacterial communities as compared with the corresponding soil background. In the \(P_c/S_1\) communities, *Chitinophaga* \((P = 0.003/P = 0.041\) for the \(P_c/S_1\) and \(P_j/S_1\) communities respectively), *Burkholderia* \((P = 0.002/P = 6.63 \times 10^{-7}\) and *Niastella* \((P = 0.006/P = 0.032\) were significantly enriched, while *Rhodanobacter* \((P = 4.57 \times 10^{-10}/P = 3.35 \times 10^{-5}\), *Paenibacillus* \((P = 0.007/P = 0.005\), *Planctomyces* \((P = 8.04 \times 10^{-6}/P = 1.98 \times 10^{-4}\), and *Sporocytophaga* \((P = 0.001/P = 3.33 \times 10^{-4}\) were significantly enriched.
in the P_c/S_5 communities. The relative abundance of *Phyllobacterium* (*P* = 0.033/*P* = 0.031), *Lysobacter* (*P* = 1.59 × 10^{-4}/*P* = 5.19 × 10^{-5}), *Paenibacillus* (*P* = 0.003/*P* = 0.02), *Agromyces* (*P* = 0.01/*P* = 0.004), and *Bacillus* (*P* = 0.02/*P* = 0.02) increased significantly in P_c/S_2. P amendment to S_3 (resulting in S_4) had a greater influence on hyphae-associated communities than on soil communities, especially for communities attracted to P_c-associated communities than on soil communities, especially *Lysobacter* to S_3 (*P* = 0.12), *Delftia* (*P* = 0.49), and *Variovorax* (*P* = 0.12) in P_c/S_3 communities was lower as compared with those in P_c/S_4, accompanied by higher relative abundance of *Paenibacillus* (*P* = 0.18), *Agromyces* (*P* = 0.47), and *Phyllobacterium* (*P* = 0.79).

Although the two fungal species used in this study showed no significant difference in their overall impact on bacterial community structure, LEfSe comparisons of combined data from hyphae-associated communities across all soils showed different discriminative taxa for P_c hyphae. The abundance of *Lysobacter* (*P* = 0.80), *Pseudomonas* (*P* = 0.43), *Pedobacter* (*P* = 0.12), *Delftia* (*P* = 0.49), and *Variovorax* (*P* = 0.12) in P_c/S_3 communities was lower as compared with those in P_c/S_4, accompanied by higher relative abundance of *Paenibacillus* (*P* = 0.18), *Agromyces* (*P* = 0.47), and *Phyllobacterium* (*P* = 0.79).

Although the two fungal species used in this study showed no significant difference in their overall impact on bacterial community structure, LEfSe comparisons of combined data from hyphae-associated communities across all soils showed different discriminative taxa for P_c and P_j in the same soil (Supplementary Figure S3). The selection of specific OTUs by the two fungal species was also visualized by ternary plot analyses showing that, compared to soil samples, only a small number of OTUs (less than 50 in each sample) were enriched by fungi (Figure 4). Furthermore, some enriched OTUs were only abundant in P_c or P_j, indicating that each *Penicillium* species could select specific bacterial species onto their hyphae (orange and green circles in Figure 4 and Supplementary Figure S4). Taxonomic assignments for enriched OTUs at order level generally revealed a more diverse assembly in P_c communities than in P_j communities (pie charts in Figure 4). A large number of OTUs were abundant in both P_c and P_j samples, indicating a general fungal effect (gray circles in Figure 4).

**Distribution and Abundance of P-Cycling Genes in Soil and Hyphae-Associated Bacterial Communities**

To determine if *Penicillium* hyphae recruit bacterial communities with a different potential for P cycling than that of soil communities, the relative abundance of selected P-cycling genes was determined by means of qPCR (Figure 5). In soil communities, *bpp*, *phoX*, and *phoD* genes varied comparably with significantly higher relative abundances in S_3, S_4, and S_5 than in S_1 and S_2 (*P* < 0.05), while *phnK* had a significantly higher abundance in S_3 and S_4 as compared to S_1, S_2, and S_5 (*P* < 0.05). In most hyphae-associated samples, the relative abundance of *phnK* was significantly higher than in soil (*P* < 0.05). The relative abundance of *bpp*, *phoX* and *phoD* genes showed a considerable variation between hyphae-associated samples in the different soils. Compared with soil samples, both *phoX* and *bpp* genes were more abundant in hyphae-associated samples from S_1, S_2, and S_5. The *phoD* gene had significantly lower relative abundances in most hyphae-associated samples than in soil (*P* < 0.05), except for samples from S_1 microcosms. For the inorganic P-cycling genes, only the *pqqC* gene was detectable, whereas the *ppX* gene was below the detection limit of the qPCR assay for both soil and hyphae-associated communities. The *pqqC* gene had a higher relative abundance in hyphae-associated samples as compared to the bulk soil in S_3 and S_4, but had a significantly lower abundance in hyphae-associated samples in S_5 (*P* < 0.05).

Network analysis showed a shift in co-occurrence patterns between bacterial taxa and phosphorus-cycling genes from soil to hyphae-associated bacterial communities (Supplementary Figure S5). In addition, a greater diversity of potential host taxa affiliating with the *phnK* gene was observed in hyphae-associated samples as compared to bulk soil samples.

**DISCUSSION**

The significance of soil origin on the composition of hyphae-associated bacterial communities is currently not well-understood. In the present study, the application of a newly developed microcosm approach (Ghodsalavi et al., 2017) provided novel insight on the importance of soil origin as a driver for hyphae-associated bacterial community assembly, both at the taxonomic level and for the functional potential in relation to P cycling. *Penicillium*, a fungal genus that encompasses important P-solubilizing biofertilizers (Antoun, 2012), as bait,
the colonization of hyphae by soil bacteria forming aggregates was observed, some of which resembled biofilms. While the ability of individual bacteria to form biofilms on hyphae is well-documented under laboratory conditions (Hogan and Kolter, 2002; Toljander et al., 2006; Brand et al., 2011), biofilm formation by bacterial communities on hyphae in soil is poorly documented (Guennoc et al., 2018).

The composition of hyphae-associated communities was distinctly different from that of the soil communities, and they were less diverse. These results are in agreement with previous studies showing shifts in bacterial communities in mycosphere soil versus bulk soil (Warmink and van Elsas, 2008; Gahan and Schmalenberger, 2015), however with a more pronounced decrease in diversity than previously observed. Consistently, only few phyla (Proteobacteria, Bacteroidetes, Firmicutes, and Actinobacteria) accounted for over 90% of all OTUs throughout all hyphae-associated microbiomes except the $P_cP_jS_5$, where these phyla accounted for 70% of the hyphae-associated microbiomes.
FIGURE 5 | Distribution and relative abundance of phosphorus-cycling genes in soil and hyphae-associated communities. The relative abundance of genes was calculated by the ratio of absolute copy numbers of target genes to 16S rRNA genes in each sample. All data were presented as the mean ± standard error of three replicates. Significant differences in relative abundance of P-cycling genes between soil and hyphae-associated samples were analyzed in each microcosm and are indicated by stars (Student’s t-test, $P < 0.05$).

(Figure 3A). Despite the low diversity observed, the hyphae-associated communities identified in the current study were more diverse than the community attached to P. bilaii hyphae in a comparable set-up, which was dominated (>90%) by Proteobacteria (Ghodsalavi et al., 2017).

Proteobacteria and Bacteriodetes are classified as copiotrophic groups which are commonly detected in carbon-enriched environments such as the rhizosphere (Yousuf et al., 2012; Aislabie and Deslippe, 2013). Here, we document the same overall recruitment of these groups to saprophytic fungi under soil conditions. When related to edaphic factors, hyphae-associated Bacteriodetes was found to correlate negatively with soil pH in the present study, while Proteobacteria and Actinobacteria were found to correlate positively with higher soil pH ($P < 0.05$). For Bacteriodetes, this correlation was mainly driven by the specific enrichment of this group in the hyphosphere of both Penicillium species in the low-pH soil from the Long Term Nutrient Depletion Trial, Denmark (S1). A strong positive correlation has previously been found between soil pH and Bacteriodetes (Lauber et al., 2009). However, in a study at the Hoosfield acid strip (Rousk et al., 2010), where pH is the main factor defining a soil gradient, no such correlation was found. The authors concluded that the often found correlation between pH and Bacteriodetes could be driven by derived factors such as C-availability. The two Penicillium species from the current study are known to excrete organic acids (Keromnes...
Penicillium species with diverging ability to solubilize P under laboratory conditions, showed no significant difference in overall composition and diversity of their respective hyphae-associated microbiomes. This result is consistent with the study of Scheublin et al. (2010) on four closely-related Glomus isolates, while less phylogenetically-related fungi, i.e., Trichoderma, Mucor and Rhizoctonia, attract different hyphae-attached bacterial communities (Ballhausen et al., 2015; Rudnick et al., 2015). The composition of hyphae-associated bacterial communities may in part reflect differences in the composition of the hyphal cell walls and hyphal exudates (Rudnick et al., 2015), which would be expected to differ increasingly with a greater phylogenetic distance between fungal species. Other cases in which fungal fruiting-bodies were not directly in contact with soils showed that fungal phylogenetic variation was more important in structuring associated microbiomes than soil characteristics (Pent et al., 2018). Nevertheless, in the current study, the hyphae of individual Penicillium species represent unique niches for several specific hyphae-associated bacterial taxa, which could be due to the differences in fungal physiology and metabolic properties such as fungal exudate production, antibiotics production and even P solubilization capacity.

Fungal hyphae are reported as potential hotspots for nutrient cycling (Furuno et al., 2012). In this study, we observed a general enrichment of phnK genes in hyphae-associated bacteria from all soils, and a greater diversity of host taxa affiliated with phnK in the hyphae-associated microbial communities. Phosphonates can serve as a phosphorus source following bacterial C-P lyase activity, although these compounds are some of the least labile organic phosphorus (Po) sources, and their utilization has primarily been reported in marine systems (Dyhrman et al., 2009; Yu et al., 2013). Potential phosphonate sources used by hyphae-associated bacteria are components of fungal exopolysaccharides, glycoproteins and membrane phosphonolipids (McGrath et al., 2013), or phosphonate-containing antimicrobial compounds released by Penicillium hyphae or Streptomyces, which are often abundant around hyphae (Watanabe et al., 1999; Metcalf and van der Donk, 2009). In addition to the increased potential for phosphonate degradation, a relatively high abundance of the bpp gene encoding a beta-propeller phytase as well as the phoX gene encoding a phosphatase were observed in hyphae-associated samples from topsoils with a greater organic content (S1, S2, and S3). Hydrolysis of phytate yields phosphoesters, which could be further hydrolyzed by a diverse group of phosphatases such as extracellular and periplasmic phosphatases (PhoX) or cytoplasmic phosphatases (PhoD) (Luo et al., 2009; Acuna et al., 2016; Neal et al., 2017). Interestingly, phoD genes were only enriched in the hyphae-associated community in S1, whereas a relative decrease was observed for all other soils. The phoX and phoD genes are important for Po recycling in rhizosphere soils (Acuna et al., 2016), but their occurrence and abundance in the hyphosphere are largely unexplored. The present study suggests a more pronounced potential ecological role of PhoX, as compared to PhoD, in the recycling of organic P around hyphae. Co-occurrence patterns identified Pseudomonas as the only potential host taxon affiliated with the bpp gene in hyphae-associated bacterial communities. This result was in accordance with a previous study showing that Pseudomonas is active in phytate-P turnover in the AMF hyphosphere (Wang et al., 2016). Indications for increased utilization of inorganic P in the hyphae-associated samples were less pronounced. Increased abundance of pqqC, one of marker genes for this process, was not consistently observed in the hyphae-associated communities as compared to the soil communities.

CONCLUSION

This study demonstrated that Penicillium hyphae recruit distinct bacterial communities with contrasting P-cycling potential compared to the surrounding bulk soil inhabitants. The complex assemblage of bacterial communities associated with hyphae, in
terms of either shifting community compositions or distribution of P-cycling genes, was mainly driven by soil origin, and only secondarily by the interactions between fungal species and soil conditions. This study highlights the importance of soil in the interaction between ecologically important fungi and their associated bacteria. The mechanisms whereby soil conditions affect hyphae-associated bacteria, however, deserve further investigation in future.

**DATA AVAILABILITY STATEMENT**

The datasets generated for this study can be found in the GenBank Sequence Read Archive with SRA accession: SRP132339.

**AUTHOR CONTRIBUTIONS**

MN and ON proposed the project. Y-GZ helped to design the experiments. XH performed the experiments and did data analysis. All authors participated in the manuscript writing and revising.

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**SUPPLEMENTARY MATERIAL**

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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